Re-evaluation of Sympoventuriaceae

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Key words

evolution lifestyle multigene analysis new taxa systematics Venturia **Abstract** *Sympoventuriaceae* (*Venturiales, Dothideomycetes*) comprises genera including saprophytes, endophytes, plant pathogens, as well as important animal or human opportunistic pathogens with diverse ecologies and wide geographical distributions. Although the taxonomy of *Sympoventuriaceae* has been well studied, generic boundaries within the family remain poorly resolved due to the lack of type materials and molecular data. To address this issue and establish a more stable and reliable classification system in *Sympoventuriaceae*, we performed multilocus phylogenetic analyses using sequence data of seven genes (SSU, ITS, LSU, *act1, tub2, tef1* and *rpb2*) with increased taxon sampling and morphological analysis. The molecular data combined with detailed morphological studies of 143 taxa resolved 22 genera within the family, including one new genus, eight new species, five new combinations and one new name. Finally, we further investigated the evolutionary history of *Sympoventuriaceae* by reconstructing patterns of lifestyle diversification, indicating the ancestral state to be saprophytic, with transitions to endophytic, animal or human opportunistic and plant pathogens.

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INTRODUCTION

Sympoventuriaceae is a large family in Venturiales (Dothideomycetes, Ascomycota) with diverse ecology, wide geographic distribution and rich species diversity (Zhang et al. 2011, Seyedmousavi et al. 2013, Liu et al. 2017, Wijayawardene et al. 2018, Crous et al. 2019a, Shen et al. 2020). Members of this family are usually hyphomycetes with conidia liberated by rhexolytic secession (Seifert et al. 2011, Machouart et al. 2014, Crous et al. 2014, Huanraluek et al. 2019). Sympoventuriaceae is mainly known as a ubiquitous environmental saprobic fungus and plant endophytes or pathogens, while a few species have been documented as opportunistic neurotropic pathogens in vertebrate hosts, including humans (Satow et al. 2008, Seyedmousavi et al. 2014, Kidd et al. 2016, Zhang et al. 2018, Samerpitak et al. 2019, Benavent 2021, Murata et al. 2022). They are also known for their thermophilic properties, such as living in hot springs (Revankar & Sutton 2010, Hao et al. 2013, Samerpitak et al. 2014, 2015b, Wang et al. 2018, Crous et al. 2020).

Sympoventuriaceae was introduced by Zhang et al. (2011) with Sympoventuria (type genus), Veronaeopsis and fusicladiumlike species included. The generic organization of sequestrate taxa within the Sympoventuriaceae has long been a subject of debate, due to a high level of morphological plasticity, and the lack of molecular data (Machouart et al. 2014, Samerpitak et al. 2016). Sympoventuria was first described for a venturialike ascomycete, typified by S. capensis, a species found on decaying leaves of *Eucalyptus*, which was characterised by its saprobic lifestyle, pseudoparaphyses, and hyaline, symmetrical ascospores and subcylindrical asci (Crous et al. 2007a, b). Veronaeopsis was introduced as a monotypic genus for V. simplex, which was previously separated from Veronaea based on its shorter conidiophores, geniculate rachis and prominent conidiogenous loci (Papendorf 1969, Arzanlou et al. 2007). Morphologically, Sympoventuria is allied to Venturia (Sivanesan 1977, Zhang et al. 2011, Zhang et al. 2016), although their asexual morphs are quite distinct. For instance, Fusicladium (asexual morph of Venturia) was established by Bonorden (1851) to accommodate F. virescens, a well-known pathogen of pears. Fusicladium is characterised by sympodial conidiogenesis, differentiated conidiophores, and melanized conidia with dark basal scars. However, the taxonomy of this genus has continued to be controversial (Baldacci & Ciferri 1937, Schubert et al. 2003, Beck et al. 2005, Koukol 2010). Shen et al. (2020) resolved Fusicladium as asexual morph of Venturia, but also introduced several additional fusicladium-like genera, namely Fuscohilum, Neofusicladium, Parafusicladium and Pinaceicola.

Since its introduction, several genera have either been included or excluded from *Sympoventuriaceae*, and many mycologists commented that there might be more unrecognized genera within the family (Machouart et al. 2014, Samerpitak et al. 2016). *Scolecobasidium* (= *Ochroconis*) and *Verruconis* are very similar genera that have sympodial conidiogenous cells and T- or Y-shaped to cylindrical or clavate conidia (Abbott 1927, De Hoog & Von Arx 1973). Due to an unusual combination of morphological and ecological characters, their systematic position has historically been controversial. Samerpitak et al. (2014) distinguished these genera based on their ecological and physiological traits and morphological differences. They

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accommodated mesophilic species with smooth-walled to verruculose conidia in Scolecobasidium (as Ochroconis), and retained the thermophilic taxa with verrucose to coarsely ornamented conidia in Verruconis. Nevertheless, morphological and ecological delimitation of Scolecobasidium and Verruconis is problematic and remains obscure (Samerpitak et al. 2016, Qiao et al. 2019). Acroconidiellina was introduced by Ellis (1971) to accommodate A. arecae, A. chloridis, A. loudetiae (type species) and A. urtiagae. Hernández-Restrepo et al. (2016) further pointed out that Acroconidiellina is allied to the Scolecobasidium/Ochroconis complex, and belonged to Sympoventuriaceae (Li et al. 2016, Wijayawardene et al. 2020). Furthermore, although Acroconidiellina currently contains four species, the taxonomic placement of only A. arecae has thus far been confirmed based on phylogenetic studies. The monotypic genus Mycosisymbrium was proposed by Carris (1994), re-described by Pratibha & Prabhugaonkar (2016), initially regarded as incertae sedis in the Pezizomycotina, and later placed in Sympoventuriaceae. Following these studies, three interesting genera, Echinocatena, Matsushimaea and Yunnanomyces were analysed phylogenetically suggesting a close relationship to Sympoventuriaceae, each of which formed a monophyletic clade with other genera in this family (Crous et al. 2018a, b, Tibpromma et al. 2018). Pseudosigmoidea (typified by P. cranei) was introduced based on species of Sigmoidea with enteroblastic conidia and phialidic conidiogenesis (Ando & Nakamura 2000), and subsequent studies showed that it also resided in Sympoventuriaceae (Diene et al. 2013, Crous et al. 2019a). Hernández-Restrepo et al. (2020) reassessed the taxonomic placement of Melnikomyces to accommodate an increasing number of emerging species, and placed it in Sympoventuriaceae, together with other genera producing septate conidia from denticulate conidiogenous cells (Crous et al. 2014, Wei et al. 2020).

The classification of *Sympoventuriaceae* includes a wide range of taxa based on morphological characters, although these are chiefly asexual genera (Zhang et al. 2011, Machouart et al. 2014). However, some genera (e.g., *Clavatispora, Neocoleroa*) of *Sympoventuriaceae* were established only based on their sexual morphology, and very few links between sexual and asexual morphs have been confirmed. *Sympoventuria capensis*

was introduced with both a sexual and asexual morph (Crous et al. 2007a, Machouart et al. 2014). Subsequently, Boonmee et al. (2014) introduced Clavatispora based on the sexual morph C. thailandica. Its unique ascospores and bitunicate asci resemble Pleosporales species but differ from most other taxa in Venturiales (Seifert et al. 2011, Hyde et al. 2013, 2020). In addition, the phylogenetic analysis of conserved genes (nuSSU, nuLSU, mtSSU and rpb2) indicated that Verruconis is distinct from Scolecobasidium, while several related sexual morphs were also included in the Sympoventuriaceae (Machouart et al. 2014). Another sexual genus, Neocoleroa (typified by N. si*birica*), is characterised by lobed to dichotomously branched, blunt-tipped setae on superficial pseudothecia (Petrak 1934, Johnston & Park 2016). Morphologically, Neocoleroa is most comparable to Wentiomyces (Koorders 1907), and they have had a tangled taxonomic history (Barr 1997, Kirk et al. 2008). It is noteworthy, except for a few species of Clavatispora, Neocoleroa, Scolecobasidium, Sympoventuria and Verruconis, that the sexual morphs of most species of the Sympoventuriaceae are unknown. Moreover, the asexual and sexual morphs of Sympoventuriaceae often develop separately, or only one morph is formed, making it difficult to confirm links between morphs of the same species.

In summary, Sympoventuriaceae has been extensively reviewed in recent years in efforts to clarify the phylogeny and taxonomic relationships of its species and allied fungi, and has resulted in a modern redefinition of the family, which provides a solid foundation to facilitate future DNA phylogenetic studies (Tibpromma et al. 2018, Crous et al. 2019a, Shen et al. 2020). In spite of this, however, many questions remain unresolved about the phylogenetic relationships of some poorly documented taxa, especially genera and species for which molecular data are not yet available. This has justified an urgent need to reconsider the species boundaries for Sympoventuriaceae based on a robust family-wide phylogenetic backbone and framework. Furthermore, Sympoventuriaceae includes approximately 164 species, is a morphologically and ecologically diverse fungal group with different lifestyles and modes of nutrition (MycoBank, April 2022). In order to adapt to changing environmental conditions, their ecological habitat varies from saprobic, animal or human opportunistic

Table 1	Primers	and	PCR	conditions.

Genes	Primers	Sequences/PCR conditions	References
ITS	ITS5 (Fw) ITS4 (Rw)	5'-GGAAGTAAAAGTCGTAACAAGG-3' 5'-TCCTCCGCTTATTGATATGC-3' 95 °C 5 min, (95 °C 35 s, 56 °C 30 s, 72 °C 1 min) 35 cycles, 72 °C 4 min	White et al. (1990) White et al. (1990)
LSU	LR0R (Fw) LR5 (Rw)	5′-ACCCGCTGAACTTAAGC-3′ 5′-TCCTGAGGGAAACTTCG-3′ 95 °C 5 min, (95 °C 45 s, 56 °C 40 s, 72 °C 2 min) 35 cycles, 72 °C 10 min	Vilgalys & Hester (1990) Vilgalys & Hester (1990)
SSU	NS1 (Fw) NS24 (Rw)	5'-GTAGTCATATGCTTGTCTC-3' 5'-AAACCTTGTTACGACTTTTA-3' 95 °C 5 min, (95 °C 45 s, 56 °C 40 s, 72 °C 2 min) 35 cycles, 72 °C 10 min	White et al. (1990) Gargas & Taylor (1992)
act1	512 (Fw) 783 (Rw)	5'-ATGTGCAAGGCCGGTTTCGC-3' 5'-TACGAGTCCTTCTGGCCCAT-3' 95 °C 5 min, (96 °C 45 s, 56 °C 30 s, 72 °C 1 min) 35 cycles, 72 °C 5 min	Carbone & Kohn (1999) Carbone & Kohn (1999)
tub2	Bt2a (Fw) Bt2b (Rw)	5-GGTAACCAAATCGGTGCTGCTTTC-3 5'-ACCCTCAGTGTAGTGACCCTTGGC-3' 95 °C 5 min, (95 °C 35 s, 56 °C 50 s, 72 °C 2 min) 35 cycles, 72 °C 7 min	Glass & Donaldson (1995) Glass & Donaldson (1995)
tef1	728 (Fw) 986 (Rw) 983 (Fw) 2218 (Fw)	5'-CATCGAGAAGTTCGAGAAGG-3' 5'-TACTTGAAGGAACCCTTAC-3' 5'-GCYCCYGGHCAYCGTGAYTTYAT-3' 5'-ATGACACCRACRGCRACRGTYTG-3' 95 °C 5 min, (96 °C 45 s, 56 °C 30 s, 72 °C 45 s) 35 cycles, 72 °C 5 min	Carbone & Kohn (1999) Carbone & Kohn (1999) Rehner & Buckley (2005) Rehner & Buckley (2005)
rpb2	5 (Fw) 7CR (Rw)	5′-GAYGAYMGWGATCAYTTYGG-3′ 5′-CCCATRGCTTGYTTRCCCAT-3′ 95 °C 5 min, (96 °C 45 s, 56 °C 30 s, 72 °C 2 min) 35 cycles, 72 °C 5 min	Liu et al. (1999) Liu et al. (1999)

Species	Strain ¹	Host and substrate	Locality		GenBank a	GenBank accession numbers ²	bers ²	
				ITS	LSU	tub2	tef1	rpb2
Acroconidiallina arecae	NECCI 3696	On little patches on the leaves of Areca Catechu	India	K X306747	K X306776	I	I	I
		Looporto obtento on Ciercio da Carolia			MI/040700		90700711	007700
						I		
Clavatispora mailandiaca	MILEUCC 10010/	Un dead stems of herbaceous plants	I nalland		NF//0458	I	NF//0459	1
Echinocatena arthrinioides	CBS 144202	Acacia crassicarpa, leaves	Malaysia	MH107890	MH107937	I	I	1
Fuscohilum rhodensis	CBS 121641*	Ceratonia siliqua, branches	Greece	MK810909	MK810796	MK926471	MK888733	MK887802
Fu. siciliana	CBS 105.85*	Chamaerops humilis	Italy	MK810910	MK810797	MK926472	MK888734	MN091924
Guizhoumvces aciculaea	GUCC 18195*	Isolated from soil	China	MZ503724	MZ503757	MZ546903	MZ546870	MZ546866
	GIICC 18152	Erom leaf litter	China	M7503733	MZEORTER	MZ546902	MZEAGREG	MZEAGREE
		Distancia putration dood moticle				1000407M		
	UBS / 20.03		Australia			I	I	I
Matsushimaea fasciculata	CBS 167.97*	On dead leaf of <i>Cinnamomum japonicum</i>	Japan	LT962397	LT962402	I	I	I
	GUCC 18239	Isolated from soil	China	MZ503725	MZ503758	MZ546904	MZ546871	MZ546867
Ma monilioides	CBS 143867*	Garden soil	Snain	I T883468	I T883469	I	I	I
					MT704004	NAT720645	01720E1E	
					16710/11M	CI CAC / I M		I
Me. thailandicus	CBS 145767*	Isolated from soil	Thailand	MN794374	MN794351	I	I	I
Me. vietnamensis	CBS 136209*	On dry leaves of broadleaved tree	Vietnam	KJ869156	KJ869213	I	I	I
Mvcosisvmbrium cirrhosum	MTCC12435	On dead leaves of Vaccinium macrocarnon	United States	KR259883	KR259884	I	I	KR349124
		leolated from decoving Camellia sinensis leaf litter		M7E03722	MZED27EE	MZEACON	MTEACOCO	MTEACREA
			CIIIIa	77/002M	CC / COC7INI		00004C7M	10004C7M
Neocoleroa cameroonensis	CBS 129041 [*]	Crematogaster sp. (ant) carton on Barteria nigritana	Cameroon	MK810902	MK810789	MINU/8219	MK888/2/	MK887797
Nc. metrosideri	ICMP 21139*	On living leaves of Metrosideros excelsa	New Zealand	KU131678	KU131677	I	I	I
Neofusicladium eucalypti	CBS 128216*	<i>Eucalyptus regnans</i> , leaf litter	Australia	MK810903	MK810790	MK926468	MK888728	MK887798
Nf. eucalvoticola	CBS 141301*	Eucalvotus robusta. leaf litter	France	MK810904	MK810791	I	MK888729	MK887799
	CBS 143427	Fucalvotus dunnii Teaves	Australia	MK810905	MK810792	I	I	I
Nf rechans	CBS 143411*	Eucalyntis radians leaves	Australia	MG386066	MG386110	MG386169	I	
								I
						MIN44 2/40	MIN442122	I
Paratusiciadium amoenum	CBS 254.95	<i>Eucalyptus</i> sp., tallen leaves	Cuba	MK810906	MK810793	MK926469	MK888730	I
Pa. intermedium	CBS 110746*	<i>Eucalyptus</i> sp., leaf litter	Madagascar	MK810907	MK810794	MK926470	MK888731	MK887800
Pa. paraamoenum	CBS 141322*	Eucalyptus regnans, leaf litter	Australia	MK810908	MK810795	I	MK888732	MK887801
Pinaceicola cordae	CBS 126959*	Pinus svivestris. litter needles	Czech Republic	MK810911	MK810798	MK926473	MK888735	I
	CBS 675.82	Pinus sylvestris, litter needles	Netherlands	MK810912	MK810799	MK926474	MK888736	I
	CBS 143494	Pinus sylvestris litter needles	Germany	MK810913	MK810800	MK926475	MK888737	I
Di nini		Dinus sulvestris, litter needles	Netherlands	MK810015	MK810802		MIK888730	MK 887804
			Nemenands	MIN0 109 14		MIN920410	IVING001 30	
Pseudosigmoidea alnicola	CBS 145034*	Leaf litter of Alnus glutinosa (Betulaceae)	Germany	MK442620	MK442556	I	I	1
Ps. excentrica	CBS 469.95*	<i>Lauraceae</i> , leaf litter	Cuba	HQ667543	KF282669	MK926478	KF155975	1
Ps. ibarakiensis	NBRC 107891*	Natural forest soil	Japan	LC146758	LC146759	I	I	I
Scolecobasidium anellii	CBS 284.64*	Stalactite	Italy	FR832477	KF156138	KF156184	KF155995	KF282684
Sc. anomala	CBS 131816*	Lascaux Cave	France	HE575201	KF156137	KF156194	KF155986	HE575205
Sc. blechni	CBS 146055*	Leaves of Blechnum capense (Blechnaceae)	South Africa	MN562134	MN567641	MN556843	MN556826	I
Sc. constricta	CBS 211,53*	Soil	Canada: Ontario	HQ667519	KF156148	KF156187	KF156005	KF282686
Sc. crassihumicola	CBS 120700	Soil	Papua New Guinea	KJ867429	KJ867430	KJ867433	KJ867428	I
Sr dameii	CBS 230 78*	Canvota niumosa leaf	Sri Lanka	KE156019	KE156150	KE156190	KF155080	
	CEC 200.1 C	Carry oranges, ican Foraet soil	Cir caina Canada: Ontario	HORE7524	KE156130	KE156174	KE156000	KE282700
ou lanauxansis	CEG 330.09	r Urest soll Black stain on cave sediment	Carrada. Ortario Erance	FD832474	KE156136	KE156183	KE155003	ED832481
Oc. Ioscaukeriolo		black statisticate sedilistic						
			Japan					
Sc. musicola	CBS 144441"	Un leaves of Musa sp. (Musaceae)	Malaysia	MH32/824	MH32/860	MH32/898	MH32/88/	MH32/8/6
sc. pnaeopnora	CBS 200.90°		Papua New Guinea	KP/98631	KP/98034	K12/2062	K12/2098	KF 282692
Sc. tshawytschae	CBS 100438	On young Oncorhynchus tshawytscha	USA: California	HQ66/562	KF156126	KF156180	KF 155990	KF 282697
Sc. verrucosa	CBS 383.81*	From Soil	India: Kerala	KF156015	KF156129	KF156185	K1272099	1
Sterila eucalypti	CBS 144019*	Eucalyptus sp.	Portugal	MK810918	MK810805	I	MK888742	MK887807
	CPC 14942	Eucalyptus sp.	Porugal	MK810916	MK810803	I	MK888/40	0188/88/W
	CPC 14943	Eucalyptus sp.	Portugal	MK810917	MK810804	1	MK888741	MK88/806
Sympoventuria capensis	CBS 120136*	On leat litter of Eucalyptus sp. (Myrtaceae)	South Africa	MK810921	MK810808	MK926481	MK888745	MK887810
	CPC 12839	On leaf litter of Eucalyptus sp. (Myrtaceae)	South Africa	MK810922	MK810809	MK926482	MK888746	MK887811
	CPC 12840	On leaf litter of <i>Eucalyptus</i> sp. (<i>Myrtaceae</i>)	South Africa	MK810923	MK810810	MK926483	MK888747	MK887812

Species	Strain ¹	Host and substrate	Locality		GenBank ac	GenBank accession numbers ²	bers ²	
				ITS	LSU	tub2	tef1	rpb2
Sy. melaleucae	CBS 143407*	<i>Melaleuca</i> sp., leaves	Australia	MG386059	MG386112	MG386168	1	1
Troposporella fumosa	CBS 351.94	On the old bark of <i>Populus tremula</i> (Salicaceae)	Finland	MK810924	MH874121	I	I	I
Tr. monilipes	MUCL 19867	On decayed wood of Quercus (Fagaceae)	United States	DQ351723	AY 856871	I	I	I
Veronaeopsis simplex	CBS 588.66*	Acacia karroo, leaf litter	South Africa	EU041820	EU041877	I	I	MN091925
Verruconis gallopava	CBS 437.64*	<i>Meleagris gallopavo</i> , brain abscess	United States	HQ667553	KF156112	KF156203	KF155968	KF282689
Ve. mangrovei	NFCCI-4390*	On decaying wood of <i>Excoecaria agallocha</i>	India	MN782361	MN241144	MN848140	I	I
Ve. panacis	CBS 142802*	From the root of Panax notoginseng	China	MF536882	MF536880	MF536883	MF536881	I
Ve. pseudotricladiata	YMF1.04915*	Leaves of a broad-leaf species in a stream	China	MK244396	MK248270	MK253013	MK248273	I
Ve. terricola	CBS 131795*	Isolated from soil	China	MK810925	MK810811	I	I	KC337072
Ve. verruculosa	CBS 119775*	Grassland soil	India	KF156014	KF156106	KF156193	KF155974	I
Yunnanomyces pandanicola	MFLUCC 17-2260*	On decaying leaves of <i>Pandanus amaryllifolius</i>	China	MH388369	MH376743	I	MH388403	MH412736
Yu. phoenicis	MFLUCC 19-0254*	On fallen rachides and leaves of Phoenix paludosa	Thailand	I	MK976738	I	MK986486	MK986484
	MFLUCC 19-0253	On fallen rachides and leaves of Phoenix paludosa	Thailand	ı	MK976737	ı	MK986485	MK986483
Tyrannosorus lichenicola	CBS 144018*	Letharia sp.	USA	MK810953	MK810838	MK926509	MK888775	MK887840
Venturia saliciperda	CBS 480.61*	Salix cordata	Switzerland	MK811007	MK810891	MK926558	MK888825	MK887886
¹ CBS: Westerdijk Fungal Biodiversity Institut Department of Plant Pathology, Agriculture (e, Utrecht, the Netherlands; CPC: Cu College, Guizhou University, China; F	CBS: Westerdijk Fungal Biodiversity Institute, Utrecht, the Netherlands; CPC: Culture collection of Pedro Crous, housed at Westerdijk Fungal Biodiversity Institute; CGMCC: Chinese General Microbiological Culture Collection Center, Beijing, China; GUCC: Culture Collection of the Department of Plant Pathology, Agriculture College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of HGUP: Herbarium of Plant Pathology, Agricultural College, Guizhou University, China; HGUP: Herbarium of HGUP; Herbarium of HGUP: Herbarium of HGUP: Herbarium of HGUP: Herbarium of HGUP: Herbariu	stitute; CGMCC: Chinese General , Guizhou University, China; ICMP	Microbiological Culture International Collecti	re Collection Cent on of Micro-organ	er, Beijing, China isms from Plants	a; GUCC: Culture	Collection of the arch, Private Bag

IRAN: Fungal Culture Collections of the Iranian Research Institute of Plant Protection; MFLU (CC); Mae Fah Luang University Culture Collection, Chiang Ria, Thailand; MUCL: Universite Catholique de Louvain-Ia-Neuve, Belgium, MTCC: actr: actin; tub2: partial B-tubulin gene; tetr: partial translation elongation factor 1-alpha gene; rpb2: partial DNA-directed RNA data numbers of sequences generated in this study are in **bold**; – indicates unavailable sequences or unknown collection SSU: partial small subunit (18S) nrRNA gene; India. Institute of Microbial Technology, Chandigarh, India; NBRC: Biological Resource Center; NFCCI: National Fungal Culture Collection of India, Pune, ITS: internal transcribed spacers and intervening 5.8S nrDNA; LSU: partial large subunit (28S) nrRNA gene; polymerase II second largest subunit gene. Accession Ex-holotype or ex-type strains. 92170, Auckland, New Zealand;

of this fungal group and its association with the environment is necessary, to elucidate the origin of this family and understand the evolutionary patterns of its lifestyles. In this study, seven DNA barcodes (SSU, ITS, LSU, act1, tub2, tef1 and rpb2) were sequenced for 33 strains representing Guizhoumyces (two isolates), Matsushimaea (one isolate), Mycosisymbrium (one isolate), Scolecobasidium (27 isolates) and Verruconis (two isolates). In addition, a multi-locus phylogenetic analysis was performed including 143 taxa of Sympoventuriaceae, and ancestral character states of Sympoventuriaceae were reconstructed. Our specific goals were as follows: i determine the taxonomic position of newly collected strains based on morphological and molecular evidence; ii provide a revised phylogram for Sympoventuriaceae; iii clarify the phylogenetic relationship between Scolecobasidium and Verruconis and other similar genera; and iv to reconstruct the ancestral state and clarify the life strategies during the evolutionary history of Sympoventuriaceae.

MATERIALS AND METHODS

Fungal materials and isolation

The soil, plant and forest litter were collected from China. Each sample or specimen was separately stored in a zip-lock bag or envelope before returning to the laboratory for isolation. Strains were isolated by dilution plate and single spore isolation methods, and subcultured on 2 % potato dextrose agar (PDA) (Crous et al. 2019b). In the present study, 33 strains from 18 species of Scolecobasidium (13 species) and the closely related genera Guizhoumyces (one species), Matsushimaea (one species), Mycosisymbrium (one species) and Verruconis (two species) were collected. The samples include five of the 22 recognised genera in Sympoventuriaceae plus one genus newly described here. The holotype specimens were deposited in the Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University (HGUP). The ex-type cultures are conserved in the Culture Collection of the Department of Plant Pathology, Agriculture College, Guizhou University, China (GUCC) and the China General Microbiological Culture Collection Center (CGMCC).

DNA extraction, amplification and sequencing

Genomic DNA was extracted after 7 d from fresh mycelial cultures grown on PDA. Approximately 50 mg mycelium was scraped off the surface of the medium and transferred to a 1.5 mL microcentrifuge tube. The DNA was extracted using the BIOMIGA Fungus Genomic DNA Extraction Kit GD2416 (Biomiga, USA) following the manufacturer's instructions. The partial nucleotide and protein coding genes were subjected to PCR amplification and sequencing of internal transcribed spacer regions and the intervening 5.8S rRNA gene (ITS) of the rDNA operon, 28S rRNA gene (LSU), 18S ribosomal RNA (SSU), actin gene (act1), translation elongation factor 1-alpha (tef1), RNA polymerase II second largest subunit (*rpb2*) and β -tubulin (*tub2*). For primers and conditions see Table 1. The PCR products were purified and sequenced by Sangon Biotech, and both directions were sequenced to ensure accuracy. The newly generated sequences in this study were deposited in GenBank, the alignments in

and plant pathogens to extremophilic species (thermophilic

fungi) (Samerpitak et al. 2019, Benavent 2021, Murata et al. 2022). Apparently, these fungi have evolved different lifestyles to exploit their environment, suggesting that adaptive radiations within *Sympoventuriaceae* was most likely driven by the ecological diversity (Martin et al. 2016, Haridas et al. 2020). Thus, to know more about the evolutionary importance of this feature, a more detailed study on the co-evolutionary history

Table 2 (cont.)

Table 3 Strains used in	the phylogenetic analysis of <i>Scolec</i>	Table 3 Strains used in the phylogenetic analysis of Scolecobasidium and Verruconis and GenBank accession numbers.	numbers.						
Species	Strain ¹	Host and substrate	Locality		Ger	GenBank accession numbers ²	on numbers ²		
				SSU	ITS	LSU	act1	tub2	tef1
S. ailanthi	MFLUCC 17-0923*	On fallen pod of <i>Ailanthus</i> sp.	Thailand	MK347838	MK347730	MK347947	MK412893	MK412883	1
	MFLU 18-2110	On fallen pod of <i>Ailanthus</i> sp.	Thailand	MK347839	MK347731	MK347948	MK412892	MK412881	I
S. anellii	CBS 284.64*	Stalactite	Italy	KF156070	FR832477	KF156138	KF155912	KF156184	KF1559
S. anomalum	CBS 131816*	Lascaux Cave	France	KF156065	HE575201	KF156137	KF155935	KF156194	KF1559
S. aquaticum	CBS 140316*	Silicone	Germany	KX668260	KX668258	KX668259	I	I	I
S. bacilliforme	CBS 100442*	On stainless steel biofilm in drinking water	Germany	KP798638	KP798632	KP798635	KT272051	KT272059	KT2720
S. blechni	CBS 146055*	Leaves of <i>Blechnum capense</i>	South Africa	I	MN562134	MN567641	I	MN556843	MN556
S. camellicola	GUCC 18242*	On decaying Camellia sinensis leaf litter	China	MZ503654	MZ503728	MZ503761	MZ546837	MZ546907	MZ5468
	GUCC 18243	On decaying Camellia sinensis leaf litter	China	MZ503655	MZ503729	MZ503762	MZ546838	MZ546908	MZ5468
	GUCC 18244	Isolated from forest litter	China	MZ503656	MZ503730	MZ503763	MZ546839	MZ546909	MZ5468
S. capsici	CBS 142096*	Leaf of Capsicum annuum	Thailand	I	KY173427	KY173518	I	I	I
S. coiledmyces	GUCC 18245*	Isolated from lawn soil	China	MZ503657	MZ503731	MZ503764	MZ546840	MZ546910	MZ5468
S. constrictum	CBS 211.53*	Soil	Canada: Ontario	KF156073	HQ667519	KF156148	KF155941	KF156187	KF1560
	CBS 131913	Human, cutaneous mycosis	Thailand	KF156071	KF156025	KF156146	KF155940	KF156176	KF1560
	GUCC 18255	Isolated from dead branches	China	MZ503667	MZ503741	MZ503774	MZ546850	MZ546920	MZ5468
	GUCC 18256	Isolated from forest litter	China	MZ503668	MZ503742	MZ503775	MZ546851	MZ546921	MZ5468

							000000000000000000000000000000000000000	0000112114	
o. allantri	MFLUCC 17-0923" MFL11 18-2110	On fallen pod of <i>Allantitus</i> sp. On fallen pod of <i>Allantitus</i> sp.	I nalland Thailand	MK347830	MK347731	MK347948	MK412893 MK412892	MK412883 MK412881	1 1
S anellii	CBS 284 64*	Stalactite	Italy	KF156070	FR832477	KF156138	KF155912	KF156184	KF155995
	CBS 131816*	Lascaux Cave	France	KF156065	HE575201	KF156137	KF155935	KF156194	KF155986
	CBS 140316*	Silicone	Germany	KX668260	KX668258	KX668259	I	I	I
S. bacilliforme	CBS 100442*	On stainless steel biofilm in drinking water	Germany	KP798638	KP798632	KP798635	KT272051	KT272059	KT272070
S. blechni	CBS 146055*	Leaves of Blechnum capense	South Africa	I	MN562134	MN567641	I	MN556843	MN556826
S. camellicola	GUCC 18242*	On decaying Camellia sinensis leaf litter	China	MZ503654	MZ503728	MZ503761	MZ546837	MZ546907	MZ546874
	GUCC 18243	On decaying Camellia sinensis leaf litter	China	MZ503655	MZ503729	MZ503762	MZ546838	MZ546908	MZ546875
	GUCC 18244	Isolated from forest litter	China	MZ503656	MZ503730	MZ503763	MZ546839	MZ546909	MZ546876
S. capsici	CBS 142096*	Leat of Capsicum annuum	Thailand	1	KY173427	KY173518	1	1	1
S. coiledmyces	GUCC 18245*	Isolated from lawn soil	China 0 · 0 · ·	MZ503657	MZ503731	MZ503764	MZ546840	MZ546910	MZ546877
S. constrictum	CBS 211.53	SOI	Canada: Untario	KF1560/3	HQ66/519	KF156148	KF155941	KF15618/	KF156005
	CBS 131913	Human, cutaneous mycosis	Thailand	KF156071	KF156025	KF156146	KF155940	KF156176	KF156006
	GUCC 18255	Isolated from dead branches	China	MZ503667	MZ503741	MZ503774	MZ546850	MZ546920	MZ546887
	GUCC 18230		Crina	MZ 500000	MZ503/42	MZ503//5	MZ546851	MZ5469/21	00004CZM
	GUCC 1825/		China	MZ503669	MZ503/43	MZ503//6	MZ546852	MZ5469Z2	MZ546889
	GUCC 18258	Isolated from lawn soil	China	MZ503670	MZ503744	MZ503777	MZ546853	MZ546923	MZ546890
o. colualiae		Not available.		KE156056		KF150122 KE156122		KE15019/	KE155000
C oraccitumicala		NUL avaliable Soil	United States	NF 130030		K 1967430	K 195301		
o. crassinarii coa	CBS 141323*	oon Leaf snots of Dracaena reflexa			KX778783	KX228334			KX228377
S. echinulatum			China	MZ503659	M7503733	MZ503766	M7546842	M7546912	M7546879
C. Commanda	GOOG 10241	isotated from soil	China	MZ503660	MZ503734	M7503767	M7546843	M7546913	M7546880
S ellinsoideum	CBS 131706*	Soil	China	-	MN077367	-	-		-
0. 0		leolated from soil	China	MZ503676	M7503750	MZ503783	M7546859	M7546929	MZ546896
	GUCC 10207	isorated inditions soll Isolated from submercied wood	China	M7503677	M7503751	M7503784	MZ546860	M7546930	M7546897
	GUCC 18266	isolated from forest litter	China	M7503678	M7503752	M7503785	M7546861	M7546931	M7546898
S. ferulica	IRAN3232C*	Root of Ferula ovina	Iran		MF186874	MH400207			
S damsii	CBS 239 78*	Carvota nlumosa leaf	Sri Lanka	KF156088	KF156019	KF156150	KF155936	KF156190	KF155982
S. alobale	CBS 119644*	Indoor sample, house	Germany	KF961108	KF961086	KF961097	KF956086	KF961065	KF961075
)	CBS 135924	Bathroom; black biofilm, sink drain	Germany	KF961107	KF961092	KF961104	KF956092	KF961070	KF961079
	GUCC 18249		China	MZ503661	MZ503735	MZ503768	MZ546844	MZ546914	MZ546881
	GUCC 18250	From soil	China	MZ503662	MZ503736	MZ503769	MZ546845	MZ546915	MZ546882
S. guangxiensis	SS23*	Soil and sugarcane root	China	MK929277	MK934570	MK956169	I	I	I
	X22	Soil and sugarcane root	China	MK961269	MK961215	MK961247	I	I	I
S. helicteris	NFCCI 4310*	Living leaves of <i>Helicteris isora</i>	India	I	MK014833	I	I	MK321318	I
S. humicola	CBS 116655*	Peat soil	Canada: Ontario	KF156068	HQ667521	KF156124	KF155904	KF156195	KF155984
S. icarus	CBS 536.69*	Forest soil	Canada: Ontario	KF156084	HQ667524	KF156132	KF155944	KF156174	KF156009
	CBS 423.64	Rhizosphere	Netherlands	KF156085	HQ667523	KF156131	KF155943	KF156173	KF156008
S. lascauxense	CBS 131815*	Black stain on cave sediment	France	KF156069	FR832474	KF156136	KF155911	KF156183	KF155994
S. leishanicola	HGUP 1808*	Soil	China	MK377071	MK377301	MK377073	I	I	1
	GUCC 18259		China	MZ503671	MZ503745	MZ503778	MZ546854	MZ546924	MZ546891
S. longiphorum	CBS 435.76*	In excrement of Insecta, and Quercus	Japan	KF156060	KF156038	KF156135	KF155908	KF156182	KF155978
S. macrozamiae	CBS 137971*	<i>Macrozamia</i> , leaf litter	Australia	I	KJ869123	KJ869180	I	I	1
	CBS 102491	On leaf litter of Macrozamia	Australia	KF156092	KF156021	KF156152	KF155938	KF156191	KF155983
S. minimum	CBS 510.71*	Gossypium arboreum, rhizosphere	Nigeria	KF156087	HQ667522	KF156134	KF155945	KF156172	KF156007
		Soll From format humin	China		NF 150UZ/	M7503770	NF 100940	NF 1501/5	NI 2/ 20/ 3
S mirabilic	GUCU 18200 CBS 413 51*	From forest numus Breathing requilator for diver	Cnina Netherlende	KE156076	M2503/46	KE156140	KE166055	KE156164	VE166001
		Decidator of diver	Netheriarius	KE166080	KE166020	KE156144	KE155019	KE156171	KE155000
o. musicola	CBS / 29.95" CBS 144441*	Regulator of alver On leaves of Musa sn (Musareae)	Netherlands Malavsia		MH327824	MH327860	NF 100440	MH327898	MH327887
0. 1110510018		OII ICAVES OI INUSA SP. (INUSACEAE)	lviaiay sia	I			I		

Species	Strain ¹	Host and substrate	Locality		Ger	GenBank accession numbers ²	in numbers ²		
				SSU	ITS	LSU	act1	tub2	tef1
S. obovoideum	GUCC 18246*	Isolated from forest litter	China	MZ503658	MZ503732	MZ503765	MZ546841	MZ546911	MZ546878
	CBS 137170*	Man, bronchoalveolar lavage fluid	USA: Utah	LM644548	LM644521	LM644564	LM644600	LM644605	KT272067
S. pandanicola	CBS 140660*	Pandanus utilis, leaves	France	I	KT950850	KT950864	I	I	I
S. phaeophorum	CBS 206.96*	Leaf in coastal rain forest	Papua New Guinea	KP798637	KP798631	KP798634	KT272054	KT272062	KT272098
S. podocarpi	CBS 143174*	Podocarpus grayae, leaves	Australia	I	MG386032	MG386085	I	I	MG386162
S. podocarpicola	CBS 146057*	Leaves of Podocarpus latifolius	South Africa	I	MN562138	MN567645	I	I	I
S. ramosum	CBS 137173*	Isolated from nail of Homo sapiens	USA: California	LM644551	LM644524	LM644567	LM644603	LM644608	KT272069
	CBS 137171	Skin	United States	LM644549	LM644522	LM644565	LM644601	LM644606	KT272068
	GUCC 18261	From forest litter	China	MZ503673	MZ503747	MZ503780	MZ546856	MZ546926	MZ546893
	GUCC 18262	From soil	China	MZ503674	MZ503748	MZ503781	MZ546857	MZ546927	MZ546894
	GUCC 18263	From forest litter	China	MZ503675	MZ503749	MZ503782	MZ546858	MZ546928	MZ546895
S. robustum	CBS 112.97*	Leaf litter of Ouercus ilex	Spain	KP798639	KP798633	KP798636	KT272052	KT272060	KT272071
S. sexuale	CBS 135765*	Swabs in a laboratory	South Africa	KF156089	KF156018	KF156118	KF155902	KF156189	KF155976
	CBS 131965	Ant	Brazil	KF156090	KF156017	KF156119	KF155903	KF156188	KF155977
S. terreum	CBS 203.27*	From soil	USA: Louisiana	I	HQ667544	I	I	HQ877665	I
S. tshawytschae	CBS 100438*	On young Oncorhynchus tshawytscha	USA: California	KF156062	HQ667562	KF156126	KF155918	KF156180	KF155990
	CBS 228.66	Peat-bog soil	Ireland	KF156064	KF156016	KF156128	KF155915	KF156179	KF155992
	GUCC 18251	From lawn soil	China	MZ503663	MZ503737	MZ503770	MZ546846	MZ546916	MZ546883
	GUCC 18252	From plant litter	China	MZ503664	MZ503738	MZ503771	MZ546847	MZ546917	MZ546884
	GUCC 18253	From soil	China	MZ503665	MZ503739	MZ503772	MZ546848	MZ546918	MZ546885
	GUCC 18254	From soil	China	MZ503666	MZ503740	MZ503773	MZ546849	MZ546919	MZ546886
S. variabile	NBRC 32268	From soil	Canada: Ontario	EU107353	DQ307334	EU107310	I	I	DQ307356
S. verrucaria	GUCC 18240*	From soil	China	MZ503652	MZ503726	MZ503759	MZ546835	MZ546905	MZ546872
S. verrucosum	CBS 383.81*	From soil	India: Kerala	KF156067	KF156015	KF156129	KF155910	KF156185	KT272099
S. zunyiense	GUCC 18241*	From forest litter	China	MZ503653	MZ503727	MZ503760	MZ546836	MZ546906	MZ546873
V. calidifluminalis	CBS 125818*	Water of a hot stream	Japan	KF156046	AB385698	KF156108	KF155901	KF156202	KF155959
	CBS 125817	Water of a hot stream	Japan	KF156045	AB385699	KF156107	KF155900	KF156201	KF155958
V. cylindricalis	GUCC 18299*	From forest humus	China	MZ503680	MZ503754	MZ503787	MZ546863	MZ546933	MZ546900
V. gallopava	CBS 437.64*	<i>Meleagris gallopav</i> o, brain abscess	United States	KF156053	HQ667553	KF156112	HQ916989	KF156203	KF155968
	CBS 118.91	Man	United States	KF156047	HQ667551	KF282655	KF155932	HQ877643	JF440539
	CBS 867.95	Sputum from patient with cardiac	United States	KF156051	HQ667561	KF282657	KF155928	KF156213	KF155972
	CBS 116660	Human, transplantation	United States	KF156048	HQ667557	KF156115	KF155929	KF156206	KF155969
V. hainanensis	YMF1.04165*	From leaves of a dicotyledonous plant	China	MK248267	MK244397	MK248269	MK248271	I	MK248272
V. heveae	MFLUCC 17-0092*	On dried latex on bark of Hevea brasiliensis	Thailand	I	MH602349	MH602348	I	I	I
V. mangrovei	NFCCI-4390*	On decaying wood of <i>Excoecaria agallocha</i>	India	MN241147	MN782361	MN241144	I	MN848140	I
	NFCCI-4391	On decaying wood of <i>Excoecaria agallocha</i>	India	MN241148	MN782362	MN241145	I	MN848141	I
V. panacis	CBS 142802*	From the root of Panax notoginseng	China	MF536879	MF536882	MF536880	I	MF536883	MF536881
V. pseudotricladiata	YMF1.04915*	Leaves of a broad-leaf species in a stream	China	MK248268	MK244396	MK248270	I	MK253013	MK248273
V. terricola	CBS 131795*	Isolated from soil	China	I	MK810925	MK810811	I	I	I
V. thailandica	CBS 145768*		Thailand	I	MN794375	MN794352	I	I	I
	GUCC 18267	Isolated from the humus soil in the stream	China	MZ503679	MZ503753	MZ503786	MZ546862	MZ546932	MZ546899
V. tricladiata	NBRC 30208	On rotten leaves	Bismarck Archipelago	EU107354	I	EU107286	I	I	DQ307352
V. verruculosa	CBS 119775*	Grassland soil	India	KF156055	KF156014	KF156106	KF155919	KF156193	KF155974
Pseudosigmoidea excentrica	CBS 469.95*	<i>Lauraceae</i> , leaf litter	Cuba	KF156096	HQ667543	KF282669	KF155934	MK926478	KF155975
Sympoventuria capensis	CBS 120136*	Eucalyptus sp., leaf litter	South Africa	KF156094	MK810921	MK810808	I	MK926481	MK888745
¹ CBS: Westerdiik Fungal Biodiversity	¹ CBS: Westerdiik Fundal Biodiversity Institute. Utrecht. the Netherlands: CPC: Culture collection of Pedro Crous.		housed at Westerdlik Fundal Biodiversity Institute: CGMCC: Chinese General Microbiological Culture Collection of the	hinese General Mic	robiological Cultu	re Collection Cen	ter. Beijina. Chini	a: GUCC: Culture	Collection of the
Department of Plant Pathology, Agric	ulture College, Guizhou University, Chin		ricultural College, Guizhou Universit	ty, China; ICMP: Inte	ernational Collecti	on of Micro-orgar	isms from Plants	s, Landcare Rese	irch, Private Bag
92170 Auckland New Zealand IRA	V. Fundal Culture Collections of the Iran	aian Besearch Institute of Plant Protection: MELLI (CC): Mae E	ah Luand Thiversity Culture Collect	ion Chiann Bia Th	ailand: MLICL - LIr	inversite Catholin	in de Louvain L	ouvain-la-Neuve	Selnium MTCC

92170. Auckland, New Zealand; IFAN: Fungal Culture Collections of the Iranian Research Institute of Plant Protection; MFLU (CC): Mae Fah Luang University Culture Collection, Chiang Ria, Thailand; MUCL: Universite Catholique de Louvain, Louvain-la-Neuve, Belgium. MTCC: Institute of Microbial Technology, Chandigam, India; NBRC: Biological Resource Center, NFCCI: National Fungal Culture Collection of India, Pune, India. ² ITS: internal transcribed spacers and intervening 5.8S nrDNA; LSU: partial large subunit (28S) nrRNA gene; SSU: partial subunit (18S) nrRNA gene; *act1*: actin, *tub2*: partial β-tubulin gene; *tef1*: partial translation elongation factor 1-alpha gene; *rpb2*: partial DNA-directed RNA polymerase II second largest subunit gene. Accession numbers of sequences generated in this study are in **bold**; – indicates unavailable sequences or unknown collection data.

Table 3 (cont.)

TreeBASE (Submission ID S29226), and all the sequences used for phylogenetic analysis are shown in Table 2 and 3.

Phylogenetic analyses

The concatenated DNA sequence dataset (SSU, ITS, LSU, act1, tub2, tef1 and rpb2) of 143 taxa was used to infer phylogenetic relationships among the new isolates and other taxa of Sympoventuriaceae. DNA sequence data were initially blast searched to determine the placement of new strains in Sympoventuriaceae. Multiple sequence alignments were carried out with MAFFT v. 7.4.9 (Rozewicki et al. 2019), and then rechecked and adjusted manually as necessary using BioEdit v. 7.1.9 (Hall 1999). The single gene datasets were combined using MEGA X (Kumar et al. 2018). Data were converted from fasta to nexus and phylip format with AliView v. 1.19 for RAxML, MrBayes and PAUP analysis (Larsson 2014). Finally, phylogenetic analyses for the individual data matrix and combined datasets were conducted by employing maximum likelihood (ML), maximum parsimony (MP) and Bayesian inference (BI). The ML analyses used RAxML-HPC2 on XSEDE v. 8.2.12 (Stamatakis 2014) via the CIPRES Science Gateway platform (Miller et al. 2012). The GTRGAMMA model was chosen and ML bootstrap analyses were estimated with 1000 replicates. Prior to Bayesian analysis, jModelTest v. 2.1.7 (Darriba et al. 2012) was used to select a best-fit model of nucleotide substitution for each data partition under the output strategy of Akaike information criterion (AIC) (Nylander 2004). The Bayesian posterior probabilities were determined by Markov Chain Monte Carlo sampling (MCMC) in MrBayes v. 3.2.7 (Ronquist et al. 2012). The six simultaneous Markov chains were run for 2 M generations, starting from random trees and sampling trees every 100th generation, and 25 % of ageing samples were discarded, running until the average standard deviation of the split frequencies dropped below 0.01. For MP, the dataset was analysed in PAUP v. 4.0b10 (Swofford 2003) using a heuristic search algorithm with 1000 random addition sequence replicates. One tree was saved at each step during stepwise addition, while tree bisection reconnection (TBR) was used to swap branches, and the maximum number of trees was set to 10000. The ambiguously aligned regions were eliminated and gaps were treated as missing data. The phylogenetic trees were visualised in FigTree v. 1.4.4 and edited using Adobe Illustrator CC 2020.

Estimating transitions in lifestyle evolution

Ancestral character states of the lifestyle (Table 2, 3) were reconstructed with the Bayesian Binary Method (BBM) of RASP v. 4.2 (Yu et al. 2015, 2020). Because BBM analysis requires a set of phylogenetic trees and a consensus topology, we generated the phylogenetic trees via Bayesian phylogenetic analysis in BEAST v. 2.6.6 (Barido-Sottani et al. 2018) using six DNA loci (SSU, ITS, LSU, act1, tub2 and tef1). The length of the MCMC chain reaction was set as 500 M generations sampled every 100000 generations; thus, a total of 5000 trees were kept. Tracer v. 1.7.2 (Rambaut et al. 2018) was used to check that the values of the mean and ESS in the log file were over 200. After removal of a proportion of each run as burn-in, the remaining trees were summarised as maximum clade credibility (MCC) trees in TreeAnnotator v. 2.6.6 (Barido-Sottani et al. 2018). For BBM analysis, the Markov chains were run for 50 000 generations, using 10 chains, with a sample frequency of 100, a temperature of 0.1, state frequencies fixed (JC), and among-site rate variation equal. We subsequently examined the resulting reconstructions of the selected characters to determine if the lifestyle changes were arising convergently or resulted from shared ancestry.

Morphological observations

The microscopic features and colony characteristics of the putative novel and known species were examined. The macromorphological characters and relevant data (colony colour and diameter, mycelium) of the isolates were examined under a dissecting microscope (Leica S9i, Germany), and images of colonies cultured on three media, malt extract agar (MEA), oatmeal agar (OA) (Crous et al. 2019b) and PDA were captured after 2 wk. The slide culture technique on OA was used for microscopic observation. If sterile on OA, morphological characters produced on other media were described. Measurements and descriptions of reproductive structures were taken from specimens mounted in lactic acid or lactophenol cotton blue. Micrographs were captured with an Olympus BX53 compound microscope. Tarosoft (R) Image Frame Work program was used to measure the lengths and widths of microscopic structures including conidiophores, conidiogenous cells, conidia and chlamydospores per isolate. At least 30 measurements were made for each microscopic structure to calculate the mean value, standard deviation, and minimum-maximum values, with the extreme measurements in parentheses (Giraldo & Crous 2019, Fan et al. 2020, Liu et al. 2022). Descriptions are based on observations after 14 d of incubation at 26 °C. Slowgrowing species were allowed to grow longer, for 20-30 d, until sporulation was observed.

RESULTS

Phylogenetic analyses

For the two datasets in the present study (Table 2, 3), phylogenetic analyses obtained from ML, MP and BI analyses resulted in trees with similar topologies, and the best scoring ML tree was selected to represent and discuss the phylogenetic relationships among taxa (Fig. 1, 2). Sympoventuriaceae phylogeny (Fig. 1): the first tree was based on a concatenated DNA sequence dataset (ITS, LSU, tef1, tub2 and rpb2) used to infer the phylogenetic position of the treated genera and species within the Sympoventuriaceae. The sequence data comprised 69 taxa for Sympoventuriaceae with Tyrannosorus lichenicola and Venturia saliciperda as the outgroup taxa. The dataset consisted of 5175 characters, of which 2399 were constant, 2173 parsimony-informative and 603 parsimony-uninformative. Based on the results of the jModelTest, GTR+I+G was estimated as the optimal nucleotide substitution model under the output strategy of AIC; Scolecobasidium and Verruconis phylogeny (Fig. 2): for the sake of revealing the phylogenetic relationship between Scolecobasidium and Verruconis species, a second analysis was performed on the six gene regions (ITS, LSU, SSU, act1, tub2 and tef1) of 97 taxa within the genus, and Pseudosigmoidea excentrica and Sympoventuria capensis were used as outgroups. The final aligned sequence matrix contained 5919 characters, of which 3054 were constant, 2152 parsimony-informative and 713 parsimony-uninformative. The optimal nucleotide substitution model HKY+I+G was used for the phylogenetic analyses.

The phylogenetic tree of *Sympoventuriaceae* distinguished 22 subclades, each subclade representing a highly supported monophyletic group (Fig. 1). *Acroconidiellina* was located at the terminal end of the phylogenetic tree, closely related to *Scolecobasidium* and *Verruconis* (Fig. 1). Phylogenetic analyses resolved 43 species of *Scolecobasidium*, which chiefly clustered in five subclades, of which 37 correspond to known species of the genus (Fig. 2). In the *Verruconis* lineage three major subclades were observed, corresponding closely to the currently recognised 11 species (Fig. 2). The 29 newly collected strains clustered in eight well-supported subclades of

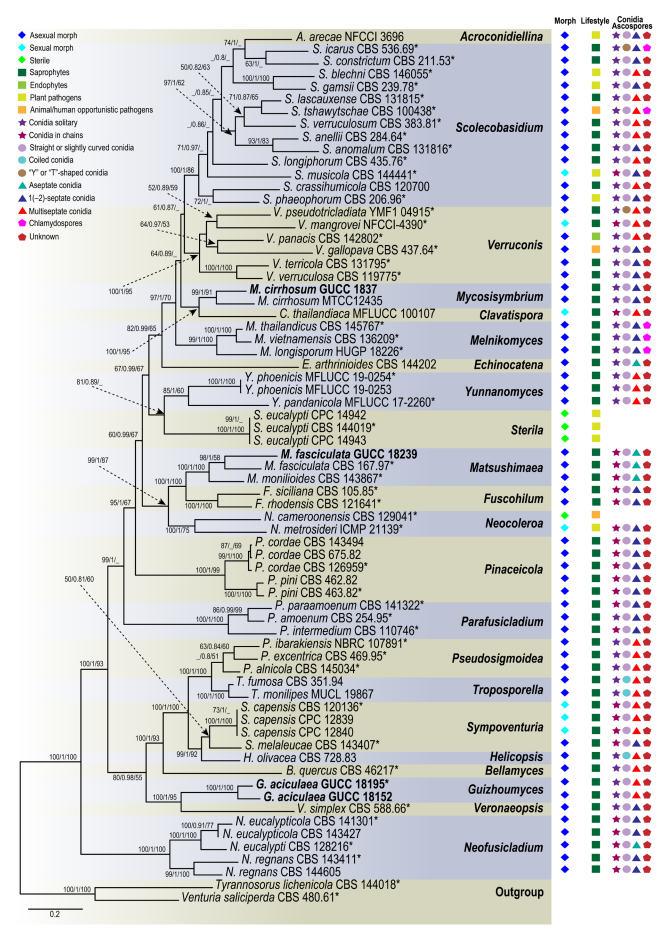


Fig. 1 Phylogenetic tree of the family *Sympoventuriaceae* based on RAxML analyses of combined DNA dataset of ITS, LSU, *tef1*, *tub2* and *rpb2* gene sequences. Bootstrap values \geq 50 % for Maximum parsimony and Maximum likelihood, and Bayesian posterior probabilities \geq 80 % are presented at the branches (ML/BI/MP). Some branches were shortened to facilitate layout, and the scale bar represents the number of changes. *Tyrannosorus lichenicola* and *Venturia saliciperda* are used as outgroup. Those in **bold** are new taxa or new combinations proposed in the current study and the strains obtained, as well as type strains are marked with an asterisk (*). Lifestyles and typical morphological characteristics of individual strains are shown at the right side of the phylogenetic tree, and the related icons plotted are explained in the legend in the upper left corner.

Scolecobasidium and Verruconis respectively, including seven new species, as well as six new combinations proposed here (Fig. 2). The monotypic genus *Mycosisymbrium* consisted of two highly supported clades, which can be distinguished from other genera in this family, and *Clavatispora* as the sister genus (Fig. 1). Three clades were distinguished in *Melnikomyces*, corresponding closely to the currently recognized three species (*M. longisporus*, *M. thailandicus* and *M. vietnamensis*), which are saprophytes in soil or plant debris (Fig. 1). Echinocatena, a monotypic genus represented by *E. arthrinioides*, formed a robust lineage at the base of the *Melnikomyces* (Fig. 1). Sterila and *Yunnanomyces* clustered in the same subclade and received moderate to strong support, which was divided into two clades; the first one included the ex-type strain of *Y. pandanicola* and two *Y. phoenicis* strains, and the second one included three sterile strains (*S. eucalypti*) (Fig. 1).

Fuscohilum, Matsushimaea and Neocoleroa grouped together and represent three monophyletic groups; the first group comprising *F. rhodense* and *F. sicilianum*, the second group *M. fasciculata* and *M. mtonilioides*, and the third group *N. cameroonensis* and *N. metrosideri*, which are plant, animal and human pathogens (Fig. 1). *Pinaceicola* (*P. cordae* and *P. pini*) and *Parafusicladium* (*P. amoenum*, *P. intermedium* and *P. paraamoenum*) formed a well circumscribed clade (Fig. 1). *Pseudosigmoidea* included *P. alnicola*, *P. excentrica* and *P. ibarakiensis*, which formed a robust clade with four other genera, viz.,

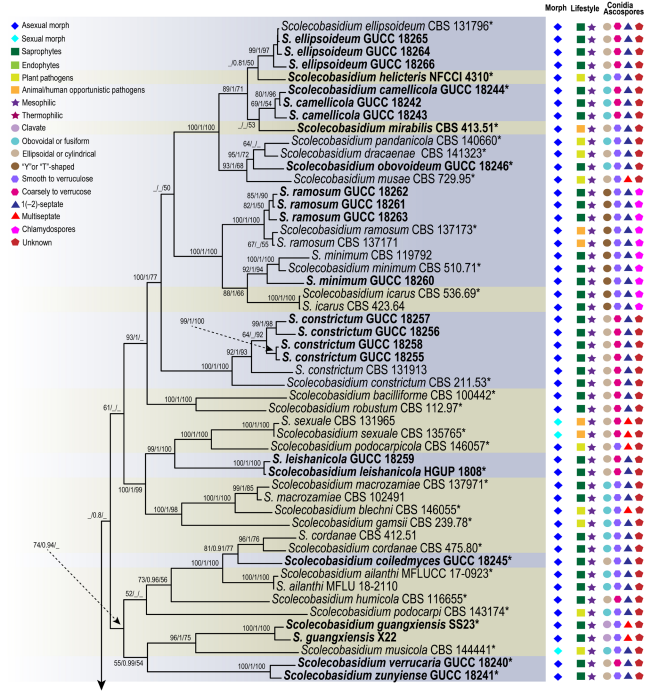


Fig. 2 Phylogenetic tree of the genera *Scolecobasidium* and *Verruconis* based on RAxML analyses of combined DNA dataset of SSU, ITS, LSU, *act1*, *tub2* and *tef1* gene sequences. Bootstrap values \geq 50 % for Maximum parsimony and Maximum likelihood, and Bayesian posterior probabilities \geq 80 % are presented at the branches (ML/BI/MP). Some branches were shortened to facilitate layout, and the scale bar represents the number of changes. *Pseudosigmoidea excentrica* and *Sympoventuria capensis* are used as outgroup. Those in **bold** are new taxa or new combinations proposed in the current study and the strains obtained, as well as type strains are marked with an asterisk (*). Lifestyles and typical morphological characteristics of individual strains are shown at the right side of the phylogenetic tree, and the related icons plotted are explained in the legend in the upper left corner.

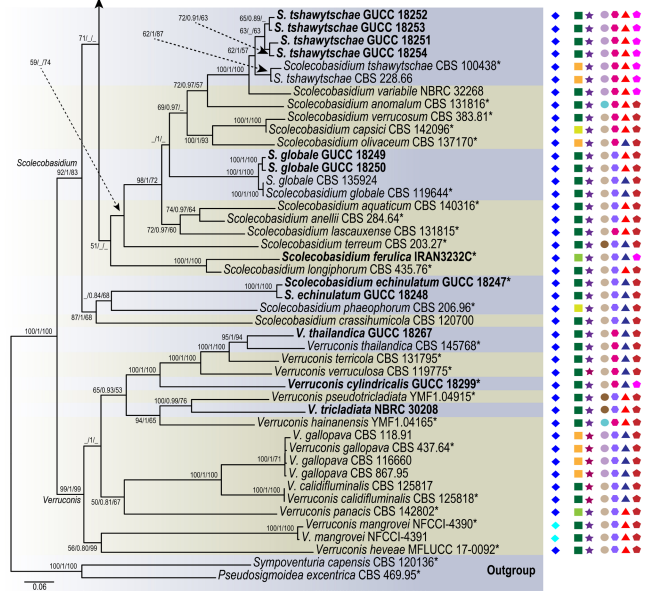


Fig. 2 (cont.)

Bellamyces, Helicopsis, Sympoventuria and Troposporella (Fig. 1). However, the phylogenetic relationship of these genera with the other clades remains poorly unresolved. *Guizhoumyces* belongs to a monophyletic clade, together with Veronaeopsis, representing a new genus in Sympoventuriaceae (Fig. 1). Neofusicladium encompassed three highly statistically supported subclades at the base of Sympoventuriaceae, which represent *N. eucalypti*, *N. eucalypticola* and *N. regnans*, respectively (Fig. 1). Relationships of the new taxa are discussed in the notes.

Lifestyle evolution analysis

Ancestral states of the lifestyles in *Sympoventuriaceae* were inferred on the reconstructed phylogeny. We defined the life strategies in four states: saprophytes, endophytes, plant pathogens, and animal/human opportunistic pathogens. Overall, the results of BBM analysis revealed that the life strategies of *Sympoventuriaceae* was based on saprophytes as the primitive state, and endophytes, plant pathogens, animal/human opportunistic pathogens were derived states, correlating with phylogeny (Fig. 3). At the genus level, *Bellamyces, Echinocatena, Guizhoumyces, Helicopsis, Neofusicladium, Parafusicladium, Pseudosigmoidea, Sympoventuria, Troposporella* and Vero-

naeopsis were basal in Sympoventuriaceae with a saprotrophic lifestyle, correlating with their ecology. The recently introduced Sterila and Neocoleroa were strongly supported as ingroup taxa of Sympoyenturiaceae, and have evolved from saprophytes to become plant pathogens (for S. eucalypti and N. metrosideri) or animal/human opportunistic pathogens (for N. cameroonensis) (Fig. 3). However, these lifestyles reverted to saprotrophic again in the well-supported Clavatispora, Fuscohilum, Matsushimaea, Melnikomyces and Mycosisymbrium clades. In the Acroconidiellina, Scolecobasidium and Verruconis clades, the number of plant pathogens and animal or human opportunistic pathogens increased from Sterila (one species) to Acroconidiellina (one species), Scolecobasidium (16 species) and Verruconis (one species), whereas the transition to endophytes occurred only twice in Scolecobasidium and Verruconis, respectively (Fig. 3). Thus, we conclude that the outstanding diversification of Scolecobasidium is related to the evolution of derived life strategies, and that the lifestyle of this genus may have been influenced by both plants and animals. Moreover, the common ancestor of the other genera except for Acroconidiellina, Neocoleroa and Sterila was saprophytic (Fig. 3), which is a derived condition from a saprotrophic ancestor of Sympoventuriaceae.

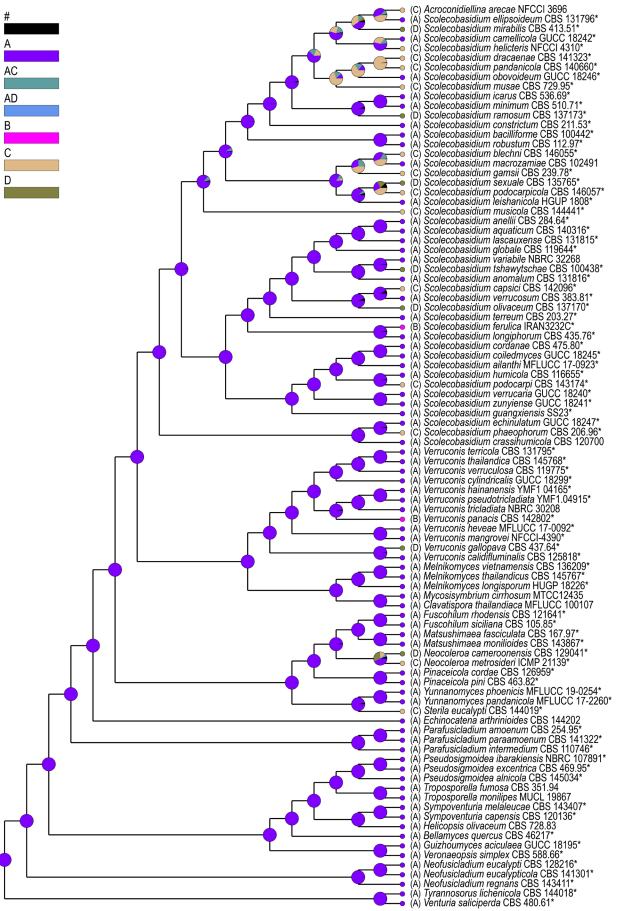


Fig. 3 Ancestral character state analysis focusing on lifestyles in Sympoventuriaceae, using Bayesian Binary MCMC as alternative method. The pie chart at each node indicates the relative probabilities of all possible ancestral states from the Bayesian analysis, and black (#) shows the pooled probabilities of estimates that each account for < 5 %. Coloured circles at the tip and letters next to taxa represent their current lifestyle. A. saprophytes; B. endophytes; C. plant pathogens; D. animal/human opportunistic pathogens. The type strains are marked with an asterisk (*). Tyrannosorus lichenicola and Venturia saliciperda are used as outgroup.

Table 4	Genera	accepted	in S	ympo\	enturiaceae.
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Zhang et al. (2011)	Wijayawardene et al. (2014)	Tibpromma et al. (2018)	Wijayawardene et al. (2020)	Shen et al. (2020)	This study (2022)
Fusicladium-like	Clavatispora	Fusicladium	Acroconidiellina	Bellamyces	Acroconidiellina
Sympoventuria	Ochroconis	Ochroconis	Clavatispora	Echinocatena	Bellamyces
Veronaeopsis	Sympoventuria	Sympoventuria	Fusicladium	Fuscohilum	Clavatispora
	Veronaeopsis	Scolecobasidium	Matsushimaea	Helicopsis	Echinocatena
		Veronaeopsis	Mycosisymbrium	Neocoleroa	Fuscohilum
		Verruconis	Ochroconis	Neofusicladium	Guizhoumyces
		Yunnanomyces	Sympoventuria	Pseudosigmoidea	Helicopsis
			Veronaeopsis	Parafusicladium	Matsushimaea
			Verruconis	Pinaceicola	Melnikomyces
			Yunnanomyces	Sympoventuria	Mycosisymbrium
				Scolecobasidium	Neocoleroa
				Sterila	Neofusicladium
				Troposporella	Pseudosigmoidea
				Veronaeopsis	Parafusicladium
				Verruconis	Pinaceicola
					Sympoventuria
					Scolecobasidium
					Sterila
					Troposporella
					Veronaeopsis
					Verruconis
					Yunnanomyces

TAXONOMY

Sympoventuriaceae Y. Zhang ter et al., Fungal Diversity 51: 255. 2011

Type genus. Sympoventuria Crous & Seifert.

Notes — Sympoventuriaceae was established by Zhang et al. (2011) with Sympoventuria designated as the type genus, which can be distinguished from the Venturiales by its saprophytic lifestyle, presence of pseudoparaphyses, and hyaline, symmetrical ascospores (Arzanlou et al. 2007, Crous et al. 2007a, b, Zhang et al. 2011). Phylogenetically, Sympoventuriaceae forms a well-supported family clade within Venturiales (Zhang et al. 2011, Machouart et al. 2014). Subsequently, Hyde et al. (2013) recognised three lineages (i.e., Sympoventuria, Veronaeopsis and fusicladium-like species) within Sympoventuriaceae and provided more details. Over the past few years, more genera have been accepted in Sympoventuriaceae, such as Ochroconis, Scolecobasidium and Verruconis (Machouart et al. 2014). The taxonomy of Sympoventuriaceae has since been widely studied and dramatically changed (Wijayawardene et al. 2014, Tibpromma et al. 2018). Despite these changes, the phylogenetic placement of many genera in the Sympoventuriaceae remains to be elucidated. Shen et al. (2020) re-described Sympoventuriaceae based on a multigene phylogenetic analysis, morphological and ecological comparisons, and included 15 genera in this family. However, generic boundaries within Sympoventuriaceae are poorly resolved and still controversial, due to lack of type materials and unresolved phylogenies. In this study, we accept 22 genera in Sympoventuriaceae (Fig. 1, Table 4).

Guizhoumyces T.P. Wei & Y.L. Jiang, gen. nov. — MycoBank MB 840922

Etymology. Named after Guizhou, where this species was collected and the Greek name for fungi (myces).

Type species. Guizhoumyces hyalinaea T.P. Wei & Y.L. Jiang.

Mycelium consisting of curved or straight, branched, pale brown, septate, smooth-walled hyphae, frequently forming hyphal coils. *Conidiophores* subcylindrical, simple, branched, straight

or slightly geniculate, pale brown, smooth, septate, sometimes reduced to conidiogenous cells. *Conidiogenous cells* integrated, simple, polyphialidic, sympodially proliferating, elongate lageniform or ampulliform, terminal or intercalary, pale brown, with an inconspicuous or distinct denticle at the conidiogenous locus after rhexolytic conidial secession. *Conidia* enteroblastic, solitary, acicular to obclavate or cylindrical, septate, straight or somewhat curved, smooth, thin-walled, subhyaline to pale brown, apex subobtuse to pointed, base truncate to short obconically truncate, with thickened and darkened hilum; anastomosis between mature conidia. *Chlamydospores* were not observed and *sexual morph* unknown.

Notes — The genus *Guizhoumyces* was established to accommodate a new species *G. aciculaea*. The phylogeny of concatenated ITS, LSU, *rpb2*, *tef1* and *tub2* DNA sequences indicated that *Guizhoumyces* formed a fully supported monophyletic lineage in *Sympoventuriaceae*, which is sister to *Bellamyces*, *Helicopsis*, *Sympoventuria* and *Veronaeopsis*. Morphologically, this genus has certain similarities with *Pseudosigmoidea* and *Sigmoidea*, but can be distinguished from them by its acicular to obclavate or cylindrical, less than four septate and smaller conidia (Crane 1968, Ando & Nakamura 2000, Crous et al. 2019a). Therefore, based on its unique morphological characteristics and phylogenetic location, a new genus name *Guizhoumyces* is introduced to accommodate this new fungus.

Guizhoumyces aciculaea T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 840923; Fig. 4

Etymology. The epithet refers to the acicular conidia.

Typus. CHINA, Guizhou Province, Shiqian County, Pingshan Township, Fodingshan National Nature Reserve, N27°40'50" E108°07'30", 1100 m a.s.l., isolated from soil, 2 Nov. 2019, *T.P. Wei* (holotype HGUP 18195, isotype CGMCC 3.20543, culture ex-type GUCC 18195).

Mycelium consisted of branched, pale brown, septate, thickwalled, 2–3 µm diam hyphae, frequently forming hyphal coils. *Conidiophores* mostly flask-shaped to subcylindrical, simple, straight or slightly geniculate, pale brown, smooth, septate, sometimes reduced to conidiogenous cells, (10-)13-46.5(-48)× 1.5–2.5(-3) µm (av. ± SD = 25.5 ± 12.7 × 2.1 ± 0.4 µm,



Fig. 4 *Guizhoumyces aciculaea* (culture ex-type GUCC 18195). a–c. Colony on PDA, OA and MEA; d. hyphal coils and conidia; e–j. conidiophores reduced to conidiogenous cells; k–l. conidiophores with conidiogenous cells and conidia; m. anastomosis between mature conidia. — Scale bars: d–m = 10 µm.

n = 30). Conidiogenous cells integrated, simple, polyphialidic, sympodially proliferating, elongate lageniform or ampulliform, terminal or intercalary, pale brown, $4-12(-15.5) \times 2-3.5 \mu m$ (av. \pm SD = 7.8 \pm 2.7 \times 2.7 \pm 0.4 μm , n = 30), with one or numerous denticles in the apex, hyaline to pale brown, $1-3 \mu m$ long. Conidia separate rhexolytically from conidiogenous cells, enteroblastic, solitary, acicular to obclavate or cylindrical, 0(-3)-septate, straight or somewhat curved, smooth, thinwalled, subhyaline to pale brown, apex subobtuse to pointed, base truncate to short obconically truncate, with thickened and darkened hilum, anastomosis between mature conidia, $(19.5-)21.5-37(-39.5) \times 1.5-2 \mu m$ (av. \pm SD = 27.8 \pm 4.2 \times 1.5 \pm 0.1 μm , n = 30).

Culture characteristics — Colonies on PDA reaching up to 14–15 mm diam after 14 d at 26 °C, compact, surface grey brown, slightly raised at centre. On OA reaching 16–19 mm diam, with sparse aerial mycelium, olivaceous grey. On MEA reaching 14–16 mm diam, raised, hairy, with abundant aerial hyphae, grey at the surface, reverse pale brown.

Additional material examined. CHINA, Guizhou Province, Shiqian County, Ganxi Township, Fodingshan National Nature Reserve, N27°40'51" E108°07'21", 1240 m a.s.l., from leaf litter, 10 June 2019, *T.P. Wei* (HGUP 18152), living culture GUCC 18152 = CGMCC 3.20542.

Notes — Guizhoumyces aciculaea somewhat resembles the type species P. cranei and S. prolifera of Pseudosigmoidea and Sigmoidea in conidial morphology, with enteroblastic conidiogenesis and phialidic conidiogenous cells, which would suggest that our taxon could be accommodated here. Unfortunately, Pseudosigmoidea and Sigmoidea (Halosphaeriaceae, Microascales) are distantly related to G. aciculaea (Fig. 1). Morphologically, G. aciculaea can also be distinguished from P. cranei and S. prolifera. The conidia of P. cranei are scolecoid, 3(-8)-septate and longer (29–116.5 × 1.5–2.5 µm) (Ando & Nakamura 2000); conidia of S. prolifera are scolecoid, 5(-11)-septate, hyaline and larger (44-110 \times 2-2.5 μ m) (Crane 1968). In contrast, G. aciculaea has acicular to obclavate or cylindrical, 0(-3)-septate, subhyaline to pale brown and smaller conidia (19.5–39.5 \times 1.5–2 µm), and anastomosis occurs among mature conidia. Moreover, the ex-type culture of G. hyalinaea and V. simplex clustered in two distinct clades representing two different genera (Fig. 1). Guizhoumyces hyalinaea is clearly distinct from V. simplex, which has oblong to subcylindrical, 0(-1)-septate and very small conidia (6-15 × 2-4 µm) (Arzanlou et al. 2007).

Scolecobasidium E.V. Abbott, Mycologia 19: 30. 1927

Synonym. Ochroconis de Hoog & Arx, Kavaka 1: 57. 1974 '1973'.

Type species. Scolecobasidium terreum E.V. Abbott.

Notes — Scolecobasidium was first described by Abbott (1927) to accommodate S. constrictum and S. terreum isolated from cotton and sugarcane soils in Louisiana, USA, with S. terreum designated as the generic type, which has Y-shaped and yellowish conidia. The salient characters of Scolecobasidium are rust-brown to olivaceous colonies producing small, brownish conidiophores bearing small numbers of dark, septate, roughwalled, rhexolytic conidia (Abbott 1927, Ellis 1976). Abbott (1927) pointed out that Scolecobasidium is distinguished from other groups by the shape of its conidia and the way conidia are arranged on its conidiophores. In the following decades, this genus received unanimous support (Barron & Busch 1962, Roy et al. 1962, Graniti 1963). Later, more species with unbranched conidia were described within Scolecobasidium, which led De Hoog & Von Arx (1973) to introduce a separate genus, Ochroconis, typified by O. constricta for hyphomycetous species with unbranched, subspherical to cylindrical or clavate, melanised conidia (Matsushima 1975, 1980, Punithalingam &

Spooner 2011). *Scolecobasidium* was restricted to species with T- or Y-shaped or bilobed, two- to multi-celled conidia (Martin-Sanchez et al. 2012). It is noteworthy that the ex-type strains of both *S. terreum* (CBS 203.27) and *O. constricta* (CBS 202.27) are now sterile (Horre et al. 1999, Gams 2015).

Samerpitak et al. (2014) revised Ochroconis and Scolecobasidium using SSU, ITS, LSU, act1, tub2 and tef1 DNA sequences. They found that Ochroconis and Scolecobasidium clustered together, while Scolecobasidium was considered as doubtful because the ex-type culture was sterile (Samerpitak et al. 2017). This opinion, however, was not shared by Gams (2015) who regarded Ochroconis as a synonym of Scolecobasidium, which was supported by Seifert et al. (2011). More recently, Shen et al. (2020) resolved Ochroconis as a synonym of Scolecobasidium based on the multi-locus (ITS, LSU, tef1, tub2 and rpb2) analysis combined with morphology and ecology, with strong support for its monophyly. We agree with Shen et al. (2020) that Solecobasidium equals Ochroconis. As noted by Gams (2015) and Shen et al. (2020), although the ex-type strain of S. terreum is sterile, there are many reliably named cultures of S. terreum globally, which clearly define the identity of this characteristic fungus. Scolecobasidium will always be applied to the clade that includes the type species S. terreum and O. constricta of Scolecobasidium and Ochroconis. This study shows that the clade defined as Scolecobasidium combines monophyly, sexual and asexual morphs, and ecological characters in a coherent way that can logically be recognised at the generic rank. Additionally, many species of Ochroconis for which DNA data are available have since been transferred to Scolecobasidium by Shen et al. (2020), except O. ferulica, O. guangxiensis, O. helicteris O. mirabilis and O. terricola, which are discussed below.

Scolecobasidium camellicola T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 840926; Fig. 5

Etymology. The epithet refers to *Camellia*, the host genus from which this fungus was collected.

Typus. CHINA, Guizhou Province, Meitan County, N27°75'09" E107°47'99", 910 m a.s.l., isolated from decaying *Camellia sinensis* leaf litter, 10 Aug. 2019, *T.P. Wei* (holotype HGUP 18242, isotype CGMCC 3.20547, culture ex-type GUCC 18242).

Mycelium partly superficial, partly immersed, hyphae branched, pale brown, septate, smooth, thick-walled, 1–2 µm wide. *Conidiophores* arising directly from superficial hyphae, mostly unbranched, subcylindrical, straight or flexuous, brown, continuous or septate, $(10-)11.5-55(-61.5) \times 2.5-4$ µm (av. ± SD = 19.4 ± 13.7 × 2.6 ± 0.4 µm, n = 30). *Conidiogenous cells* integrated, polyblastic, intercalary or terminal, sympodial extensions, pale brown to brown, bearing 1–6 conidia at the apex, $(6-)7.5-13(-14) \times (2-)2.5-3.5$ µm (av. ± SD = $9.2 \pm 2.1 \times 2.6 \pm 0.4$ µm, n = 30). *Conidia* secession rhexolytic from conidiogenous cells, subcylindrical or fusoid, 1-septate, minutely echinulate, pale brown, slightly constricted at the septum, with hilum bearing a marginal frill, $(7-)7.5-10.5(-11.5) \times (2.5-)3-4.5$ µm (av. ± SD = $8.7 \pm 0.9 \times 3.4 \pm 0.5$ µm, n = 30).

Culture characteristics — Colonies on PDA attaining 21– 22 mm diam after 14 d at 26 °C, growing slow, isabelline, raised in the centre. On OA reaching up to 23–24 mm diam, flat, spreading, immersed, dark brown. On MEA reaching 22–24 mm diam, raised, with sparse to moderate aerial hyphae, pale to medium brown.

Additional materials examined. CHINA, Guizhou Province, Meitan County, N27°75'09" E107°47'99", 910 m a.s.l., on decaying *Camellia sinensis* leaf litter, 10 Aug. 2019, *T.P. Wei* (HGUP 18243), living culture GUCC 18243 = CGMCC 3.20548; Leishan County, N26°24'02" E107°77'22", 1178 m a.s.l., from forest litter, 12 Mar. 2018, *T.P. Wei* (HGUP 18244), living culture GUCC 18244 = CGMCC 3.20549.

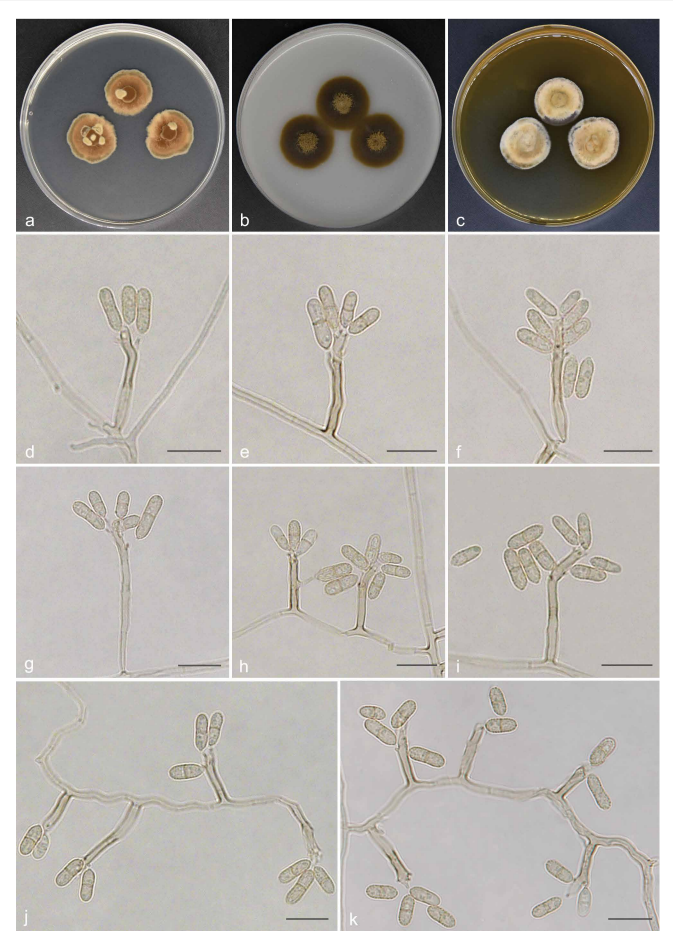


Fig. 5 Scolecobasidium camellicola (culture ex-type GUCC 18242). a–c. Colony on PDA, OA and MEA; d–h. conidiogenous cells giving rise to conidia; i. aging conidia forming conidiogenous loci; j–k. hypha with conidiogenous cells and conidia. — Scale bars: $d-k = 10 \mu m$.



Fig. 6 Scolecobasidium coiledmyces (culture ex-type GUCC 18245). a-c. Colony on PDA, OA and MEA; d-g. conidiophores arising from hyphal coils; h. conidiophores with conidiogenous cells and conidia; i. branched conidiophores. — Scale bars: $d-i = 10 \ \mu m$.

Notes — Scolecobasidium camellicola is introduced as a new species based on morphological and phylogenetic differences to other Scolecobasidium species. Phylogenetically, S. camellicola shares a sister relationship with S. mirabilis and S. helicteris with high statistical support (Fig. 2). Nevertheless, S. camellicola showed high heterogeneity, forming a wellseparated clade, which was genetically distant from all species. Morphologically, S. mirabilis differs from S. camellicola by its smooth-walled to verruculose and larger conidia (9.0–13.5 × 4.8–6.7 µm vs 7–11.5 × 2.5–4.5 µm) (Samerpitak et al. 2014); S. helicteris differs by its smooth to verruculose, ellipsoid or pyriform and smaller conidia (4–8 × 2–3.4 µm vs 7–11.5 × 2.5–4.5 µm) (Singh et al. 2019).

Scolecobasidium coiledmyces T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 840927; Fig. 6

Etymology. The epithet refers to the frequently forming hyphal coils.

Typus. CHINA, Guizhou Province, Guiyang City, Huaxi Wetland Park, N26°43'92" E106°67'76", 1140 m a.s.l., isolated from lawn soil, 16 Nov. 2018, *T.P. Wei* (holotype HGUP 18245, isotype CGMCC 3.20550, culture ex-type GUCC 18245).

Mycelium consisting of smooth, septate, branched, subhyaline or medium brown, $1.5-3 \mu m$ diam hyphae, forming hyphal coils. *Conidiophores* erect, 0(-4)-septate, occasionally branched, brown to dark brown, smooth and thick walled, subcylindrical, $(12-)13-43(-56) \times 2.5-4(-4.5) \mu m$ (av. \pm SD = $23.9 \pm 9.5 \times$ $3.1 \pm 0.5 \mu m$, n = 30). *Conidiogenous cells* terminal, subhyaline or pale brown, sympodially proliferating, producing several cylindrical denticles in the apical region, $(5-)7-17.5(-19) \times 2.5-3 \mu m$ (av. \pm SD = $12.4 \pm 3.9 \times 2.6 \pm 0.3 \mu m$, n = 30). *Conidia* solitary, medianly 1-septate, subcylindrical, apex obtuse, frills remaining on denticle and on conidial hilum, $0.5 \mu m$ long, medium brown, verruculose, released by rhexolytic secession, $8-11.5(-12) \times$ $2.5-3.5 \mu m$ (av. \pm SD = $9.6 \pm 1.1 \times 2.7 \pm 0.3 \mu m$, n = 30).

Culture characteristics — Colonies on PDA attaining 24– 26 mm diam after 14 d at 26 °C, brown, slightly raised in the centre, with moderate aerial mycelium and smooth. On OA reaching up to 20–22 mm diam, flat, spreading, dark brown. On MEA reaching 21–22 mm diam, raised, with moderate aerial mycelium, olivaceous.

Notes — *Scolecobasidium coiledmyces* is phylogenetically related to *S. cordanae* and *S. ailanthi*, but *S. coiledmyces* forms a single branch as the sister clade to the other two species with high support from three independent algorithms (Fig. 2). Furthermore, *S. coiledmyces* is distinct based on its morphology. The conidia of *S. cordanae* are smaller (5–10 × $2.5-3.5 \mu$ m vs $8-12 \times 2.5-3.5 \mu$ m), obovoidal to broadly fusoid and constricted at the median septum (Samerpitak et al. 2014); *S. ailanthic* differed from our strain in having fusoid, longitudinally striate and smaller conidia (9–10 × $2.4-2.6 \mu$ m vs $8-12 \times 2.5-3.5 \mu$ m) with a thick septum, as well as unbranched conidiophores (Jayasiri et al. 2019).

Scolecobasidium echinulatum T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 842082; Fig. 7

Etymology. The epithet refers to the conidia with minutely echinulate cell walls.

Mycelium superficial or immersed, hyphae brown, smooth, thinwalled, septate, $1.5-3 \mu m$ wide. *Conidiophores* clearly differentiated, arising at right angles from creeping hyphae, branched, erect, straight or slightly flexuous, brown, smooth, septate, (12–) 13.5–70(–87) × (2.5–)3–4 µm (av. \pm SD = 29.7 \pm 18.1 × 3.1 \pm 0.3 µm, n = 30). *Conidiogenous cells* integrated, terminal or intercalary, elongate to cylindrical, with some scattered denticles in the apical region, pale brown, smooth, (4–)5–15(–20.5) × 2.5–4 µm (av. \pm SD = 9.8 \pm 3.9 × 3.1 \pm 0.4 µm, n = 30). *Conidia* ellipsoidal to cylindrical, verruculose or minutely echinulate, dark brown to black, 1(–2)-septate, slightly narrower around the middle, frills remaining on denticle and on conidial hilum, released by rhexolytic secession, 8.5–10(–11.5) × 4–5 µm (av. \pm SD = 9.5 \pm 0.9 × 4.3 \pm 0.3 µm, n = 30).

Culture characteristics — Colonies on PDA attaining 13– 18 mm diam after 14 d at 26 °C, spreading, dark olivaceous brown. On OA reaching up to 18–20 mm diam, colonies moderately expanding, immersed, flat, dark olivaceous brown. On MEA reaching 13–15 mm diam, olivaceous, raised, with moderate aerial mycelium.

Additional material examined. CHINA, Guizhou Province, Qingzhen City, Red maple lake scenic area, N26°54'35" E106°38'74", 1272 m a.s.l., from soil, 07 Aug. 2020, *T.P. Wei* (HGUP 18248), living culture GUCC 18248 = CGMCC 3.20553.

Notes — The proposed new species, *S. echinulatum*, is phylogenetically related to *S. phaeophorum* and *S. crassihumicola*, but they can be distinguished by their morphological characteristics and DNA sequence data. *Scolecobasidium echinulatum* differs from *S. phaeophorum* as it has ellipsoidal to cylindrical, verruculose or minutely echinulate, dark brown to black and 1(–2)-septate conidia (the cylindrical to fusoid conidia of *S. phaeophorum* are smooth-walled, pale brown and 1-septate) (Samerpitak et al. 2015b); *S. crassihumicola* differs from *S. echinulatum* by having 1(–3)-septate, non-constricted, ovoid to cylindrical and larger conidia (7.5–13 × 4.2–5.5 µm vs $8.5-11.5 \times 4-5 µm$) (Matsushima 1971).

Scolecobasidium obovoideum T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 842083; Fig. 8

Etymology. The epithet refers to the obovoidal conidia.

Typus. CHINA, Guizhou Province, Guiyang City, Tianhetan Tourist Holiday Resort, N26°43'95" E106°57'64", 1164 m a.s.l., isolated from forest litter, 16 Dec. 2019, *T.P. Wei* (holotype HGUP 18246, isotype CGMCC 3.20551, culture ex-type GUCC 18246).

Mycelium composed of hyaline to pale brown, septate, branched, smooth, thick-walled, $1.5-3 \mu m$ wide hyphae. *Conidiophores* arising directly from vegetative hyphae, subcylindrical, branched, multi-septate, brown, erect, straight or flexuous, $(7-)9.5-39.5(-43) \times (2-)2.5-3.5 \mu m$ (av. \pm SD = $20.0 \pm 9.9 \times 2.6 \pm 0.4 \mu m$, n = 30). *Conidiogenous cells* polyblastic, terminal or intercalary, subcylindrical to subclavate, pale brown, producing conidia sympodially on long open denticles, $6-14(-15.5) \times 2-3.5 \mu m$ (av. \pm SD = $9.4 \pm 2.1 \times 2.6 \pm 0.4 \mu m$, n = 30). *Conidia* solitary, obovoidal to fusoid, sometimes slightly apiculate at the base, finely verruculose, 1-septate, constricted at the septum, brown, released by rhexolytic secession, $5.5-8.5 \times 2.5-4 \mu m$ (av. \pm SD = $7.2 \pm 0.6 \times 3.1 \pm 0.3 \mu m$, n = 30).

Culture characteristics — Colonies on PDA reaching up to 19–20 mm diam after 14 d at 26 °C, with moderate aerial mycelium, dark brown. On OA reaching 24–26 mm diam, flat, immersed, olivaceous. On MEA reaching 19–21 mm diam, raised, aerial mycelium moderate to abundant, olive grey.

Notes — Scolecobasidium obovoideum is phylogenetically closely related to *S. pandanicola* and *S. dracaenae* and can be differentiated from that species by DNA sequences of ITS, LSU, SSU, act1, tub2 and tef1 gene regions. Morphologically, *S. pandanicola* can be distinguished from *S. obovoideum* as it has subhyaline to hazel brown, thin-walled, fusoid or ellipsoid

Typus. CHINA, Guizhou Province, Guiyang City, Huaxi Wetland Park, N26°43'92" E106°67'76", 1140 m a.s.l., isolated from soil, 16 Nov. 2018, *T.P. Wei* (holotype HGUP 18247, isotype CGMCC 3.20552, culture ex-type GUCC 18247).

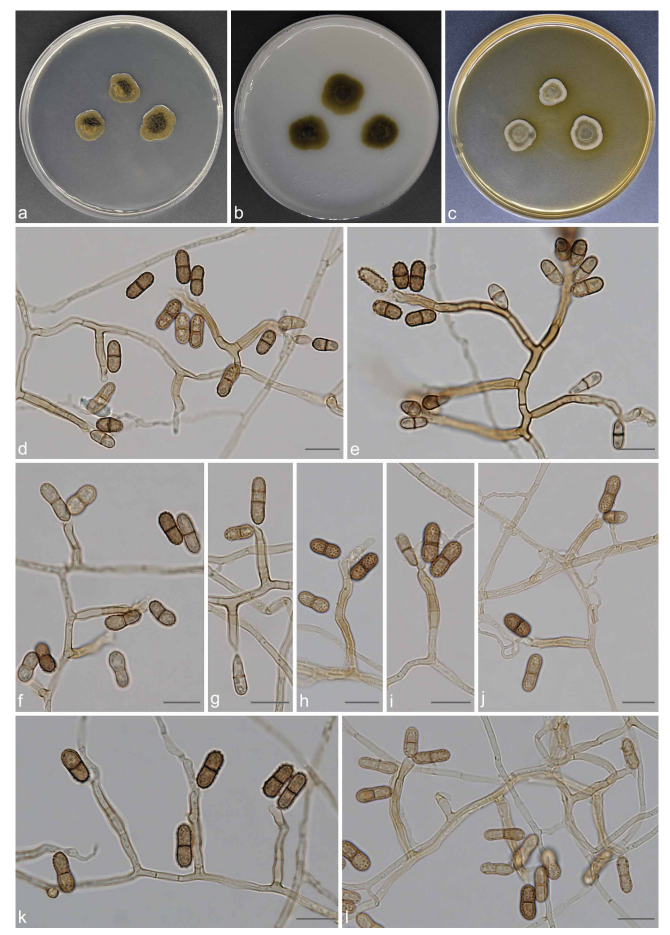


Fig. 7 Scolecobasidium echinulatum (culture ex-type GUCC 18247). a–c. Colony on PDA, OA and MEA; d–f. conidial apparatus with rhexolytic conidia, produced from sympodial conidiogenous cells; g. immature and 1(–2) septate conidia; h–I. conidiophores with conidiogenous cells and conidia. — Scale bars: $d-I = 10 \ \mu m$.

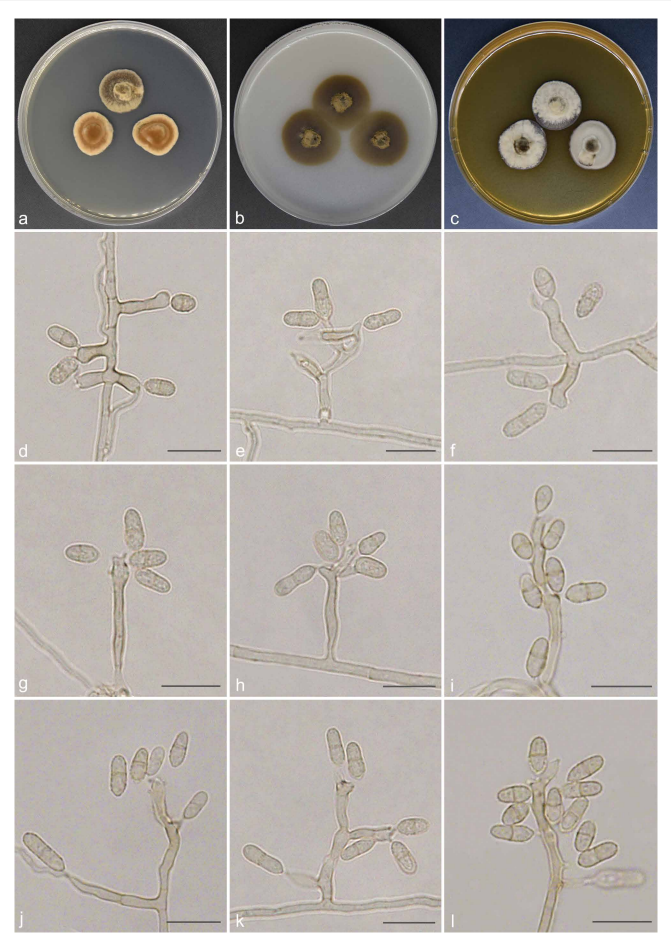


Fig. 8 Scolecobasidium obovoideum (culture ex-type GUCC 18246). a–c. Colony on PDA, OA and MEA; d–i. conidial apparatus with rhexolytic conidia, produced from sympodial conidiogenous cells; j–k. branched conidiophores; l. conidiophores with conidiogenous cells and conidia. — Scale bars: d–I = 10 μm.



Fig. 9 Scolecobasidium verrucaria (culture ex-type GUCC 18240). a–c. Colony on PDA, OA and MEA; d–f. conidiophores with conidiogenous cells and conidia; g. 1(-2) septate and coarsely verrucose conidia; h–p. maturation process of conidia. — Scale bars: d–p = 10 μ m.

and larger conidia (6–10 × 3–4.5 µm vs 5.5–8.5 × 2.5–4 µm) (Crous et al. 2015); S. *dracaenae* differs from S. *obovoideum* in having subcylindrical and larger conidia (6.5–10 × 3–4 µm vs 5.5–8.5 × 2.5–4 µm), and shorter conidiophores (10–30 × 2–3 µm vs 7–43 × 2–3.5 µm) (Crous et al. 2016). Phylogenetically, our new isolate GUCC 18246 forms a single clade separated from other *Scolecobasidium* species (Fig. 2).

Scolecobasidium verrucaria T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 840924; Fig. 9

Etymology. The epithet refers to its verrucose conidia.

Typus. CHINA, Guizhou Province, Qingzhen City, Red maple lake scenic area, N26°54'35" E106°38'74", 1272 m a.s.l., from soil, 16 Apr. 2018, *T.P. Wei* (holotype HGUP 18240, isotype CGMCC 3.20545, culture ex-type GUCC 18240).

Mycelium mostly superficial or semi-immersed, hyphae pale brown, smooth, branched, 1-2 µm wide. Conidiophores arising directly from vegetative hyphae, occasionally branched, continuous or septate, dark brown, straight or slightly geniculate, cylindrical, $(12.5-)14.5-51.5(-54) \times 3-3.5 \ \mu m$ (av. ± SD = 30.1 ± 11.8 × 2.9 ± 0.2 µm, n = 30). Conidiogenous cells integrated, terminal, polyblastic, sympodial, cylindrical, subhyaline, (6.5–)7–16.5(–17.5) × 2.5–3.5 µm (av. ± SD = 10.2 ± $2.8 \times 2.7 \pm 0.3 \mu m$, n = 30), with one or more denticles in the apical region, denticles 1-3 µm long. Conidia acropleurogenous, broadly ellipsoidal, 1(-2)-septate, strongly constricted at the septum, $8.5-11(-11.5) \times 4.5-5.5(-6) \mu m$ (av. ± SD = $9.3 \pm$ $0.8 \times 5.0 \pm 0.3 \,\mu$ m, n = 30), the colour of immature conidia changed from brown to yellow to dark brown, and gradually from smooth to coarsely verrucose, verrucous protrusions up to 2.7 µm long.

Culture characteristics — Colonies on PDA reaching up to 20–22 mm diam after 14 d at 26 °C, slightly raised at centre, isabelline, lobate margin. On OA reaching 24–26 mm diam, flat, spreading, immersed, olivaceous. On MEA reaching 16–18 mm diam, raised, hairy, grey brown.

Notes — *Scolecobasidium verrucaria* was collected from natural forest soil, and has the same lifestyle as some species of *Scolecobasidium*. Morphologically, the polyblastic sympodial conidiogenous cells, conidial apparatus with rhexolytic conidiogenesis, and olivaceous colonies point to *Scolecobasidium*. It is noteworthy that the ellipsoidal, 1(–2)-septate, yellow to dark brown, and coarsely verrucose conidia of *S. verrucaria* differ from other reported members of *Scolecobasidium*. Phylogenetically, *S. verrucaria* nests in the *Scolecobasidium* clade, being closely related to *S. verrucaria* by its brown grey, verruculose, 0(–1)-septate and larger conidia (8.5–14 × 4–5.5 µm vs 8.5–11.5 × 4.5–6 µm).

Scolecobasidium zunyiense T.P. Wei & Y.L. Jiang, sp. nov. — MycoBank MB 840925; Fig. 10

Etymology. Named after Zunyi, where this species was collected.

Typus. CHINA, Guizhou Province, Zunyi City, Phoenix Mountain National Forest Park, N27°36'27" E106°45'07", 1024 m a.s.l., from forest litter, 10 June 2019, *T.P. Wei* (holotype HGUP 18241, isotype CGMCC 3.20546, culture ex-type GUCC 18241).

Mycelium consisting of pale brown, smooth, branched, septate, 1.5–2.5 µm diam hyphae, often giving rise to hyphal coils. *Conidiophores* branched, straight to irregularly curved, solitary or at times two arising from the same basal cell, smooth, subcylindrical, septate, pale brown to brown, $(9-)11-60(-78) \times 3-4$ µm (av. ± SD = 32.1 ± 15.9 × 3.2 ± 0.3 µm, n = 30). *Conidiogenous cells* polyblastic, sympodial, terminal or intercalary, subhyaline to brown, smooth, subcylindrical, $(5-)5.5-15.5(-17.5) \times 3-4 \mu m$ (av. \pm SD = 9.8 $\pm 2.9 \times 3.3 \pm 0.3 \mu m$, n = 30), with 1–9 terminal denticles, 1–2 μm . *Conidia* released by rhexolytic secession, solitary, acropleurogenous, ellipsoidal with rounded ends, 0(–1)-septate, brown grey, verruculose, prominently constricted at the septum, $8.5-13(-14) \times 4-5.5 \mu m$ (av. \pm SD = $10.2 \pm 1.2 \times 4.4 \pm 0.4 \mu m$, n = 30).

Culture characteristics — Colonies on PDA attaining 18– 20 mm diam after 14 d at 26 °C, slightly raised at centre, immersed, isabelline. On OA reaching up to 20–23 mm diam, flat, spreading, dark brown, reverse brown. On MEA reaching 20–22 mm diam, raised, hairy, pale brown at the surface, reverse olivaceous.

Notes — Scolecobasidium zunyiense is phylogenetically related to *S. verrucaria*, being fully supported in three independent algorithms (Fig. 2). Nevertheless, they can be distinguished based on their morphological characteristics as *S. zunyiense* is characterised by brown grey, 0(-1)-septate and verruculose conidia, while the conidia of *S. verrucaria* are yellow to dark brown, 1(-2)-septate, and verrucous protrusions can be up to 2.7 µm long, a feature not observed for *S. zunyiense*.

Scolecobasidium ferulica (Z. Tazik & K. Rahnama) T.P. Wei & Y.L. Jiang, comb. nov. — MycoBank MB 840930

Basionym. Ochroconis ferulica Z. Tazik & K. Rahnama, Nova Hedwigia 110: 374. 2020.

Description — Tazik et al. (2020).

Notes — *Scolecobasidium ferulica* was initially reported as an endophyte on roots of *Ferula ovina* in northeast Iran (Tazik et al. 2020). Multi-locus phylogenetic analyses indicate that this species is sister to *S. longiphorum* and is fully supported as phylogenetically distinct (Fig. 2). Morphologically, the ellipsoidal and smaller conidia of *S. ferulica* (8–10.5 × 6–7.5 µm) distinguishes this species from *S. longiphorum* (12–19 × 3.5–4.5 µm), which is characterised by long cylindrical and larger conidia (Samerpitak et al. 2014).

Scolecobasidium guangxiensis (Xie et al.) T.P. Wei & Y.L. Jiang, comb. nov. — MycoBank MB 840931

Basionym. Ochroconis guangxiensis Xie et al., Mycoscience 61: 308. 2020

Description — Chen et al. (2020).

Notes — Scolecobasidium guangxiensis was collected from rhizosphere soil of sugarcane in Guangxi, China, and was introduced by Chen et al. (2020). Based on the phylogeny presented here, despite the *S. guangxiensis* shares a sister relationship with *S. musicola* (Fig. 2), the considerably high number of variable positions in the ITS (88 bp, 17 %) and LSU (55 bp, 6 %) alignments supports the split into two distinct taxa. No SSU, *act1*, *tub2* and *tef1* data are currently available for the ex-type of *S. guangxiensis* and *S. musicola*. Morphologically, it is characterised by forming flask-shaped or cylindrical conidio-phores and smooth-walled to verruculose, yellow brown, clavate or cylindrical, 1(–3)-septate conidia; *S. musicola* differs by its sexual morph forming fusoid to ellipsoid, hyaline to pale brown and straight to slightly curved ascospores (Crous et al. 2018b), whereas *S. guangxiensis* lacks a sexual morph.

Scolecobasidium helicteris (Singh et al.) T.P. Wei & Y.L. Jiang, comb. nov. — MycoBank MB 840932

Basionym. Ochroconis helicteris Singh et al., Phytotaxa 427: 192. 2019.

Description — Singh et al. (2019).

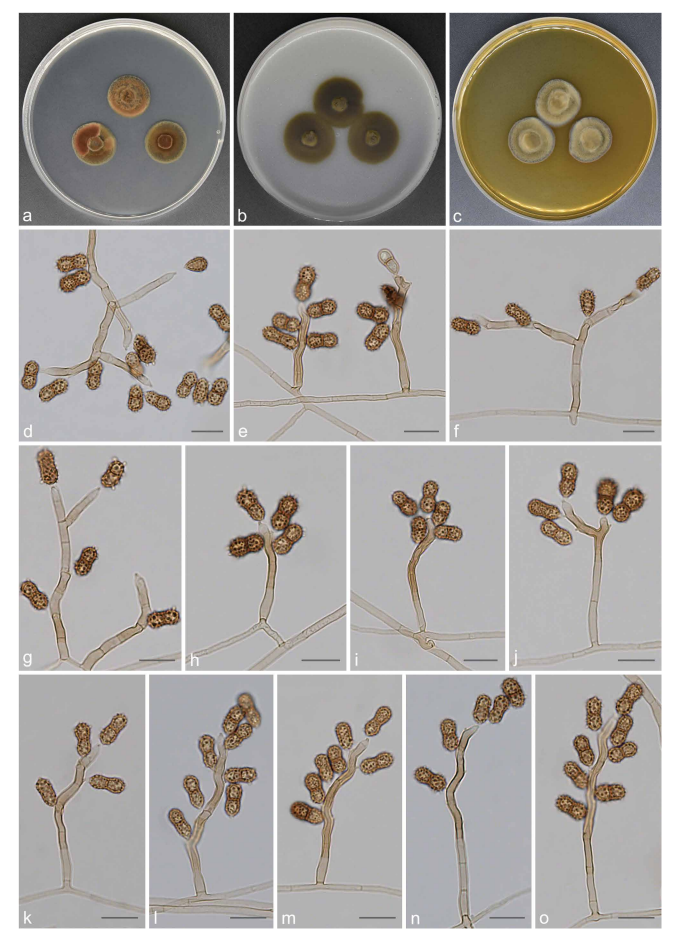


Fig. 10 Scolecobasidium zunyiense (culture ex-type GUCC 18241). a-c. Colony on PDA, OA and MEA; d-e. conidiophores and conidiogenous cells bearing conidia; f-g. branched conidiophores; h-o. acropleurogenous, ellipsoidal, 1-septate and verruculose conidia. — Scale bars: d-o = 10 μ m.



Fig. 11 Scolecobasidium leishanicola (GUCC 18259). a–c. Colony on PDA, OA and MEA; d–e. conidial apparatus with rhexolytic conidia, produced from sympodial conidiogenous cells; f–I. germinating conidia; j–m. maturation process of conidia. — Scale bars: d–m = 10 µm.

Notes — Scolecobasidium helicteris was previously recorded as pathogen associated with leaf spots on *Helicteris isora* in India (Singh et al. 2019). According to our analysis, *S. helicteris* resolved as the closest phylogenetic relative to *S. ellipsoideum* (Fig. 2). *Scolecobasidium helicteris* is, however, clearly distinguished morphologically by its obovoid to fusoid or pear shaped and smooth to verruculose conidia, and shorter conidiophores (ellipsoidal to oblong and spinulose conidia in *S. ellipsoideum*) (Fig. 13; Ren et al. 2013).

Scolecobasidium leishanicola (X. Zhang & Y.L. Jiang) T.P. Wei & Y.L. Jiang, comb. & nom. nov. — MycoBank MB 840929; Fig. 11

Basionym. Ochroconis terricola X. Zhang & Y.L. Jiang, Mycotaxon 135: 146. 2020.

Mycelium consisting of branched, septate, subhyaline to pale brown, smooth-walled, $1.5-2.5 \mu m$ diam hyphae. *Conidiophores* arising directly from vegetative hyphae, erect, subcylindrical, unbranched, medium brown, multi-septate, smoothwalled, straight to flexuous, $(9.5-)11-41(-108) \times 2.5-4 \mu m$ (av. \pm SD = 26.1 \pm 18.6 \times 3.2 \pm 0.4 μm , n = 30). *Conidiogenous cells* integrated, terminal, subcylindrical, subhyaline to pale brown, with one to several sympodial denticle-like loci, $(7.5-)9-23.5(-25) \times (2.5-)3-4 \mu m$ (av. \pm SD = 13.4 \pm 4.6 \times 3.1 \pm 0.3 μm , n = 30). *Conidia* solitary, brown, ellipsoidal to cylindrical or fusoid, 1-septate, minutely echinulate, sometimes slightly constricted at the septum, apex obtuse, base narrowly truncated with hilum bearing a marginal frill, $9.5-12(-13.5) \times$ $3-4.5 \mu m$ (av. \pm SD = 11.6 \pm 1.5 \times 3.5 \pm 0.4 μm , n = 30).

Culture characteristics — Colonies on PDA attaining 22– 24 mm diam after 14 d at 26 °C, effuse, grey brown, reverse dark brown, growing slowly. On OA reaching up to 24–26 mm diam, olivaceous brown, immersed, with sparse aerial mycelium. On MEA reaching 18–23 mm diam, spreading, hairy, isabelline, raised in the centre.

Material examined. CHINA, Guizhou Province, Leishan County, N26°24'02" E107°77'22", 1178 m a.s.l., isolated from soil, 12 Mar. 2018, *T.P. Wei* (HGUP 18259), living culture GUCC 18259 = CGMCC 3.20564.

Notes — Scolecobasidium leishanicola (as Ochroconis terricola) was previously recorded from soil in China (Zhang et al. 2020). The phylogenetic tree constructed based on six gene loci showed that one isolate (GUCC 18259) from the present study clusters in a clade closely related to *S. leishanicola*, and was sister to *S. sexualis* and *S. podocarpicola* (Fig. 2). Morphologically, the conidial dimensions in the present study fit exactly with those in Zhang et al. (2020). Therefore, this species was placed in *Scolecobasidium*, for which a new name had to be introduced (*S. leishanicola*), as *S. terricola* was already occupied.

Scolecobasidium mirabilis (Samerp. & de Hoog) T.P. Wei & Y.L. Jiang, comb. nov. — MycoBank MB 840933

Basionym. Ochroconis mirabilis Samerp. & de Hoog, Fungal Diversity 65: 114. 2014.

Description — Samerpitak et al. (2014).

Notes — *Scolecobasidium musae* was originally isolated from the fruit surface of *Musa basjoo* in Hainan, China (Hao et al. 2013). Almost simultaneously, Samerpitak et al. (2014) described *S. mirabilis* from the regulator of a scuba diver in the Netherlands. Later, Samerpitak et al. (2015a) compared DNA sequence data between the two species and thought that the LSU sequences of *S. mirabilis* and *S. musae* were almost identical. To solve this taxonomic dilemma, they proposed *S. mirabilis* as a synonym for *S. musae*. However, our phylogenetic analysis indicates that *S. mirabilis* and *S. musae* cluster apart, and are phylogenetically distant (Fig. 2). They are genetically distinct in 2 bp (1 %), 21 bp (3 %), 3 bp (1 %), 10 bp (4 %), 46 bp (10 %) and 16 bp (4 %) in SSU, ITS, LSU, *act1*, *tub2* and *tef1* loci. Based on these results as well as their morphology, we thus resurrect *S. mirabilis* and recognise *S. mirabilis* and *S. musae* as two different species of *Scolecobasidium*.

Scolecobasidium constrictum E.V. Abbott, Mycologia 19: 30. 1927 — Fig. 12

Synonym. Ochroconis constricta (E.V. Abbott) de Hoog & Arx, Kavaka 1: 57. 1973.

Mycelium composed of hyaline or pale brown, septate, branched, smooth-walled, 1.5-2 µm diam hyphae. Conidiophores differentiated arising directly from vegetative hyphae, branched, sparingly septate, brown, smooth and thick-walled, cylindrical, mostly geniculate-sinuous, (11-)12-46.5(-56) × (2-)2.5- $3.5 \,\mu\text{m}$ (av. $\pm \text{SD} = 23.8 \pm 10.3 \times 2.5 \pm 0.3 \,\mu\text{m}$, n = 30). Conidiogenous cells integrated, terminal, flask-shaped or ampulliform to cylindrical, hyaline or pale brown, with single or several sympodial apical loci with rhexolytic conidiogenesis, $(3.5-)5.5-14(-15) \times (2-)2.5-4(-4.5) \ \mu m \ (av. \pm SD = 8.8 \pm 10^{-1})$ $3.1 \times 3.0 \pm 0.5 \,\mu\text{m}$, n = 30). Conidia solitary, subhyaline to pale brown, 1-septate, broadly ellipsoidal to cylindrical, finely echinulate to verruculose, usually constricted at the septum, frills remaining on denticle and on conidial base, ends obtusely rounded, $9.5-12 \times 3.5-4.5 \mu m$ (av. $\pm SD = 10.3 \pm 0.6 \times 3.9 \pm$ $0.3 \,\mu m, n = 30$).

Culture characteristics — Colonies on PDA attaining 14– 15 mm diam after 14 d at 26 °C, effuse, hairy, olivaceous brown, raised, usually slow-growing. On OA reaching up to 18–20 mm diam, cottony to floccose at centre, glabrous at periphery, grey olivaceous. On MEA reaching 14–15 mm diam, aerial mycelium moderate, olivaceous black, raised in the centre.

Materials examined. CHINA, Guizhou Province, Shibing County, Fodingshan National Nature Reserve, N27°06'47" E108°10'82", 1180 m a.s.l., from dead branches, 12 Sept. 2020, *T.P. Wei* (HGUP 18255), living culture GUCC 18255 = CGMCC 3.20560; Guiyang City, Huaxi District, N26°42'55" E106°68'07", 1140 m a.s.l., from forest litter, 6 May 2018, *T.P. Wei* (HGUP 18256), living culture GUCC 18256 = CGMCC 3.20561; Qingzhen City, N26°54'35" E106°38'74", 1272 m a.s.l., from soil, 16 Apr. 2018, *T.P. Wei* (HGUP 18257), living culture GUCC 18257 = CGMCC 3.20562; Guiyang City, Guanshan Lake Park, N26°64'28" E106°63'06", 1260 m a.s.l., from lawn soil, 5 Apr. 2021, *T.P. Wei* (HGUP 18258), living culture GUCC 18258 = CGMCC 3.20563.

Notes — *Scolecobasidium constrictum* was reported by Abbott (1927) as the type species of *Scolecobasidium*, which is characterised by having ampulliform or cylindrical conidiogenous cells and broadly ellipsoidal to cylindrical conidia. The phylogenetic result shows that our four isolates cluster together with *S. constrictum* and are well-supported (Fig. 2). Therefore, we identified our isolates as *S. constrictum* and provide an illustration of the species.

Scolecobasidium ellipsoideum Ren et al., Mycoscience 54: 422. 2013 — Fig. 13

Mycelium partly superficial or semi-immersed, hyphae simple branched, septate, pale brown, smooth-walled, $1.5-2 \mu m$ wide. *Conidiophores* erect, branched, slightly to distinctly geniculate-sinuous, cylindrical, multi-septate, brown, smooth, $(10.5-)11-64.5(-77.5) \times (2-)2.5-3.5 \mu m$ (av. \pm SD = $30.6 \pm 19.7 \times 2.6 \pm 0.3 \mu m$, n = 30). *Conidiogenous cells* terminal or intercalary, subcylindrical, pale to medium brown, producing conidia sympodially on long open denticles, $(4-)6-10(-11) \times 2-2.5(-3) \mu m$ (av. \pm SD = $8.2 \pm 2.2 \times 2.3 \pm 0.2 \mu m$, n = 30). *Conidia* solitary, ellipsoidal to oblong, 1-septate, pale to dark brown, spinulose, apex



Fig. 12 Scolecobasidium constrictum (GUCC 18256). a–c. Colony on PDA, OA and MEA; d–m. conidial apparatus with rhexolytic conidia, produced from sympodial conidiogenous cells; n–o. hyphal coil; p. cylindrical to fusoid conidia. — Scale bars: $d-p = 10 \mu m$.



Fig. 13 *Scolecobasidium ellipsoideum* (GUCC 18264). a–c. Colony on PDA, OA and MEA; d–f. conidiophores and conidiogenous cells bearing conidia; g. branched conidiophores; h-p. medium brown, 1-septate and ellipsoidal to cylindrical conidia. — Scale bars: $d-p = 10 \mu m$.



Fig. 14 Scolecobasidium globale (GUCC 18249). a–c. Colony on PDA, OA and MEA; d–k. hyphae, conidiophores with sympodially proliferating conidiogenous cells and cylindrical to pyriform and rhexolytic succession conidia; I. germinating conidia. — Scale bars: $d-l = 10 \ \mu m$.

subobtuse, base truncate with basal marginal frill, sometimes slightly constricted at the septum, released by rhexolytic secession, $7.5-9.5 \times 3-4 \mu m$ (av. ± SD = $8.2 \pm 0.7 \times 3.3 \pm 0.3 \mu m$, n = 30).

Culture characteristics — Colonies on PDA attaining 17– 25 mm diam after 14 d at 26 °C, spreading, yellow brown, with moderate aerial mycelium. On OA reaching up to 26–32 mm diam, immersed, olivaceous, with sparse aerial mycelium. On MEA reaching 20–24 mm diam, raised in the centre, grey olivaceous.

Materials examined. CHINA, Guizhou Province, Leishan County, N26°24'02" E107°77'22", 1178 m a.s.l., from soil, 12 Mar. 2018, *X. Zhang* (HGUP 18264), living culture GUCC 18264 = CGMCC 3.20569; Meitan County, N27°75'09" E107°47'99", 910 m a.s.l., from submerged wood, 10 Aug. 2019, *T.P. Wei* (HGUP 18265), living culture GUCC 18265 = CGMCC 3.20570; Zunyi City, N27°36'27" E106°45'07", 1024 m a.s.l., from forest litter, 10 June 2019, *T.P. Wei* (HGUP 18266), living culture GUCC 18266 = CGMCC 3.20571.

Notes — Scolecobasidium ellipsoideum was originally described as a saprophyte from soils in Guizhou Province, China (Ren et al. 2013). In this study, phylogenetic inference revealed that our three newly collected isolates clustered together with the ex-type strains of *S. ellipsoideum* (Fig. 2). The morphological characters of our studied specimens fit well with *S. ellipsoideum* (Fig. 13). Moreover, *S. ellipsoideum* differs from the phylogenetically related species *S. helicteris* by nucleotide differences in ITS (7 bp, 1 %) and *tub2* (GUCC 18264: 3 bp, 2 %). No LSU, SSU, act1 and tef1 are available for the ex-type of *S. ellipsoideum* and *S. helicteris*.

Scolecobasidium globale (Samerpitak et al.) Crous et al., Stud. Mycol. 96: 211. 2020 — Fig. 14

Basionym. Ochroconis globalis Samerpitak et al., Mycol. Progr. 14: 3. 2015.

Mycelium superficial or immersed, hyphae septate, hyaline to pale brown, smooth and thin-walled, $1.5-2 \mu m$ wide. *Conidiophores* differentiated arising directly from vegetative hyphae, straight to flexuous, cylindrical, multi-septate, branched, pale to dark brown, $(20.5-)27.5-190(-211.5) \times (2.5-)3-4.5(-5.5) \mu m$ (av. \pm SD = 99.7 \pm 59.8 \times 3.7 \pm 0.8 μm , n = 30). *Conidiogenous cells* terminal or intercalary, proliferating sympodially, with a single or more denticle-like conidiogenous loci, subhyaline to brown, $(6.5-)7.5-22(-30.5) \times 3.5-4.5(-5) \mu m$ (av. \pm SD = 12.7 \pm 6.8 \times 3.7 \pm 0.4 μm , n = 30). *Conidia* solitary, ellipsoidal to cylindrical, brown, 1(-3)-septate, smooth or somewhat verrucose, constricted at the septum, frills remaining visible on denticle and on conidial base, $(7-)8.5-10(-11.5) \times 4-5.5(-6) \mu m$ (av. \pm SD = 9.7 \pm 1.6 \times 5.1 \pm 0.7 μm , n = 30).

Culture characteristics — Colonies on PDA attaining 10– 11 mm diam after 14 d at 26 °C, surface yellow brown, growing slowly, with sparse aerial mycelium. On OA reaching up to 13–16 mm diam, moderately expanding, immersed, olivaceous. On MEA reaching 12–13 mm diam, raised, hairy, brownish olive green.

Materials examined. CHINA, Guizhou Province, Shiqian County, Ganxi Township, Fodingshan National Nature Reserve, N27°40'51" E108°07'21", 1240 m a.s.l., from forest humus, 2 Nov. 2019, *T.P. Wei* (HGUP 18249), living culture GUCC 18249 = CGMCC 3.20554; Guiyang City, Tianhetan Tourist Holiday Resort, N26°43'95" E106°57'64", 1164 m a.s.l., from soil, 26 Oct. 2020, *T.P. Wei* (HGUP 18250), living culture GUCC 18250 = CGMCC 3.20555.

Notes — Scolecobasidium globale was introduced by Samerpitak et al. (2015a). In this study, two newly collected isolates clustered together with *S. globale* and were fully supported phylogenetically (Fig. 2). The comparison of morphological characteristics of the three strains found them to be similar. However, our newly obtained isolates differ from the ex-type culture of *S. globale* in having more septate conidia (mostly 1-septate in CBS 119644 vs up to 1(–3)-septate in GUCC 18249) (Samerpitak et al. 2015a; Fig. 14), which may depend on the state (mature or immature) of the specimen being observed. Importantly, the ex-type culture (CBS 119644) of *S. globale* had the following nucleotide similarities with the sequences of our newly collected strain (GUCC 18249). On ITS, LSU, SSU, *act1*, *tub2* and *tef1*, respectively: 677/687 (99 %, including six gaps), 791/794 (99 %, including two gaps), 1463/1464 (99 %, including one gap), 334/348 (96 %, including 12 gaps), 459/488 (94 %, including seven gaps) and 508/513 (99 %, including one gap). Therefore, we identified our isolates as *S. globale*, and provide an illustration from a different host, representing a new record from China.

Scolecobasidium minimum (Fassat.) Crous et al., Stud. Mycol. 96: 212. 2020 — Fig. 15

Basionym. Humicola minima Fassat., Ceská Mykol. 21: 87. 1967.

Synonym. Ochroconis minima (Fassat.) Samerpitak & de Hoog, Fungal Diversity 65: 110. 2013.

Mycelium consisting of hyaline to yellow brown, smooth, septate, branched, 1.5-2.5 µm diam hyphae. Conidiophores hyaline to brown, cylindrical, septate, straight to flexuous, branched, mostly reduced to conidiogenous cells, $7-29(-36.5) \times 2.5-4 \,\mu m$ (av. ± SD = 14.6 ± 8.4 × 3.1 ± 0.5 μm, n = 30). Conidiogenous cells flask-shaped to clavate, pale brown, mostly standing at right angles from undifferentiated hyphae, with some scattered denticles in the apical region, $5.5-11.5(-15) \times 3-4 \ \mu m$ (av. ± SD = $8.6 \pm 2.1 \times 3.3 \pm 0.4 \mu m$, n = 30). Conidia acrogenous, yellow brown, 1-septate, smooth, somewhat T- or Y- shaped, composed of the main axis and two branches, branch form an angle of 45° with the apex of main axis; main axis $9-14.5 \times$ $3-4 \mu m$ (av. ± SD = 10.9 ± 1.3 × 3.3 ± 0.2 μm , n = 30), secondary branches $1.5-5 \times 2.5-4.5 \mu m$ (av. ± SD = $3.4 \pm 0.7 \times 10^{-5}$ km s = $3.4 \pm 0.7 \times 10^{-5}$ $3.6 \pm 0.3 \mu m$, n = 30). Chlamydospores spherical, dark brown, aseptate, smooth, arising directly from vegetative hyphae, $5-6 \mu m$ (av. $\pm SD = 5.4 \pm 0.2$, n = 30).

Culture characteristics — Colonies on PDA attaining 17– 21 mm diam after 14 d at 26 °C, dark brown at centre, pale yellowish at periphery, fluffy aerial mycelium. On OA reaching up to 20–23 mm diam, pale olivaceous brown, with sparse aerial mycelium. On MEA reaching 19–21 mm diam, raised, centre olivaceous to grey olivaceous and white toward the periphery.

Material examined. CHINA, Guizhou Province, Shiqian County, Fodingshan National Nature Reserve, N27°40'49" E108°07'35", 1100 m a.s.l., from forest humus, 2 Nov. 2019, *T.P. Wei* (HGUP 18260), living culture GUCC 18260 = CGMCC 3.20565.

Notes — In the current study, the phylogenetic result shows that our new collection GUCC 18260 clusters together with *S. minimum*, sharing a sister relationship to *S. ramosum* and *S. icarus* with high statistical support from three independent algorithms (Fig. 2). All three species have T- or Y-shaped conidia. Chen et al. (2020) obtained *S. minimum* from sugarcane and banana rhizospheres in Guangxi. Therefore, this is the second report of this species from China.

Scolecobasidium ramosum (Giraldo et al.) Crous et al., Stud. Mycol. 96: 212. 2020 — Fig. 16

Basionym. Ochroconis ramosa Giraldo et al., J. Clinical Microbiol. 52: 4197. 2014.

Mycelium composed of branched, septate, hyaline to pale brown, smooth-walled, $1.5-2 \mu m$ diam hyphae. *Conidiophores* pale brown, clavate or cylindrical with beaked apex, simple or sympodially branched, often reduced to conidiogenous cells, $7.5-64 (-69.5) \times 3-4 \mu m$ (av. \pm SD = $21.2 \pm 14.6 \times 3.1 \pm 0.3 \mu m$, n = 30).



Fig. 15 Scolecobasidium minimum (GUCC 18260). a–c. Colony on PDA, OA and MEA; d–l. conidial apparatus with rhexolytic conidia, produced from sympodial conidiogenous cells; n. T- or Y- shaped conidia; m, o. chlamydospores. — Scale bars: d–o = 10 μm.

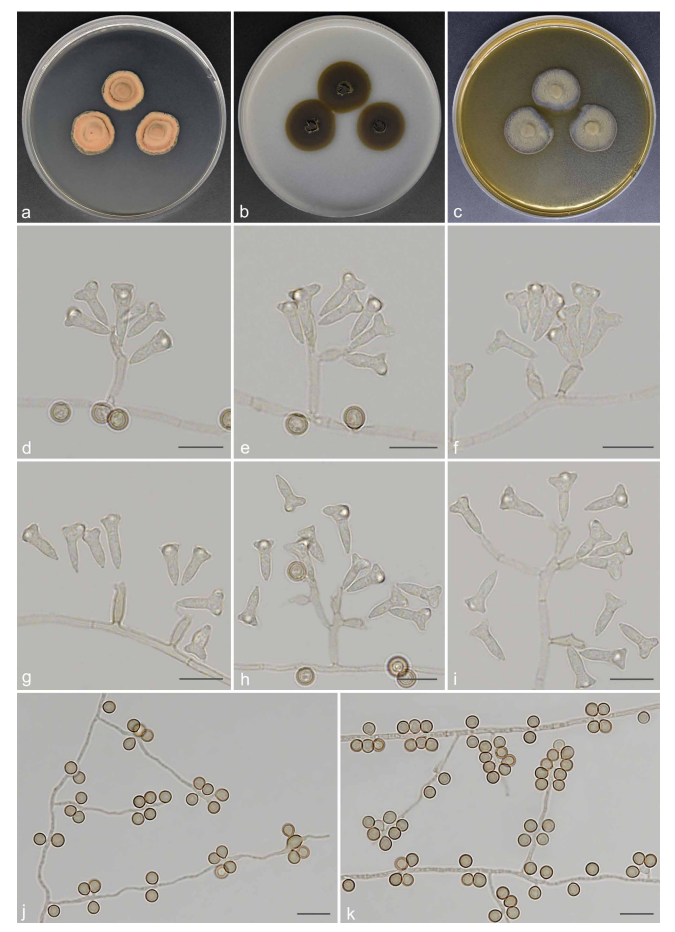


Fig. 16 Scolecobasidium ramosum (GUCC 18261). a-c. Colony on PDA, OA and MEA; d-i. hyphae, conidiophores with sympodially proliferating conidiogenous cells and T- or Y- shaped and rhexolytic succession conidia; j-k. chlamydospores. — Scale bars: $d-k = 10 \ \mu m$.

Conidiogenous cells subhyaline to pale brown, bearing one or more denticles in the apical region, ampulliform to cylindrical, $5.5-14.5(-17.5) \times 2.5-4 \ \mu m$ (av. $\pm SD = 9.1 \pm 2.9 \times 3.1 \pm$ 0.4 μ m, n = 30). Conidia solitary, pale brown, 1-septate, trilobate, T- or Y-shaped, smooth or verrucose, base slightly tapered to truncate hilum, main axis $10-13.5 \times 3-4 \ \mu m$ (av. $\pm SD = 11.3 \pm$ 0.9 $\times 3.5 \pm 0.2 \ \mu m$, n = 30), secondary branches $1.5-6.5 \times$ $3-5.5 \ \mu m$ (av. $\pm SD = 3.5 \pm 1.2 \times 4 \pm 0.4 \ \mu m$, n = 30), released by rhexolytic secession. *Chlamydospores* solitary, growing directly on vegetative hyphae, globose or subglobose, aseptate, dark brown, sessile, smooth, $4.5-5.5 \ \mu m$ (av. $\pm SD = 4.5 \pm$ 0.2 μm , n = 30).

Culture characteristics — Colonies on PDA attaining 17– 20 mm diam after 14 d at 26 °C, slightly raised, yellow-brown, spreading. On OA reaching up to 21–23 mm diam, flat, chocolate brown, felty at center, membranous toward the periphery. On MEA reaching 19–20 mm diam, olivaceous grey, velvety to cottony, slightly raised to umbonate.

Materials examined. CHINA, Guizhou Province, Zunyi City, Phoenix Mountain National Forest Park, N27°36'27" E106°45'07", 1024 m a.s.l., from forest litter, 10 June 2019, *T.P. Wei* (HGUP 18261), living culture GUCC 18261 = CGMCC 3.20566; Qingzhen City, N26°54'35" E106°38'74", 1272 m a.s.l., from soil, 16 Apr. 2018, *T.P. Wei* (HGUP 18262), living culture GUCC 18262 = CGMCC 3.20567; Leishan County, N26°24'02" E107°77'22", 1178 m a.s.l., from forest litter, 12 Mar. 2018, *T.P. Wei* (HGUP 18263), living culture GUCC 18263 = CGMCC 3.20568.

Notes — *Scolecobasidium ramosum* is phylogenetically closely related to *S. minimum* and *S. icarus* (Fig. 2). The sequences of ITS, LSU, SSU, *act1*, *tub2* and *tef1* gene regions can differentiate *S. ramosum* from these two species. This species was originally collected from human nails in the USA (Giraldo et al. 2014). Recently, Chen et al. (2020) isolated this species from the soil rhizosphere of sugarcane, and the strain we obtained was derived from forest litter and soil, which indicates that *S. ramosum* has both a parasitic and saprophytic lifestyle.

Scolecobasidium tshawytschae (Doty & D.W. Slater) McGinnis & Ajello, Trans. Brit. Mycol. Soc. 63: 202. 1974 — Fig. 17

Basionym. Heterosporium tshawytschae Doty & D.W. Slater, Amer. Midl. Naturalist 36: 663. 1946.

Synonym. Ochroconis tshawytschae (Doty & D.W. Slater) Kiril. & Al-Achmed, Mykrobiol. Zhurn. 39: 305. 1977.

Mycelium superficial or immersed, hyphae branched, septate, hyaline to pale brown, smooth and thin-walled, 1.5-2.5 µm wide. Conidiophores erect, cylindrical, straight to gently curved, septate, unbranched, mostly reduced to conidiogenous cells, $(5.5-)6.5-18.5(-24) \times 3-3.5(-4) \mu m (av. \pm SD = 11.6 \pm 5.1 \times 10^{-10})$ 3.2 ± 0.3 µm, n = 30). Conidiogenous cells integrated, terminal, often somewhat inflated, lageniform or ampulliform, with one to several conidiogenous loci, leaving minute collarettes on denticulate loci, $5-7.5(-9) \times 4-5 \mu m$ (av. $\pm SD = 6.5 \pm 1.1 \times 4.1 \pm$ 0.4 µm, n = 30). Conidia solitary, verrucose, pale to dark brown, 1(-3)-septate, cylindrical or slightly clavate, the colour of immature conidia changed from greenish olivaceous to yellow to dark brown, $12.5-22(-24.5) \times (4.5-)5-6(-6.5) \mu m$ (av. $\pm SD = 16.4$ \pm 1.6 \times 5.2 \pm 0.4 μ m, n = 30). Chlamydospores acrogenous, brown, smooth, 0(-1)-septate, ellipsoidal to fusoid, arising directly from vegetative hyphae, $7.5-13 \times (4-)4.5-6(-6.5) \mu m$ $(av. \pm SD = 9.7 \pm 1.9 \times 4.8 \pm 0.6 \mu m, n = 30).$

Culture characteristics — Colonies on PDA attaining 15– 18 mm diam after 14 d at 26 °C, olivaceous at centre, dark brown at periphery, slightly raised to umbonate. On OA reaching up to 20–22 mm diam, felty, olivaceous brown to olive, submerged mycelium. On MEA reaching 20–22 mm diam, slightly domed, woolly, with dense dark hairs at the centre and grey loose mycelium near the edge. *Materials examined*. CHINA, Guizhou Province, Guiyang City, Guanshan Lake Park, N26°64'28" E106°63'06", 1260 m a.s.l., from lawn soil, 5 Apr. 2021, *T.P. Wei* (HGUP 18251), living culture GUCC 18251 = CGMCC 3.20556; Zunyi City, N27°36'27" E106°45'07", 1024 m a.s.l., from plant litter, 10 June 2019, *T.P. Wei* (HGUP 18252), living culture GUCC 18252 = CGMCC 3.20557; Guiyang City, Huaxi District, N26°42'55" E106°68'07", 1140 m a.s.l., isolated from soil, 6 May 2018, *T.P. Wei* (HGUP 18253), living culture GUCC 18253 = CGMCC 3.20558; Shiqian County, Pingshan Township, N27°40'50" E108°07'30", 1100 m a.s.l., isolated from soil, 2 Nov. 2019, *T.P. Wei* (HGUP 18254), living culture GUCC 18254 = CGMCC 3.20559.

Notes — Scolecobasidium tshawytschae was originally isolated as the etiologic agent of kidney mycosis in chinook salmon (Oncorhynchus tshawytscha) smolts (Doty & Slater 1946). The four strains (GUCC 18251 to GUCC 18254) obtained in the present study and many strains that have been reported so far originate in soil and plant litter (Barron & Busch 1962, Hamayun et al. 2009, Samerpitak et al. 2014), indicating that this species can also be saprophytic, corroborating the supposition that its infective ability in fish is purely opportunistic.

Verruconis Samerpitak et al., Fungal Diversity 65: 117. 2013 '2014'

Type species. Verruconis gallopava (W.B. Cooke) Samerpitak & de Hoog.

Notes - Verruconis was established by Samerpitak et al. (2014) to accommodate thermophilic species separated from Ochroconis (O. calidifluminalis and O. gallopava) and Scolecobasidium (S. verruculosum), with the type species, V. gallopava being an opportunistic neurotropic pathogen (Salkin et al. 1990, Seyedmousavi et al. 2013, Wang et al. 2018, Samerpitak et al. 2019). Therefore, thermophilicity and unbranched conidia were the main characteristics distinguishing this genus from Ochroconis. Subsequently, Zhang et al. (2018) and Qiao et al. (2019) successively placed the mesophilic species V. panacis, V. hainanensis and V. pseudotricladiata with Y- or T-shaped and cylindrical conidia under Verruconis. They found that Verruconis is not limited to thermophilic species with clavate to cylindrical conidia (Hernández-Restrepo et al. 2020). However, the addition of these species blurred the major distinguishing feature between Verruconis and Ochroconis. In contrast, the molecular systematics has played an important role in the taxonomy of these two genera (Machouart et al. 2014, Huanraluek et al. 2019, Shen et al. 2020). Samerpitak et al. (2016) revealed that the concatenated dataset of SSU, ITS, LSU, act1, tub2 and tef1 served as a reference for genus and species delimitations of Ochroconis and Verruconis (Lackner et al. 2014, Al-Hatmi et al. 2016). It is noteworthy that V. mangrovei is the first reported sexual species in Verruconis (Hyde et al. 2020).

Verruconis cylindricalis T.P. Wei & Y.L. Jiang, sp. nov. — Myco-Bank MB 840928; Fig. 18

Etymology. The epithet refers to its cylindrical chlamydospores.

Typus. CHINA, Guizhou Province, Shiqian County, Fodingshan National Nature Reserve, N27°40'49" E108°07'35", 1100 m a.s.l., from forest humus, 2 Nov. 2019, *T.P. Wei* (holotype HGUP 18299, isotype CGMCC 3.20573, culture ex-type GUCC 18299).

Mycelium consisting of branched, subhyaline or pale brown, smooth, 1–3 µm thick hyphae, frequently forming hyphal coils. *Conidiophores* solitary, straight or flexuous, subcylindrical, pale brown, sparsely septate, $(21-)27-57.5(-87.5) \times 2.5-$ 3(-3.5) µm (av. ± SD = 44.3 ± 22.2 × 2.6 ± 0.3 µm, n = 30). *Conidiogenous cells* integrated, terminal, sympodial, denticulate, bearing 1–2 conidium at the apex, $(8-)9.5-16(-17.5) \times 2.5-3(-3.5)$ µm (av. ± SD = $12.0 \pm 3.6 \times 2.5 \pm 0.4$ µm, n = 30). *Conidia* sparse on PDA and MEA, broadly ellipsoidal with prominent hila, 1-septate, minutely echinulate, brown to olivaceous brown, slightly constricted at the septum, $(7-)7.5-12 \times 4-5$ µm



Fig. 17 Scolecobasidium tshawytschae (GUCC 18254). a–c. Colony on PDA, OA and MEA; d–h. conidial apparatus with rhexolytic conidia; i. conidiophores reduced to conidiogenous cells; j–l. maturation process of conidia; m. chlamydospores. — Scale bars: d–m = 10 µm.



Fig. 18 *Verruconis cylindricalis* (culture ex-type GUCC 18299). a–c. Colony on PDA, OA and MEA; d–g, j–k, m. conidial apparatus with rhexolytic conidia, produced from sympodial conidiogenous cells; h–i. chlamydospores; l. hyphal coil. — Scale bars: h-i, $I = 20 \ \mu m$, all others = 10 $\ \mu m$.



Fig. 19 *Verruconis thailandica* (GUCC 18267). a–c. Colony on PDA, OA and MEA; d–e. conidiophores reduced to conidiogenous cells; f–l. hyphae, conidiophores with sympodially proliferating conidiogenous cells and ellipsoidal and rhexolytic succession conidia. — Scale bars: d–l = 10 µm.

(av. \pm SD = 9.4 \pm 1.2 \times 4.3 \pm 0.2 μ m, n = 30). *Chlamydospores* intercalary, verruculose, cylindrical or clavate, slightly tapered at both ends, 2-septate, often asymmetric with smaller middle cells, olivaceous brown, usually constricted at the septum, 25–29.5 \times 4.5–6 μ m (av. \pm SD = 26.9 \pm 2.2 \times 5.1 \pm 0.6 μ m, n = 30).

Culture characteristics — Colonies on PDA attaining 17– 20 mm diam after 14 d at 26 °C, brown, margin irregular, submerged mycelium. On OA reaching up to 23–25 mm diam, velvety, dark brown, woolly at centre. On MEA reaching 20–23 mm diam, hairy, olivaceous brown. On PDA and MEA reverse blood colour with diffuse blood pigment spreading into agar.

Notes — *Verruconis cylindricalis* shares a few morphological similarities with *V. thailandica* and *V. calidifluminalis* in having two-celled conidia with protuberant hila. However, *V. thailandica* produces smaller, verrucose conidia ($5-7 \times 2.2-3.1 \mu m vs 7-12 \times 4-5 \mu m$), with a wing-like gelatinous brown sheath (Hernández-Restrepo et al. 2020). The conidia of the *V. calidifluminalis* are cylindrical to clavate, pale to dark brown and larger ($9.5-20.5 \times 2.5-5.0 \mu m vs 7-12 \times 4-5 \mu m$) (Yarita et al. 2010). Additionally, *V. cylindricalis* has longer conidiophores, cylindrical or clavate and 2-septate chlamydospores. These characters form the most notable differences with respect to *V. thailandica* and *V. calidifluminalis*. Phylogenetically, *V. cylindricalis* clustered in a distinct clade with full support from three independent algorithms (Fig. 2).

Verruconis tricladiata (Matsush.) T.P. Wei & Y.L. Jiang, comb. nov. — MycoBank MB 842456

Basionym. Scolecobasidium tricladiatum Matsush., Microfungi Solomon Isl. Papua-New Guinea: 52. 1971.

Description — Matsushima (1971).

Notes — Scolecobasidium tricladiatum was introduced based on its Y- or T-shaped or ellipsoidal to fusoid conidia, a species previously isolated from rotten leaves in Papua New Guinea (Matsushima 1971). Although we did not locate the type specimen, we have examined the DNA sequence data of another culture lodged under this species name in GenBank. Multi-locus phylogenetic analysis showed it to cluster in Verruconis, as sister to V. pseudotricladiata (Fig. 2). Nevertheless, S. tricladiatum and V. pseudotricladiata are genetically distinct in 6 bp (1 %), 87 bp (10 %) and 85 bp (9 %) in SSU, LSU and tef1 loci. No ITS, act1 and tub2 data are currently available for S. tricladiatum. Morphologically, S. tricladiatum differs from V. pseudotricladiata by its moniliform, irregularly branched conidiophores and mostly unbranched, pale olivaceous to brown, verruculose conidia (Matsushima 1971, Qiao et al. 2019). Based on morphological characters and phylogenetic analyses, we therefore transferred S. tricladiatum to Verruconis.

Verruconis thailandica Giraldo López & Crous, Fung. Syst. Evol. 6: 21. 2020 — Fig. 19

Mycelium composed of branched, septate, pale brown, smooth, thin-walled, 2–2.5 µm diam hyphae. *Conidiophores* erect, arising directly from vegetative hyphae, sometimes reduced to conidiogenous cells, simple or branched, pale brown, septate, subcylindrical, straight or slightly curved, $(7.5-)9.5-45(-52.5) \times 3-4 \mu m$ (av. $\pm SD = 19.3 \pm 10.9 \times 3.2 \pm 0.3 \mu m$, n = 30). *Conidiogenous cells* terminal or intercalary, flask-shaped to clavate, $5-13.5(-14) \times 3-4 \mu m$ (av. $\pm SD = 7.5 \pm 2.2 \times 3.3 \pm 0.3 \mu m$, n = 30), producing conidia sympodially on long open denticles; denticles cylindrical, pale brown, up to 1 µm long. *Conidia* solitary, broadly ellipsoidal with a protuberant hilum, strongly constricted at the septum, 1-septate, brown, finely echinulate to verrucose, sometimes with a wing-like gelatinous brown sheath,

released by rhexolytic secession, $7-11(-11.5) \times 3.5-5 \ \mu m$ (av. ± SD = 8.4 ± 1.0 × 3.9 ± 0.3 μm , n = 30).

Culture characteristics — Colonies on PDA attaining 16– 18 mm diam after 14 d at 26 °C, felty, growing slowly, margin irregular, grey olivaceous. On OA reaching up to 17–18 mm diam, cottony to floccose, immersed, dark brown. On MEA reaching 24–26 mm diam, grey olivaceous, aerial mycelium moderate. On PDA and MEA with ochreous diffusible pigment.

Material examined. CHINA, Guizhou Province, Guiyang City, Huaxi Wetland Park, N26°43'92" E106°67'76", 1140 m a.s.l., isolated from the humus soil in the stream, 16 Nov. 2020, *T.P. Wei* (HGUP 18267), living culture GUCC 18267 = CGMCC 3.20572.

Notes — Multi-locus phylogenetic analyses indicate that our newly obtained isolates GUCC 18267 clustered together with *V. thailandica*, and were sister to *V. terricola*, *V. verruculosa* and *V. cylindricalis*. Morphologically, *V. thailandica* can be readily distinguished from other members of *Verruconis* by its two-celled ellipsoidal conidia with a wing-like gelatinous sheath (Hernández-Restrepo et al. 2020).

Matsushimaea Subramanian, Kavaka 5: 96. 1977 '1978'

Type species. Matsushimaea fasciculata (Matsush.) Subramanian.

Notes — Matsushimaea was introduced by Subramanian (1977) to accommodate species segregated from *Torula (T. fasciculata)*, which are characterised by the production of sessile, branched and aseptate conidia from polyblastic sympodial conidiogenous cells. This genus was formerly placed in the *Pezizomycotina* as *incertae sedis* (Castañeda-Ruiz et al. 1996, Matsushima 1996). Later, Crous et al. (2018b) using the rDNA (ITS and LSU) sequence data elucidated the phylogenetic position of *Matsushimaea* and placed it in *Sympoventuriaceae*. Presently, *Matsushimaea* includes four species, *M. fasciculata*, *M. fertilis*, *M. magna* and *M. monilioides* (Castañeda-Ruiz et al. 1996, Matsushima 1996, Crous et al. 2018b). Among them, *M. fertilis* and *M. magna* lack authentic cultures and DNA sequence data, thus their phylogenetic position remains unknown.

Matsushimaea fasciculata (Matsush.) Subram., Kavaka 5: 96. 1977 — Fig. 20

Basionym. Torula fasciculata Matsush., Icon. Microfung. Matsush. Lect. (Kobe): 153. 1975.

Mycelium superficial or immersed, hyphae branched, septate, pale brown, smooth, occasionally with irregular swellings, $1.5-2 \mu m$ wide. *Conidiophores* reduced to conidiogenous cells arising directly from vegetative hyphae. *Conidiogenous cells* integrated, terminal or intercalary, mono- or polyblastic, pale brown, cylindrical to inflated, smooth-walled, $4-5.5 \times 4-5 \mu m$ (av. $\pm SD = 4.4 \pm 0.4 \times 4.1 \pm 0.3 \mu m$, n = 30). *Conidia* catenate, usually formed in branched chains, straight to sometimes curved, up to $(9.5-)14-50.5(-53) \mu m$ (av. $\pm SD = 27.5 \pm 9.5 \mu m$, n = 30) long; cells subglobose or ellipsoidal to somewhat pyriform, sessile or on short protrusions, aseptate, smooth, pale to dark brown, $3.5-5 \times 3.5-5 \mu m$ (av. $\pm SD = 3.9 \pm 0.4 \times 4.1 \pm 0.4 \mu m$, n = 30).

Culture characteristics — Colonies on PDA attaining 14– 15 mm diam after 14 d at 26 °C, velvety, olivaceous at centre, dark brown at periphery, margin entire. On OA reaching up to 15–19 mm diam, dark brown, dusty, flat, with sparse aerial mycelium. On MEA reaching 15–16 mm diam, olivaceous grey, floccose to loosely cottony.

Material examined. CHINA, Guizhou Province, Guiyang City, Tianhetan Tourist Holiday Resort, N26°43'95" E106°57'64", 1164 m a.s.l., from soil, 16 Apr. 2018, *T.P. Wei* (HGUP 18239), living culture GUCC 18239 = CGMCC 3.20544.



Fig. 20 Matsushimaea fasciculata (GUCC 18239). a–c. Colony on PDA, OA and MEA; d–l. conidia in simple or branched chains arising from conidiogenous cells; m–o. olivaceous to plate brown, subglobose or pyriform and smooth-walled conidia. — Scale bars: d, n = 20 μ m, all others = 10 μ m.



Fig. 21 *Mycosisymbrium cirrhosum* (GUCC 1837). a–c. Colony on PDA, OA and MEA; d–e. filiform appendages terminating from conidiophores; f–i. conidia stained with lactic acid phenol cotton blue; j–l. conidiophore aggregates with conidia. — Scale bars: $d-e = 20 \mu m$, all others = 10 μm .

Notes — In this study, the phylogenetic analysis is partly consistent with the morphological comparison, and our isolate GUCC 18239 and *M. fasciculata* have basically the same morphological characteristics (Fig. 1). All members of *Matsushimaea* have been isolated from forest litter and soil, which indicates that the lifestyle of this genus is probably saprophytic (Fig. 3). Moreover, this second report of this poorly known taxon extends its distribution to southwest China from its original location in Japan (Matsushima 1975).

Mycosisymbrium Carris, Mycologia 86: 132. 1994

Type species. Mycosisymbrium cirrhosum Carris.

Notes — The monotypic genus *Mycosisymbrium* was first described by Carris (1994), based on *M. cirrhosum* collected from dead leaves of *Vaccinium* macrocarpon in Massachusetts. It is characterised by discrete aggregates of conidiophores terminating in sterile, filiform appendages and brown, 1-septate conidia. Initially, this genus was treated as *incertae sedis* in *Pezizomycotina*. Pratibha & Prabhugaonkar (2016) confirmed the phylogenetic placement of *Mycosisymbrium*, which is a well-supported sister genus to *Ochroconis* and *Verruconis* in *Sympoventuriaceae*. It is noteworthy that since *Mycosisymbrium* was described, only Pratibha & Prabhugaonkar (2016) have reported on this species.

Mycosisymbrium cirrhosum Carris, Mycologia 86: 132. 1994 — Fig. 21

Mycelium consisting of brown, septate, branched, smooth, thick-walled, $1.5-2 \mu m$ diam hyphae. *Conidiophores* in determinate clusters, discrete, infundibuliform, pale to dark brown, smooth, branched, each branch terminating in a filiform appendage, $(21.5-)24-50(-57.5) \times 3-4 \mu m$ (av. \pm SD = $37.9 \pm 9.3 \times 3.2 \pm 0.3 \mu m$, n = 30); appendages hyaline, flexuous, up to $(29-)47-85.5(-90.5) \mu m$ (av. \pm SD = $63.4 \pm 16.9 \mu m$, n = 30) long. *Conidiogenous cells* terminal or intercalary, mono- to polyblastic, cylindrical, pale brown, with one to two denticle-like conidiogenous loci inconspicuous to slightly prominent, $(4.5-)5.5-10.5(-11) \times (2-)2.5-3.5(-4) \mu m$ (av. \pm SD = $7.8 \pm 1.2 \times 2.6 \pm 0.3 \mu m$, n = 30). *Conidia* solitary, oblong, 1-septate, smooth-walled, with bluntly rounded ends, constricted at median septum, $9-12 \times (3.5-)4-5 \mu m$ (av. \pm SD = $9.8 \pm 0.8 \times 4.0 \pm 0.3 \mu m$, n = 30).

Culture characteristics — Colonies on PDA attaining 24– 27 mm diam after 14 d at 26 °C, margin effuse, brown, with moderate aerial mycelium. On OA reaching up to 16–25 mm diam, mycelium immersed, dark brown, felty or granulose. On MEA reaching 23–24 mm diam, woolly to loosely cottony, olivaceous grey.

Material examined. CHINA, Guizhou Province, Meitan County, N27°41'08" E107°25'41", 910 m a.s.l., isolated from decaying *Camellia sinensis* leaf litter, 10 Aug. 2019, *T.P. Wei* (HGUP 1837), living culture GUCC 1837 = CGMCC 3.20541.

Notes — Multi-locus phylogenetic analyses indicate that our isolate GUCC 1837 clusters with *M. cirrhosum* with high statistical support. Furthermore, the two strains are morphologically similar (Fig. 21). This is the third report of *M. cirrhosum*, which was previously isolated in Massachusetts, USA (Carris 1994) and Goa, India (Pratibha & Prabhugaonkar 2016). In our study we extended its distribution to southwest China and added another new host record.

DISCUSSION

The controversy of Ochroconis, Scolecobasidium and Verruconis

Ochroconis, Scolecobasidium and Verruconis are morphologically and phylogenetically similar. Historically, there is some disagreement and incongruence about the phylogenetic placement, circumscription and classification of these three genera (De Hoog & Von Arx 1973, Ren et al. 2013, Giraldo et al. 2014, Samerpitak et al. 2014). Although several taxonomic revisions of these genera have been made based on morphology and phylogeny, their species boundaries have not been completely resolved due to historical confusion and limited molecular data (Samerpitak et al. 2016, Qiao et al. 2019, Shen et al. 2020). In this study, our multi-locus phylogenetic analyses indicated that Scolecobasidium and Ochroconis are synonymous and sister to Verruconis, and reside in Sympoventuriaceae (Fig. 1, 2). Morphologically, Verruconis and Scolecobasidium are distinguished mainly based on their conidial shape; in Verruconis the conidia are pale to dark brown, verrucose to coarsely ornamented, with protuberant hila, while in Scolecobasidium the conidial shape is more variable from ellipsoidal, cylindrical, bilobate, to T- or Y-shaped, and conidia are smooth-walled to verruculose. Additionally, the dark brown diffuse red colony pigmentation on PDA of Verruconis also distinguishes this genus from Scolecobasidium, which tends to have a pale luteus pigment. The chlamydospores seem to be another taxonomically important feature, in Scolecobasidium they are spherical or subcylindrical, smooth-walled, but in Verruconis they are cylindrical or clavate, verruculose and larger. Furthermore, Verruconis includes thermophilic and mesophilic species, while Scolecobasidium only has mesophilic species.

The species boundaries drawn in this study are mainly based on the multi-locus phylogeny as well as morphology. Compared to previous studies, more samples and additional gene markers were used to provide a better understanding of the phylogenetic relationships among species of Scolecobasidium and Verruconis. By comparing morphological characteristics and related DNA sequence data, seven new species were proposed in Scolecobasidium and Verruconis, namely S. camellicola, S. coiledmyces, S. echinulatum, S. obovoideum, S. verrucaria, S. zunyiense and V. cylindricalis. We also proposed six new combinations in Scolecobasidium and Verruconis as S. ferulica. S. guangxiensis, S. helicteris, S. leishanicola, S. mirabilis and V. tricladiata. The number of species in Scolecobasidium and Verruconis has increased significantly over the years (Crous et al. 2019c, Shen et al. 2020). Scolecobasidium and Verruconis are rather common genera of saprotrophic soil hyphomycetes, some of which are opportunistic neurotropic pathogens of humans, fish or other animals, and some are also known for their thermophilic properties (Samerpitak et al. 2014, 2019). Scolecobasidium is the largest genus of Sympoventuriaceae with 88 accepted species names recorded in MycoBank (http:// www.mycobank.org, April 2022), 39 of which are devoid of DNA sequence data in GenBank. Therefore, to resolve the phylogenetic position of these species in Scolecobasidium, they need to be re-collected, sequenced and epitypified.

Genera of Sympoventuriaceae

Sympoventuriaceae was originally established for three groups, namely Sympoventuria, Veronaeopsis and fusicladium-like species (Zhang et al. 2011). However, as discussed by Zhang et al. (2011), the fusicladium-like morphs are polyphyletic and include species residing in another different family, Venturiaceae. Machouart et al. (2014) transferred Scolecobasidium (= Ochroconis) and Verruconis to the Sympoventuriaceae, thereby expanding the concept of the family. Subsequent phylogenetic studies further expanded the concept of Sympoventuriaceae, making it the largest family of Venturiales (Johnston & Park 2016, Tibpromma et al. 2018, Crous et al. 2019a, Shen et al. 2020). Although these advances have allowed the resolution of several long-standing questions concerning the generic boundaries of Sympoventuriaceae, many questions remain unresolved about the phylogenetic relationships of some taxa, especially genera and species for which molecular data are not yet available. To better define the generic boundaries and reveal the evolutionary relationship of Sympoventuriaceae, we carried out a more comprehensive analysis of this group based on a hitherto most complete sequence dataset consisting of seven loci (SSU, ITS, LSU, act1, tub2, tef1 and rpb2). The present results resolved 22 well-supported monophyletic lineages, representing 22 genera (Fig. 1), viz., Acroconidiellina, Bellamyces, Clavatispora, Echinocatena, Fuscohilum, Guizhoumyces, Helicopsis, Matsushimaea, Melnikomyces, Mycosisymbrium, Neocoleroa, Neofusicladium, Parafusicladium, Pinaceicola, Pseudosigmoidea, Scolecobasidium, Sterila, Sympoventuria, Troposporella, Veronaeopsis, Verruconis and Yunnanomyces. These included fungi with a broad spectrum of morphology, lifestyles and modes of nutrition, accommodating saprophytes, endophytes, plant pathogens, and animal or human opportunistic pathogens (Samerpitak et al. 2014, 2019, Wang et al. 2018, Crous et al. 2020).

The number of known taxa in Sympoventuriaceae is increasing at a steady pace as more geographic areas and habitats are investigated, and its taxonomy has also changed dramatically (Wijayawardene et al. 2014, Tibpromma et al. 2018, Shen et al. 2020). In this study, in addition to the 15 genera previously placed in Sympoventuriaceae (Table 4), we also reassessed species of Acroconidiellina, Clavatispora, Guizhoumyces, Matsushimaea, Melnikomyces, Mycosisymbrium and Yunnanomyces. The morphological characters and molecular data of these genera provide significant evidence for their taxonomic placement in Sympoventuriaceae. In the multi-locus phylogenetic tree of the current study, A. arecae is allied with Scolecobasidium at the top of the clade (Fig. 1). Nevertheless, A. arecae can be distinguished morphologically from the species in Scolecobasidium by its multi-septate, obclavate, large conidia with a truncate base and macronematous conidiophores (Li et al. 2016). The placement of Acroconidiellina in Sympoventuriaceae contradicts earlier placements in the order Pleosporales due to similarities in the type of conidiophores and conidium morphology (Ellis 1971, Zhang et al. 2009, Bhat 2010). Therefore, our results are more in line with the proposal of Hernández-Restrepo et al. (2016) about a closer affinity with members of the Sympoventuriaceae. Furthermore, our findings support the placement of Mycosisymbrium in Sympoventuriaceae, as suggested by previous molecular studies (Pratibha & Prabhugaonkar 2016). The extended taxon sampling and the use of five markers allow us to strongly corroborate these findings. Species of two genera were included in the analyses, Mycosisymbrium and Clavatispora, which formed two different sister subclades (Fig. 1). Mycosisymbrium can be distinguished by having discrete aggregates of conidiophores terminating in sterile, filiform appendages and 1-septate conidia (Fig. 21), while Clavatispora has 1(-3)-septate, guttulate conidia that are deeply constricted at their septa (Boonmee et al. 2014). Clavatispora also produced a sexual morph in culture characterised by ostiolate ascomata, bitunicate asci and muriformly septate ascospores.

According to the multigene phylogeny and morphology in the present study, a new genus, *Guizhoumyces*, is now recognised within *Sympoventuriaceae* (Fig. 1). Morphologically, members of *Guizhoumyces* are quite distinct from those of *Sympoventuriaceae*, having straight or curved, smooth and acicular to

obclavate conidia, which contrasts with the mostly verrucose to denticulate and subcylindrical to fusoid-ellipsoidal conidia typical for species of other genera; as well as by the presence of anastomosis between mature conidia in Guizhoumyces. Moreover, Matsushimaea forms a robust clade with another two genera, Fuscohilum and Neocoleroa (Fig. 1). As we mentioned before, our results confirm its placement within Sympoventuriaceae, although the DNA sequences of M. fertilis and M. magna were not available. The polyblastic and sympodial conidiogenous cells of several other genera in Sympoventuriaceae share similar morphologies with those of Matsushimaea. However, Matsushimaea differs in having sessile, branched and aseptate conidia arising directly from vegetative hyphae (Crous et al. 2018b; Fig. 20), which strongly supports the genus as monophyletic. Based on the general characteristics of Melnikomyces, previous studies hypothesized its close relationship with Scolecobasidiella and Scolecobasidium (Crous et al. 2014, Hernández-Restrepo et al. 2020). In our phylogenetic tree, Melnikomyces formed a separate clade in Sympoventuriaceae distinct from other genera (Fig. 1). Morphologically, this genus has certain similarities with Scolecobasidiella and Scolecobasidium, but can be distinguished from them by its fusoid-ellipsoidal conidia and the chlamydospores that are subglobose, occurring in branched chains (Wei et al. 2020). Thus, our analysis shows that the phylogenetic isolation of Melnikomyces, Scolecobasidiella and Scolecobasidium may be supported by the unique morphological features and genetic differences. Yunnanomyces is characterised by its globose to broadly oval, yellow to brown and muriformly septate conidia (Tibpromma et al. 2018). This genus is phylogenetically related to the monotypic genus Sterila, and formed a well-supported clade within Sympoventuriaceae (Fig. 1). Morphologically it is not possible to compare these two genera, as the latter is sterile in culture. Nonetheless, Yunnanomyces differs from its closest phylogenetic neighbour Sterila by unique fixed alleles in four loci of their type species, by 6 bp in ITS (3 %), 52 bp in LSU (7%), 278 bp in tub2 (57%) and 191 bp in rpb2 (24%). More specifically, these two genera represent significantly different

Sympoventuria currently includes three species, namely S. africana, S. capensis and S. melaleucae, with the sexual species S. capensis designated as the generic type (Crous et al. 2007a, 2017). Sympoventuria is phylogenetically closely related to the asexual genus Helicopsis with high support from three independent algorithms (Fig. 1). Sympoventuria is, however, clearly distinguished from other asexual morphs by its fusoid-ellipsoid or cylindrical, simple or branched conidial chains (helically coiled conidia with thick conidial filaments in Helicopsis) (Karsten 1888, Crous et al. 2007b, Tsui & Berbee 2010). Furthermore, Neocoleroa metrosideri, a little-known sexual morph is here shown to be closely related to the sterile N. cameroonensis. However, N. metrosideri differs from its closest phylogenetic neighbour N. cameroonensis by unique fixed alleles in two loci, by 56 bp in ITS (14 %) and 26 bp in LSU (3 %). Notably, species with sexual morphs were scattered throughout the phylogenetic tree, which indicates that sexual reproduction may have evolved more than once within the family (Fig. 1). Overall, the results of this study have provided a robust overview of the species boundaries in Sympoventuriaceae. This is based largely on seven gene regions inferring an updated phylogram of all concerned genera in the family. On the other hand, multi-gene phylogenetic analyses combined with morphological features provide a robust means to delimit fungal species boundaries (Woudenberg et al. 2017, Lücking et al. 2020, Crous et al. 2021). Results of this study revealed that SSU, ITS, LSU, act1, tub2, tef1 and rpb2 gene regions can provide stable and reliable resolution for species delimitation

lineages in Sympoventuriaceae.

in *Sympoventuriaceae*. Although each of these loci proved to be suitable barcoding markers for species identification, a combined analysis is highly recommended.

Evolution of lifestyles in Sympoventuriaceae

The transition from saprophytes to endophytes, plant pathogens and animal or human opportunistic pathogens is a notable feature within Sympoventuriaceae (Fig. 3). Studies have shown that the saprotrophic fungal ancestors experienced a largescale loss of plant cell wall degrading enzymes, and obtained effector-like secreted proteins to fit a plant-fungal associated lifestyle (Bödeker et al. 2014, Kohler et al. 2015, Martin et al. 2016, Haridas et al. 2020, Shen et al. 2020, Benavent 2021). In our reconstruction analyses, although the Sympoventuriaceae clade most conspicuously includes saprophytes, members of the Neocoleroa and Sterila clades convergently evolved towards plant pathogens and animal/human opportunistic pathogens, with a shift towards endophytes in Verruconis (Fig. 3). In addition, we found the Scolecobasidium clade to have a high species richness, including saprophytes, endophytes, animal/human opportunistic and plant pathogens, and to be associated with significant increasing shifts in diversification rate (Fig. 3). It is worth noting that transition from saprophytes to plant pathogens and animal or human opportunistic pathogens has independently occurred multiple times during the evolution of Sympoventuriaceae, which is possibly driven by habitat selection (Fig. 3). The present study has provided a wealth of data about the lifestyle and phylogeny of Sympoventuriaceae, showing some trends in the evolution of species in the family. For example, saprophytes were mainly at the early diverged clades in phylogenetic trees, e.g., Neofusicladium, Parafusicladium, Pseudosigmoidea and Veronaeopsis. In contrast, the endophytes, plant pathogens and animal/human opportunistic pathogens were only found in the Acroconidiellina, Neocoleroa, Scolecobasidium, Sterila and Verruconis species in more recently diverged clades, supporting a strong correlation between the evolution of Sympoventuriaceae life strategies and its phylogeny.

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REFERENCES

- Abbott EV. 1927. Scolecobasidium, a new genus of soil fungi. Mycologia 19: 29–31.
- Al-Hatmi AMS, Gerrits van den Ende AHG, Stielow JB, et al. 2016. Evaluation of two novel barcodes for species recognition of opportunistic pathogens in Fusarium. Fungal Biology 120: 231–245.
- Ando K, Nakamura N. 2000. Pseudosigmoidea: a new genus for a hyphomycete (ATCC 16660) formerly identified as Sigmoidea prolifera. The Journal of General and Applied Microbiology 46: 51–57.
- Arzanlou M, Groenewald JZ, Gams W, et al. 2007. Phylogenetic and morphotaxonomic revision of Ramichloridium and allied genera. <u>Studies in</u> Mycology 58: 57–93.
- Baldacci E, Ciferri R. 1937. Un nuovo genere di micete parassita del pioppo Pollaccia radiosa (Lib.) Baldacci e Ciferri, Revisione dei G. Stigmella e Stigmina. I. Pollaccia radiosa (Lib.) Baldacci e Ciferri. Atti dell'Istituto Botanico della Universita e Laboratorio Crittogamico di Pavia 10: 55–72.
- Barido-Sottani J, Bošková V, Plessis LD, et al. 2018. Taming the BEAST A community teaching material resource for BEAST 2. Systematic Biology 67: 170–174.
- Barr ME. 1997. Notes on some 'Dimeriaceous' fungi. Mycotaxon 64: 149–171. Barron GL, Busch LV. 1962. Studies on the soil hyphomycete Scolecobasidium. Canadian Journal of Botany 40: 77–84.
- Beck A, Ritschel A, Schubert K, et al. 2005. Phylogenetic relationships of the anamorphic genus Fusicladium s. lat. as inferred by ITS nrDNA data. Mycological Progress 4: 111–116.

- Benavent IG. 2021. Macro-and microevolutionary perspectives on Seynesiella juniperi, a fungus in the Venturiales (Dothideomycetes, Ascomycota). Botanica complutensis 45: 115–126.
- Bhat DJ. 2010. Fascinating microfungi (hyphomycetes) of Western Ghats, India. Department of Botany, Goa University.
- Bödeker IT, Clemmensen KE, De Boer W, et al. 2014. Ectomycorrhizal Cortinarius species participate in enzymatic oxidation of humus in northern forest ecosystems. New Phytologist 203: 245–256.
- Bonorden HF. 1851. Handbuch der allgemeinen Mykologie. Schweizerbart'sche Verlagshandlung, Stuttgart.
- Boonmee S, Bhat DJ, Maharachchikumbura SS, et al. 2014. Clavatispora thailandica gen. et sp. nov., a novel taxon of Venturiales (Dothideomycetes) from Thailand. Phytotaxa 176: 92–101.
- Carbone I, Kohn L. 1999. A method for designing primer sets for speciation studies in filamentous ascomycetes. Mycologia 91: 553–556.
- Carris LM. 1994. Vaccinium fungi: Mycosisymbrium cirrhosum gen. et sp. nov. Mycologia 86: 31–133.
- Castañeda-Ruiz RF, Guarro J, Cano J. 1996. Notes on conidia fungi. Two new dematiaceous hyphomycetes from Cuba. Mycotaxon 57: 463–469.
- Chen Y, Xie L, Long Y, et al. 2020. A new species and two new Chinese records of Ochroconis from sugarcane and banana rhizosphere in Guangxi, China. Mycoscience 61: 307–314.
- Crane JL. 1968. Freshwater hyphomycetes of the northern Appalachian Highland including New England, and three coastal plain states. American Journal of Botany 55: 996–1002.
- Crous PW, Carlier J, Roussel V, et al. 2020. Pseudocercospora and allied genera associated with leaf spots of banana (Musa spp.). Fungal Systematics and Evolution 7: 1–19.
- Crous PW, Lombard L, Sandoval-Denis M, et al. 2021. Fusarium: more than a node or a foot-shaped basal cell. Studies in Mycology 98: 100116.
- Crous PW, Mohammed C, Glen M, et al. 2007a. Eucalyptus microfungi known from culture. 3. Eucasphaeria and Sympoventuria genera nova, and new species of Furcaspora, Harknessia, Heteroconium and Phacidiella. Fungal Diversity 25: 19–36.
- Crous PW, Schubert K, Braun U, et al. 2007b. Opportunistic, human-pathogenic species in the Herpotrichiellaceae are phenotypically similar to saprobic or phytopathogenic species in the Venturiaceae. Studies in Mycology 58: 185–217.
- Crous PW, Schumacher RK, Akulov A, et al. 2019a. New and interesting fungi. 2. Fungal Systematics and Evolution 3: 57–134.
- Crous PW, Schumacher RK, Wingfield MJ, et al. 2018a. New and interesting fungi. 1. Fungal Systematics and Evolution 1: 169–215.
- Crous PW, Shivas RG, Quaedvlieg W, et al. 2014. Fungal Planet description sheets: 214–280. Persoonia 32: 184–306.
- Crous PW, Verkley GJM, Groenewald JZ, et al. 2019b. Fungal Biodiversity. [Westerdijk Laboratory Manual Series no.1.] Utrecht, Westerdijk Fungal Biodiversity Institute, Utrecht, the Netherlands.
- Crous PW, Wingfield MJ, Burgess TI, et al. 2017. Fungal Planet description sheets: 625–715. Persoonia 39: 270–467.
- Crous PW, Wingfield MJ, Burgess TI, et al. 2018b. Fungal Planet description sheets: 716–784. Persoonia 40: 239–392.
- Crous PW, Wingfield MJ, Le Roux JJ, et al. 2015. Fungal Planet description sheets: 371–399. Persoonia 35: 264–327.
- Crous PW, Wingfield MJ, Lombard L, et al. 2019c. Fungal Planet description sheets: 951–1041. Persoonia 43: 223–425.
- Crous PW, Wingfield MJ, Richardson DM, et al. 2016. Fungal Planet description sheets: 400–468. Persoonia 36: 316–458.
- Darriba D, Taboada GL, Doallo R, et al. 2012. jModelTest 2: more models, new heuristics and parallel computing. Nature Methods 9: 772–772.
- De Hoog GS, Von Arx JA. 1973. Revision of Scolecobasidium and Pleurophragmium. Kavaka 1: 55–60.
- Diene O, Wang W, Narisawa K. 2013. Pseudosigmoidea ibarakiensis sp. nov., a dark septate endophytic fungus from a cedar forest in Ibaraki, Japan. Microbes and Environments 28: 381–387.
- Doty MS, Slater DW. 1946. A new species of Heterobasidium tshawytschae pathogenic on young chinook salmon. American Midland Naturalist 36: 663–665.
- Ellis MB. 1971. Dematiaceous Hyphomycetes. Commonwealth Mycological Institute, Kew, Surrey.
- Ellis MB. 1976. More Dematiaceous Hyphomycetes. Commonwealth Mycological Institute, Kew, England.
- Fan XL, Bezerra JDP, Tian CM, et al. 2020. Cytospora (Diaporthales) in China. Persoonia 45: 1–45.
- Gams W. 2015. An ex-type culture cannot always tell the ultimate truth. IMA Fungus 6: A69.
- Gargas A, Taylor JW. 1992. Polymerase chain reaction (PCR) primers for amplifying and sequencing 18S rDNA from lichenized fungi. Mycologia 84: 589–592.

Giraldo A, Crous PW. 2019. Inside plectosphaerellaceae. Studies in Mycology 92: 227–286.

- Giraldo A, Sutton DA, Samerpitak K, et al. 2014. Occurrence of Ochroconis and Verruconis species in clinical specimens from the United States. Journal of Clinical Microbiology 52: 4189–4201.
- Glass NL, Donaldson GC. 1995. Development of primer sets designed for use with the PCR to amplify conserved genes from filamentous ascomycetes. Applied & Environmental Microbiology 61: 1323–1330.
- Graniti A. 1963. Scolecobasidium anellii nsp. agente di annerimenti superficiali di stalattiti (Scolecobasidium anellii n. sp. causing a superficial darkening of stalacitites). Nuovo Giornale Botanico Italiano 69: 360–365.
- Hall TA. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium Series 41: 95–98.
- Hamayun M, Khan SA, Kim HY, et al. 2009. Gibberellin production and plant growth enhancement by newly isolated strain of Scolecobasidium tshawytschae. Journal of Microbiology and Biotechnology 19: 560–565.
- Hao L, Chen C, Zhang R, et al. 2013. A new species of Scolecobasidium associated with the sooty blotch and flyspeck complex on banana from China. Mycological Progress 12: 489–495.
- Haridas S, Albert R, Binder M, et al. 2020. 101 Dothideomycetes genomes: a test case for predicting lifestyles and emergence of pathogens. <u>Studies</u> in Mycology 96: 141–153.
- Hernández-Restrepo M, Giraldo A, Van Doorn R, et al. 2020. The genera of fungi – G6: Arthrographis, Kramasamuha, Melnikomyces, Thysanorea, and Verruconis. Fungal Systematics and Evolution 6: 1–24.
- Hernández-Restrepo M, Schumacher RK, Wingfield MJ, et al. 2016. Fungal Systematics and Evolution: FUSE 2. Sydowia 68: 193–230.
- Horré R, De Hoog GS, Kluczny C, et al. 1999. rDNA diversity and physiology of Ochroconis and Scolecobasidium species reported from humans and other vertebrates. Studies in Mycology 43: 194–205.
- Huanraluek N, Phukhamsakda C, Senwanna C, et al. 2019. Verruconis heveae, a novel species from Hevea brasiliensis in Thailand. Phytotaxa 403: 47–54.
- Hyde KD, Dong Y, Phookamsak R, et al. 2020. Fungal diversity notes 1151– 1276: taxonomic and phylogenetic contributions on genera and species of fungal taxa. Fungal Diversity 100: 5–277.
- Hyde KD, Jones EBG, Liu JK, et al. 2013. Families of Dothideomycetes. Fungal Diversity 63: 1–313.
- Jayasiri SC, Hyde KD, Jones EBG, et al. 2019. Diversity, morphology and molecular phylogeny of Dothideomycetes on decaying wild seed pods and fruits. Mycosphere 10: 1–186.
- Johnston PR, Park D. 2016. Neocoleroa metrosideri sp. nov. (Sympoventuriaceae, Venturiales). Phytotaxa 253: 214–218.
- Karsten PA. 1888. Diagnoses fungorum nonnullorum novorum, in Fennia detectorum. Revue Mycologique Toulouse 10: 73–75.
- Kidd S, Halliday C, Alexiou H, et al. 2016. Descriptions of medical fungi (3rd ed.). Newstyle Printing, Adelaide, Australia.
- Kirk PM, Cannon PF, Minter DW, et al. 2008. Ainsworth & Bisby's dictionary of the fungi. CAB International, Wallingford.
- Kohler A, Kuo A, Nagy LG, et al. 2015. Convergent losses of decay mechanisms and rapid turnover of symbiosis genes in mycorrhizal mutualists. Nature Genetics 47: 410–415.
- Koorders SH. 1907. Botanische Untersuchungen. Verhandelingen Koninklijke Nederlandse Akademie van Wetenschappen Afdeling Natuurkunde 13: 1–264.
- Koukol O. 2010. Revision of "Septonema ochraceum" revealed three new species of Venturiaceae and Herpotrichiellaceae. Mycological Progress 9: 369–378.
- Kumar S, Stecher G, Li M, et al. 2018. MEGA X: Molecular Evolutionary Genetics Analysis across Computing Platforms. Molecular Biology and Evolution 35: 1547–1549.
- Lackner M, De Hoog GS, Yang L, et al. 2014. Proposed nomenclature for Pseudallescheria, Scedosporium and related genera. Fungal Diversity 67: 1–10.
- Larsson A. 2014. AliView: a fast and lightweight alignment viewer and editor for large data sets. Bioinformatics 30: 3276–3278.
- Li JF, Bhat DJ, Phookamsak R, et al. 2016. Sporidesmioides thailandica gen. et sp. nov. (Dothideomycetes) from northern Thailand. Mycological Progress 15: 1169–1178.
- Liu F, Ma ZY, Hou LW, et al. 2022. Updating species diversity of Colletotrichum, with a phylogenomic overview. Studies in Mycology 101: 1–56.
- Liu JK, Hyde KD, Jeewon R, et al. 2017. Ranking higher taxa using divergence times: a case study in Dothideomycetes. Fungal Diversity 84: 75–99.
- Liu YJ, Whelen S, Hall BD. 1999. Phylogenetic relationships among ascomycetes: evidence from an RNA polymerase II subunit. <u>Molecular Biology</u> and Evolution 16: 1799–1808.

- Lücking R, Aime MC, Robbertse B, et al. 2020. Unambiguous identification of fungi: where do we stand and how accurate and precise is fungal DNA barcoding. IMA Fungus 11: 1–32.
- Machouart M, Samerpitak K, De Hoog GS, et al. 2014. A multigene phylogeny reveals that Ochroconis belongs to the family Sympoventuriaceae (Venturiales, Dothideomycetes). Fungal Diversity 65: 77–88.
- Martin F, Kohler A, Murat C, et al. 2016. Unearthing the roots of ectomycorrhizal symbioses. Nature Reviews Microbiology 14: 760–773.
- Martin-Sanchez PM, Nováková A, Bastian F, et al. 2012. Two new species of the genus Ochroconis, O. lascauxensis and O. anomala isolated from black stains in Lascaux Cave, France. Fungal Biology 116: 574–589.
- Matsushima T. 1971. Microfungi of the Solomon Islands and Papua-New Guinea. Kobe, Japan.
- Matsushima T. 1975. Icones microfungorum: a Matsushima lectorum. Kobe, Japan.
- Matsushima T. 1980. Saprophytic microfungi from Taiwan part 1. Hyphomycetes. Matsushima Mycological Memoirs 1: 1–82.
- Matsushima T. 1996. Matsushima Mycological Memoirs 9: 1-30.
- Miller MA, Pfeiffer W, Schwartz T. 2012. Creating the CIPRES Science Gateway for inference of large phylogenetic trees. Proceedings of the Gateway Computing Environments Workshop (GCE). New Orleans, LA. CA, San Diego Supercomputer Center: 1–8.
- Murata K, Ogawa Y, Kusama K, et al. 2022. Disseminated Verruconis gallopava infection in a patient with systemic lupus erythematosus in Japan: A case report, literature review, and autopsy case. Medical Mycology Case Reports 35: 35–38.
- Nylander JAA. 2004. MrModeltest v2.2. Program distributed by the author: 2. Evolutionary Biology Centre, Uppsala University.
- Papendorf MC. 1969. New South African soil fungi. Transactions of the British Mycological Society 52: 483–489.
- Petrak F. 1934. Mykologische Beiträge zur Flora von Sibirien. Hedwigia 74: 30–78.
- Pratibha J, Prabhugaonkar A. 2016. Distribution and phylogeny of Mycosisymbrium cirrhosum. Mycosphere 7: 44–50.
- Punithalingam E, Spooner BM. 2011. A new fungicolous Scolecobasidium (hyphomycetes) and Caducirostrum gen. nov. (coelomyces) from leaf litter in the UK and Italy. Kew Bulletin 66: 309–324.
- Qiao M, Tian W, Castañeda-Ruiz RF, et al. 2019. Two new species of Verruconis from Hainan, China. MycoKeys 48: 41–53.
- Rambaut A, Drummond AJ, Xie D, et al. 2018. Posterior summarization in Bayesian phylogenetics using Tracer 1.7. Systematic Biology 67: 901.
- Rehner SA, Buckley E. 2005. A Beauveria phylogeny inferred from nuclear ITS and EF1-α sequences: evidence for cryptic diversification and links to Cordyceps teleomorphs. Mycologia 97: 84–98.
- Ren J, Jie CY, Zhou QX, et al. 2013. Molecular and morphological data reveal two new species of Scolecobasidium. Mycoscience 54: 420–425.
- Revankar SG, Sutton D. 2010. Melanized fungi in human disease. Clinical Microbiology Reviews 23: 884–928.
- Ronquist F, Teslenko M, Van der Mark P, et al. 2012. MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. Systematic Biology 61: 539–542.
- Roy RY, Dwivedi RS, Mishra RR. 1962. Two new species of Scolecobasidium from soil. Lloydia 25: 164–166.
- Rozewicki J, Li S, Amada KM, et al. 2019. MAFFT-DASH: integrated protein sequence and structural alignment. Nucleic Acids Research 47: 5–10.
- Salkin IF, Dixon DM, Kemna ME, et al. 1990. Fatal encephalitis caused by Dactylaria constricta var. gallopava in a snowy owl chick (Nectea scandiaca). Journal of Clinical Microbiology 28: 2845–2847.
- Samerpitak K, Alfjorden A, Seyedmousavi S, et al. 2019. Ochroconis globalis infecting Atlantic salmon (Salmo salar), with a review of Ochroconis species in cold-blooded animals. Journal of Fish Diseases 42: 947–957.
- Samerpitak K, Duarte APM, Attili-Angelis D, et al. 2015a. A new species of the oligotrophic genus Ochroconis (Sympoventuriaceae). Mycological Progress 14: 1–10.
- Samerpitak K, Gerrits van den Ende BH, Stielow JB, et al. 2016. Barcoding and species recognition of opportunistic pathogens in Ochroconis and Verruconis. Fungal Biology 120: 219–230.
- Samerpitak K, Gerrits van den Ende AH, Menken SBJ, et al. 2015b. Three new species of the genus Ochroconis. Mycopathologia 180: 7–17.
- Samerpitak K, Gloyna K, De Hoog GS. 2017. A novel species of the oligotrophic genus Ochroconis colonizing indoor wet cells. Mycoscience 58: 290–296.
- Samerpitak K, Van der Linde E, Choi HJ, et al. 2014. Taxonomy of Ochroconis genus including opportunistic pathogens on humans and animals. Fungal Diversity 65: 89–126.
- Satow MM, Attili-Angelis D, De Hoog GS, et al. 2008. Selective factors involved in oil flotation isolation of black yeast from the environment. Studies in Mycology 61: 157–163.

Schubert K, Ritschel A, Braun U. 2003. A monograph of Fusicladium s. lat. (Hyphomycetes). Schlechtendalia 9: 1–132.

- Seifert KA, Morgan-Jones G, Gams W, et al. 2011. The genera of hyphomycetes. CBS-KNAW Fungal Biodiversity Centre, Utrecht, the Netherlands.
- Seyedmousavi S, Guillot J, De Hoog GS. 2013. Phaeohyphomycoses, emerging opportunistic diseases in animals. Clinical Microbiology Reviews 26: 19–35.
- Seyedmousavi S, Samerpitak K, Rijs AJMM, et al. 2014. Antifungal susceptibility patterns of opportunistic fungi in the genera Verruconis and Ochroconis. Antimicrobial Agents and Chemotherapy 58: 3285–3292.
- Shen M, Zhang JQ, Zhao LL, et al. 2020. Venturiales. Studies in Mycology 96: 185–308.
- Singh A, Singh NK, Singh PN, et al. 2019. Additions to Ochroconis from India. Phytotaxa 427: 186–199.
- Sivanesan A. 1977. The taxonomy and pathology of Venturia species. Lubrecht & Cramer Ltd, Vaduz.
- Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30: 1312–1313.
- Subramanian CV. 1977. Revisions of Hyphomycetes I. Kavaka 5: 93-98.
- Swofford DL. 2003. PAUP* 4.0b10. Phylogenetic Analysis Using Parsimony (*and other methods). Version 4. Sinauer Associates, Sunderland, MA, USA.
- Tazik Z, Rahnama K, Iranshahi M, et al. 2020. Ochroconis ferulica sp. nov. (Venturiales), a fungal endophyte from Ferula ovina. Nova Hedwigia 110: 369–381.
- Tibpromma S, Hyde KD, McKenzie EH, et al. 2018. Fungal diversity notes 840–928: micro-fungi associated with Pandanaceae. Fungal Diversity 92: 1–160.
- Tsui CKM, Berbee ML. 2010. Transfer of two Helicoma species to Troposporella based on molecular and morphological data. Mycoscience 51: 144–148.
- Vilgalys R, Hester M. 1990. Rapid genetic identification and mapping of enzymatically amplified ribosomal DNA from several Cryptococcus species. Journal of Bacteriology 172: 4238–4246.
- Wang X, Cai W, Gerrits van den Ende AH, et al. 2018. Indoor wet cells as a habitat for melanized fungi, opportunistic pathogens on humans and other vertebrates. Scientific Reports 8: 1–10.

- Wei TP, Zhang X, Ren PP, et al. 2020. A novel Melnikomyces species from forest litter in China. Phytotaxa 471: 61–68.
- White TJ, Bruns T, Lee S, et al. 1990. Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. In: Innis MA, Gelfand DH, Sninsky JJ, et al. (eds), PCR Protocols: a guide to methods and applications: 315–322. Academic Press, San Diego, California, USA.
- Wijayawardene NN, Crous PW, Kirk PM, et al. 2014. Naming and outline of Dothideomycetes – 2014 including proposals for the protection or suppression of generic names. Fungal Diversity 69: 1–55.
- Wijayawardene NN, Hyde KD, Al-Ani LKT, et al. 2020. Outline of fungi and fungus-like taxa. Mycosphere 11: 1060–1456.
- Wijayawardene NN, Hyde KD, Lumbsch HT, et al. 2018. Outline of ascomycota: 2017. Fungal Diversity 88: 167–263.
- Woudenberg JHC, Hanse B, Van Leeuwen GCM, et al. 2017. Stemphylium revisited. Studies in Mycology 87: 77–103.
- Yarita K, Sano A, Samerpitak K, et al. 2010. Ochroconis calidifluminalis, a sibling of the neurotropic pathogen O. gallopava, isolated from hot spring. Mycopathologia 170: 21–30.
- Yu Y, Blair C, He XJ. 2020. RASP 4: Ancestral State Reconstruction Tool for Multiple Genes and Characters. Molecular Biology and Evolution 37: 604–606.
- Yu Y, Harris AJ, Blair C, et al. 2015. RASP (Reconstruct Ancestral State in Phylogenies): a tool for historical biogeography. Molecular Phylogenetics and Evolution 87: 46–49.
- Zhang JQ, Dou ZP, Zhou YP, et al. 2016. Venturia chinensis sp. nov. a new venturialean ascomycete from Khingan Mountains. Saudi Journal of Biological Sciences 23: 592–597.
- Zhang TY, Yu Y, Zhang MY, et al. 2018. Verruconis panacis sp. nov., an endophyte isolated from Panax notoginseng. International Journal of Systematic and Evolutionary Microbiology 68: 2499–2503.
- Zhang X, Wang KY, Ren PP, et al. 2020. Ochroconis terricola sp. nov. from China. Mycotaxon 135: 143–150.
- Zhang Y, Crous PW, Schoch CL, et al. 2011. A molecular, morphological and ecological re-appraisal of Venturiales a new order of Dothideomycetes. Fungal Diversity 51: 249–277.
- Zhang Y, Schoch CL, Fournier J, et al. 2009. Multi-locus phylogeny of Pleosporales: a taxonomic, ecological and evolutionary re-evaluation. Studies in Mycology 64: 85–102.