

Research Article

Taxonomic novelties and global biogeography of *Montagnula* (Ascomycota, Didymosphaeriaceae)

Dhanushka N. Wanasinghe¹⁰, Thilina S. Nimalrathna^{2,3,4,50}, Li Qin Xian¹⁰, Turki Kh. Faraj⁶⁰, Jianchu Xu^{1,70}, Peter E. Mortimer¹⁰

- 1 Honghe Center for Mountain Futures, Kunming Institute of Botany, Chinese Academy of Sciences, Honghe County 654400, Yunnan, China
- 2 CAS Key Laboratory of Tropical Forest Ecology, Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, Menglun, Mengla, Yunnan, China
- 3 Southeast Asia Biodiversity Research Institute, Chinese Academy of Sciences & Center for Integrative Conservation, Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, Mengla, Yunnan 666303, China
- 4 Yunnan International Joint Laboratory of Southeast Asia Biodiversity Conservation & Yunnan Key Laboratory for Conservation of Tropical Rainforests and Asian Elephants, Menglun, Mengla, Yunnan 666303, China
- 5 International College, University of Chinese Academy of Sciences, Beijing, China
- 6 Department of Soil Science, College of Food and Agriculture Sciences, King Saud University, P.O. Box 145111, Riyadh 11362, Saudi Arabia
- 7 CIFOR-ICRAF China Country Program, Kunming, Yunnan, China

Corresponding authors: Jianchu Xu (jxu@mail.kib.ac.cn); Peter E. Mortimer (peter@mail.kib.ac.cn)

Abstract

Whilst conducting surveys of lignicolous microfungi in Yunnan Province, we collected a large number of taxa that resemble Montagnula (Didymosphaeriaceae, Pleosporales). Our phylogenetic study on Montagnula involved analysing sequence data from ribosomal RNA genes (nc18S, nc28S, ITS) and protein-coding genes (rpb2, tef1-q). We present a biphasic approach (morphological and molecular phylogenetic evidence) that supports the recognition of four new species in Montagnula viz., M. lijiangensis, M. menglaensis, M. shangrilana and M. thevetiae. The global diversity of Montagnula is also inferred from metabarcoding data and published records based on field observations. Metabarcoding data from GlobalFungi and field observations provided insights into the global diversity and distribution patterns of Montagnula. Studies conducted in Asia, Australia, Europe, and North America revealed a concentration of Montagnula species, suggesting regional variations in ecological preferences and distribution. Montagnula species were found on various substrates, with sediments yielding a high number of sequences. Poaceae emerged as a significant contributor, indicating a potential association between Montagnula species and grasses. Culture-based investigations from previously published data revealed Montagnula species associations with 105 plant genera (in 45 plant families), across 55 countries, highlighting their wide ecological range and adaptability. This study enhances our understanding of the taxonomy, distribution, and ecological preferences of Montagnula species. It emphasizes their role in the decomposition of organic matter in grasslands and savannah systems and suggests further investigation into their functional roles in ecosystem processes. The global distribution patterns and ecological interactions of Montagnula species underscore the need for continued research and conservation efforts.

Key words: Global distribution, microfungi, molecular phylogeny, taxonomy, Yunnan

Academic editor: Nalin Wijayawardene Received: 25 September 2023 Accepted: 17 December 2023 Published: 19 January 2024

Citation: Wanasinghe DN, Nimalrathna TS, Qin Xian L, Faraj TK, Xu J, Mortimer PE (2024) Taxonomic novelties and global biogeography of *Montagnula* (Ascomycota, Didymosphaeriaceae). MycoKeys 101: 191–232. https://doi.org/10.3897/ mycokeys.101.113259

Copyright: © Dhanushka N. Wanasinghe et al. This is an open access article distributed under terms of the Creative Commons Attribution License (Attribution 4.0 International – CC BY 4.0).

Introduction

Fungi are the second largest group of eukaryotes, performing vital ecological functions such as decomposition, mutualism, and pathogenesis to plants and animals (Tedersoo et al. 2014). Ascomycota, which forms the largest phylum of Fungi, and includes the genus Montagnula, is an incredibly diverse group, with an estimated global species richness of ~154,500 species (Bánki et al. 2023). Despite their ecological and economic importance, many Ascomycota species remain undescribed, and their distribution and diversity have yet to be properly determined (Maharachchikumbura et al. 2021a, b; Wijayawardene et al. 2022). This is somewhat due to the fact that many Ascomycota species are microscopic and inconspicuous, making them difficult to find and subsequently study, or sometimes these smaller species can be overlooked with studies focussing on more charismatic species of macrofungi (Wanasinghe et al. 2022a). The investigation of taxonomic and phylogenetic systematics in Ascomycota is bridging crucial knowledge gaps and enhancing our understanding of this particular group of fungi. Montagnula (typified with M. infernalis), is an example of a relatively understudied genus within Ascomycota, and many species remain undescribed. Understanding the taxonomic, phylogenetic and host relationships between Montagnula species will help us better understand how they have diversified and adapted to different habitats in various ecological zones. These data are useful to make predictions about the ecology and biology of the genus and to guide future research into their interactions with other organisms and their roles in ecosystem processes. Understanding the taxonomy and phylogeny of *Montagnula* is also important for conservation purposes. With ongoing habitat destruction and climate change, it is more important than ever to understand the current diversity and distribution of fungi around the world (Wanasinghe et al. 2022a).

Therefore, our research group at the Center for Mountain Futures (CMF), has been conducting investigations into the microfungal diversity and biogeography in Yunnan Province, Southwest China. Specifically, we are focusing on various substrates such as leaf and woody litter, aiming to clarify the taxonomy of fungi on these substrates, using morphology in conjunction with multigene phylogeny. As a result, we have successfully isolated numerous anamorphic and teleomorphic Ascomycota species in Yunnan, and we have published our findings based on different themes, including their relationship with hosts, substrates, and localities (Thiyagaraja et al. 2019, 2020, 2021; Abeywickrama et al. 2020; Wanasinghe et al. 2020, 2021, 2022b, 2023; Yasanthika et al. 2020; Bundhun et al. 2021; Dissanayake et al. 2021; Gao et al. 2021; Monkai et al. 2021; Mortimer et al. 2021; Ren et al. 2021a, b, 2022a, b; Aluthmuhandiram et al. 2022; Maharachchikumbura et al. 2022; Wanasinghe and Mortimer 2022). The objectives of this study are (1) to identify the lignicolous Montagnula species collected from Yunnan using both morphological and phylogenetic approaches, and (2) to utilize metabarcoding data and published records based on field observations to infer the global diversity and biogeography of Montagnula. The analyses conducted in this study revealed four new species and four existing species of Montagnula, in Yunnan. The discovery of several previously undescribed Ascomycota species in the genus Montagnula in Yunnan Province is a significant advancement in our understanding of the diversity and distribution of this group of fungi. Furthermore, the utilization of metabarcoding data and published records based on field observations to infer the global diversity of *Montagnula* demonstrates the potential of these approaches in elucidating the biogeography of fungi on a large scale. By studying and documenting the diversity of *Montagnula* species, we can enhance our appreciation for the importance of conserving these fungi and their habitats, and take appropriate measures to mitigate the threats they face.

Materials and methods

Sample collecting

Fresh fungal materials were collected from dead woody twigs from Honghe, Kunming, Mengla, Shangri-La and Yulong Counties, all within Yunnan Province, China, during the dry season (January, March, April) and wet season (August, September). To preserve their integrity, the specimens were transported to the laboratory in Zip lock plastic bags during the dry season and in paper bags during the wet season.

Morphological observations

The morphology of external and internal macro-/micro-structures were observed as described in Wanasinghe et al. (2017, 2018a, 2020). Hand sections of the ascomata were mounted in distilled water and the following characteristics were evaluated and measured: ascomata diameter, height, color and shape; width of peridium; and height and diameter of ostioles. Length and width (at the widest point) of asci and ascospores. Images were captured with a Canon EOS 600D digital camera fitted to a Nikon ECLIPSE Ni compound microscope. Macroscopic images of colonies were documented using an iPhone XS Max (Apple Inc., Cupertino, CA, USA) with daylight. Measurements were made with the Tarosoft (R) Image Frame Work program, and images used for figures were processed with Adobe Photoshop CS5 Extended version 10.0 software (Adobe Systems, San José, CA, USA).

Isolation

Single spore isolation was conducted by following the methods described in Wanasinghe et al. (2018b). Germinated spores were individually transferred to potato dextrose agar (PDA: 39 g/L distilled water, Difco potato dextrose) plates and grown at 20 °C in the daylight.

Deposition of specimens, cultures and registering names

The living cultures were deposited at the Kunming Institute of Botany Culture Collection (KUNCC), Kunming, China. Dry herbarium materials were deposited in the herbarium of Cryptogams Kunming Institute of Botany, Academia Sinica (KUN-HKAS). MycoBank numbers have been obtained as outlined in MycoBank (http://www.MycoBank.org accessed on 21 September 2023) for the novel taxa.

DNA extraction, PCR amplifications and sequencing

Genomic DNA was extracted from the axenic mycelium as described by Phookamsak et al. (2017). Mycelia for DNA extraction from each isolate were grown on PDA for 3–4 weeks at 20 °C and total genomic DNA was extracted from approximately 150 ± 50 mg axenic mycelium scraped from the edges of the growing culture. Mycelium was ground to a fine powder with liquid nitrogen and DNA extracted using the Biospin Fungus Genomic DNA Extraction Kit-BSC14S1 (BioFlux, P.R. China) following the instructions of the manufacturer. When fungi failed to grow in culture, DNA extraction was carried out directly from fruiting bodies, adhering to the protocol outlined by Wanasinghe et al. (2018b). DNA to be used as templates for Polymerase Chain Reaction (PCR) were stored at 4 °C for use in regular work and duplicated at -20 °C for long-term storage.

We used primers ITS5/ITS4 (White et al. 1990), LR0R/LR5 (Vilgalys and Hester 1990; Rehner and Samuels 1994), NS1/NS4 (White et al. 1990), EF1-983F/EF1-2218R (Liu et al. 1999; Rehner and Buckley 2005), and fRPB2-5f/fRPB2-7cR (Sung et al. 2007) to amplify sequence data for a total of five markers: the internal transcribed spacers (ITS), partial 28S large subunit rDNA (LSU), partial 18S small subunit rDNA (SSU), translation elongation factor 1- α (tef1- α), and RNA polymerase II second largest subunit (*rpb2*). PCR amplifications were performed following the methods described in Wanasinghe et al. (2021). We sequenced complementary strands with the same primers used for PCR amplifications and sequencing was done from a commercial sequencing provider (BGI, Ltd Shenzhen, P.R. China). The nucleotide sequence data obtained were deposited in GenBank (Table 2).

Sequencing assembly and alignments

Sequences generated from different primers of the five genes were analysed with other sequences retrieved from GenBank (Table 2). Sequences with high similarity indices were determined from a BLAST search to find the closest matches with taxa in Didymosphaeriaceae, using recently published data (Du et al. 2021; Ren et al. 2022a; Sun et al. 2023). The multiple alignments of all consensus sequences, as well as the reference sequences were automatically generated with MAFFT v. 7 (Katoh et al. 2019), and manually corrected where necessary using BioEdit v. 7.0.5.2 (Hall 1999).

Phylogenetic inference

The single-locus datasets were examined for topological incongruence among loci for members of the analyses. The alignments were concatenated into a multi-locus alignment that was analyzed with maximum likelihood (ML) and Bayesian (BI) phylogenetic methods in the CIPRES Science Gateway (Miller et al. 2010). ML tree was obtained using RAxML-HPC2 on XSEDE v. 8.2.10 (Stamatakis 2014) with applying GTR+G+I model. Support values were obtained with 1,000 bp replicates (Felsenstein 1985). ML bootstrap values equal or greater than 75% are given above each node. The best-fit model was selected with respect to Bayesian Information Criterion (BIC) scores using the IQ-TREE web application at http://iqtree.cibiv.univie.ac.at (Trifinopoulos et al. 2016). For model selection, we restricted the pool of available models to JC, F81, HKY, SYM and GTR (Ronquist et al. 2011). BI

were performed with two parallel runs of 2 M generations, using four chains in each, and retaining one tree every 100 generations. The dataset was partitioned by gene region, and a GTR + G + I model was applied to each partition, ending the run automatically when standard deviation of split frequencies dropped below 0.01 with a burn-in fraction of 0.25. A fifty percent majority rule consensus tree was obtained after discarding the first 25% of trees, and posterior probabilities were used as a measure of nodal support. The posterior probability in BI (BYPP) greater than 0.95 are given above each node. Phylograms were visualized with FigTree v1.4.0 program (Rambaut 2012) and reorganized in Microsoft power point (2019).

The biogeographical distribution of Montagnula

In our initial approach, we obtained detailed geographical distribution information for the Montagnula genus. This data was extracted from the GlobalFungi database (https://globalfungi.com, accessed on 04 December 2023), as outlined by Větrovský et al. (2020). The database provided information on the countries and precise geographical coordinates of recorded Montagnula occurrences. To visualize these occurrences, we employed a range of packages in R version 4.2.1 (R Core Team 2022), including 'sf' (Pebesma and Bivand 2023), 'raster' (Hijmans 2023), 'rgdal' (Bivand et al. 2022), and 'ggplot2' (Wickham 2011). In our map, each marker signifies an individual occurrence of Montagnula. These occurrences are visually distinguished by a color scheme, with each color denoting the specific biome from which the samples were collected, as illustrated in Fig. 2a. Additionally, we have developed two donut charts, showcased in Fig. 2b, c, which effectively illustrate the distribution of Montagnula sequences. These charts present the sequence abundance as a percentage of the total, segmented across various biomes and continents, providing a clear visual breakdown of their distribution. Furthermore, we have gathered Environmental DNA (eDNA) data from diverse sources in metabarcoding studies focusing on fungi, as found in the GlobalFungi database (Fig. 3). This dataset included specifics about eDNA sources, locations of the studies, and the sequence abundance of Montagnula sequences. It is important to note that the sequence abundance in metabarcoding studies might not always accurately represent the actual abundance of species in a habitat. Nonetheless, these data can provide valuable insights into the potential rarity or prevalence of the group in the eDNA source. We analyzed the sequence abundance in diverse eDNA samples from different continents. Before visualization, the abundance values were normalized via a logarithmic transformation to ensure a standardized and comparable presentation of Montagnula sequence abundance. Post-transformation abundance data were visualized using the 'ggplot2' package, aiding in highlighting the focus areas of metabarcoding and identifying the environmental sample types from which Montagnula sequences were derived across various continents (Figs 2, 3).

The host relations of Montagnula

To illustrate the host specificity of *Montagnula* species, we utilized detailed information regarding host species from the literature (Table 1). This enabled us to create informative bar plots displaying the host preferences of *Montagnula* species (Fig. 4). This information was visualized using the 'ggplot2' package in R.

Species	Host species	Host family	Country	Reference		
Montagnula acaciae	Acacia auriculiformis	Fabaceae Thailand		Tennakoon et al. (2022)#		
Montagnula aloes	Aloe sp.	Asphodelaceae South Africa		Crous et al. (2012)#		
Montagnula appendiculata	Zea mays	Poaceae China		Aptroot (2004)#		
Montagnula aquatica	Submerged wood	NA	Thailand	Sun et al. (2023)#		
	Dead woody litter	NA	China	This study#		
Montagnula aquilariae	Aquilaria sinensis	Thymelaeaceae	China	Hyde et al. (2023)#		
	Dead woody litter	NA	China	This study#		
Montagnula baatanensis	Agave sp.	Asparagaceae	USA	Crivelli (1983)		
Montagnula bellevaliae	Bellevalia romana	Asparagaceae	Italy	Hongsanan et al. (2015)#		
Montagnula camporesii	Dipsacus sp.	Caprifoliaceae	Italy	Hyde et al. (2020)#		
Montagnula camarae	Cytisus scoparius	Fabaceae	Portugal	Checa (2004)		
Montagnula chiangraiensis	Chromolaena odorata	Asteraceae	Thailand	Mapook et al. (2020)#		
Montagnula chromolaenae	Chromolaena odorata	Asteraceae	Thailand	Mapook et al. (2020)#		
Montagnula chromolaenicola	Chromolaena odorata	Asteraceae	Thailand	Mapook et al. (2020)#		
	Lagerstroemia sp.	Lythraceae	China	This study#		
Montagnula cirsii	Cirsium sp.	Asteraceae	Italy	Hyde et al. (2016)#		
Montagnula cylindrospora	Human skin##	NA	USA	Crous et al. (2020)#		
Montagnula dasylirionis	Dasylirion sp.	Asparagaceae	USA	Ramaley and Barr (1995)		
Montagnula donacina	Acacia reficiens	Fabaceae	Namibia	Aptroot (1995)		
	Acacia sp.	Fabaceae	India	Aptroot (1995)		
	Adhatoda vasica	Acanthaceae	India	Aptroot (1995)		
	Ailanthus altissima	Simaroubaceae	India	Aptroot (1995)		
	Althaea rosea	Malvaceae	China	Aptroot (1995)		
	Annona squamosa	Annonaceae	India	Aptroot (1995)		
	Arundo donax	Poaceae	Portugal	Aptroot (1995)		
	Bambusoideae	Poaceae	Brazil	Aptroot (1995)		
	Bambusoideae	Poaceae	Papua New Guinea	Aptroot (1995)		
	Cajanus cajan	Fabaceae	India	Aptroot (1995)		
	Calamus australis	Arecaceae	Australia	Hyde et al. (1999)		
	Careya arborea	Lecythidaceae	India	Aptroot (1995)		
	Citrus aurantiifolia	Rutaceae	India	Aptroot (1995)		
	Clerodendrum infortunatum	Lamiaceae	India	Aptroot (1995)		
	Clerodendrum multiflorum	Lamiaceae	India	Aptroot (1995)		
	Coffea arabica	Rubiaceae Paraguay		Aptroot (1995)		
	Coffea robusta	Rubiaceae Central African Republic		Aptroot (1995)		
	Craterellus odoratus##	Cantharellaceae China		Zhao et al. (2018)#		
	Duranta repens	Verbenaceae	India	Aptroot (1995)		
	Ficus glomerata	Moraceae	India	Aptroot (1995)		
	Funtumia africana	Apocynaceae	Sierra Leone	Aptroot (1995)		
	Hibiscus sp.	Malvaceae	India	Aptroot (1995)		
	lpomoea carnea	Convolvulaceae	India	Aptroot (1995)		
	Mallotus philippinensis	Euphorbiaceae	India	Aptroot (1995)		
	Morus alba	Moraceae	India	Aptroot (1995)		
	Litchi litchi	Sapindaceae	Myanmar	Thaung (2008)		

 Table 1. Accepted species in Montagnula including their host and geographic location.

Species	Host species	Host family	Country	Reference
Montagnula donacina	Nerium odorum	Apocynaceae	India	Aptroot (1995)
	Paeonia suffruticosa	Paeoniaceae	China	Li et al. (2023)#
	Phyllostachys bambusoides	Poaceae	Japan	Wang et al. (2004)
	Pistacia sp.	Anacardiaceae	India	Aptroot (1995)
	Platanus sp.	Platanaceae	USA	Wang et al. (2004)
	Premna cumingiana	Lamiaceae	Philippines	Aptroot (1995)
	Pseudosasa japonica	Poaceae	France	Aptroot (1995)
	Saccharum officinarum	Poaceae	Brazil	Aptroot (1995)
	Unknown stem	NA	India	Aptroot (1995)
	Tectona grandis	Lamiaceae	India	Aptroot (1995)
	Terminalia tomentosa	Combretaceae	India	Aptroot (1995)
	Trachycarpus fortunei	Arecaceae	China	Hyde et al. (1999)
	Unknown bark	NA	India	Aptroot (1995)
	Unknown branches	NA	Sierra Leone	Aptroot (1995)
	Unknown plant	NA	Colombia	Aptroot (1995)
	Dead wood	NA	China	Sun et al. (2023)#
	Dead wood	NA	Thailand	Ren et al. (2022a)#
	Dead wood	NA	China	This study#
	Vitis vinifera	Vitaceae	Australia	Pitt et al. (2014)#
	Wikstroemia sp.	Thymelaeaceae	USA	Aptroot (1995)
	Zea mays	Poaceae	Georgia	Aptroot (1995)
Montagnula dura	Aconitum septentrionale	Ranunculaceae	Sweden	Eriksson (1992)
	Lonicera etrusca	Caprifoliaceae	Spain	Checa (2004)
Montagnula gilletiana	Fraxinus ornus	Oleaceae	Bulgaria	Fakirova (2004)
	Retama sphaerocarpa	Fabaceae	Spain	Checa (2004)
	Ulex europaeus	Fabaceae	Spain	Checa (2004)
Montagnula graminicola	Poaceae	Poaceae	Italy	Liu et al. (2015)#
Montagnula guiyangensis	Helwingia himalaica	Helwingiaceae China		Sun et al. (2023) [#]
Montagnula hirtula	Cerastium latifolium	Caryophyllaceae	Austria	Leuchtmann (1984)
	Cerastium sp.	Caryophyllaceae	Italy	Leuchtmann (1984)
	Epilobium parviflorum	Onagraceae	Switzerland	Leuchtmann (1984)
	Rubus idaeus	Rosaceae	Finland	Leuchtmann (1984)
	Rubus sp.	Rosaceae	Sweden	Eriksson (1992)
Montagnula infernalis	Agave americana	Asparagaceae	Portugal	Checa (2004)
	Agave americana	Asparagaceae	Spain	Checa (2004)
	Fourcroya sp.	Asparagaceae	Portugal	Ariyawansa et al. (2014)
	Furcraea gigantea	Asparagaceae	Portugal	Checa (2004)
	Furcraea gigantea	Asparagaceae	Spain	Checa (2004)
	Furcraea longaeva	Asparagaceae	Portugal	Checa (2004)
	Furcraea longaeva	Asparagaceae	Spain	Checa (2004)
Montagnula infernalis	Furcraea macrophylla	Asparagaceae	Bahamas	Barr (1990)
Montagnula jonesii	Fagus sylvatica	Fagaceae	Italy	Tennakoon et al. (2016)#
	Ficus benjamina	Moraceae	Thailand	Tennakoon et al. (2022)#
Montagnula krabiensis	Pandanus sp.	Pandanaceae	Thailand	Tibpromma et al. (2018)#
Montagnula lijiangensis	Quercus sp.	Fagaceae	China	This study [#]
Montagnula longipes	Agave americana	Asparagaceae	Algeria	Aptroot (1995)
Montagnula melanorhabdos	Agave sp.	Asparagaceae	Turkey	Aptroot (2006)

Species	Host species	Host family	Country	Reference	
Montagnula menglaensis	Indocalamus tessellatus	Poaceae	China	This study#	
Montagnula mohavensis	Yucca mohavensis	Asparagaceae USA		Ramaley and Barr (1995)	
Montagnula obtusa	llex sp.	Aquifoliaceae	USA	French (1989)	
	Juglans sp.	Juglandaceae	USA	French (1989)	
	Pinus pinaster	Pinaceae	Portugal	Checa (2004)	
	Sorbus aucuparia	Rosaceae	Sweden	Eriksson (1992)	
Montagnula opaca	Phalaris	Poaceae Switzerland		Crivelli (1983)	
Montagnula opulenta	Ammophila arenaria	Poaceae	France	Aptroot (1995)	
	Ammophila arenaria	Poaceae	Germany	Aptroot (1995)	
	Ammophila arenaria	Poaceae	Sweden	Aptroot (1995)	
	Festuca brachyphylla	Poaceae	Canada	Aptroot (1995)	
	Opuntia ficus-indica	Cactaceae	Canary Islands	Aptroot (1995)	
	Opuntia ficus-indica	Cactaceae	France	Aptroot (1995)	
	Opuntia ficus-indica	Cactaceae	Italy	Aptroot (1995)	
	Opuntia ficus-indica	Cactaceae	Malta	Aptroot (1995)	
	Opuntia ficus-indica	Cactaceae	Tunisia	Aptroot (1995)	
	Opuntia sp.	Cactaceae	Cyprus	Aptroot (1995)	
	<i>Opuntia</i> sp.	Cactaceae	Israel	Aptroot (1995)	
	<i>Opuntia</i> sp.	Cactaceae	Italy	Aptroot (1995)	
	<i>Opuntia</i> sp.	Cactaceae	Tunisia	Aptroot (1995)	
	Opuntia tuna	Cactaceae	USA	Aptroot (1995)	
	Poa abbreviata	Poaceae	Canada	Aptroot (1995)	
	Puccinellia angustata	Poaceae	Greenland	Aptroot (1995)	
	Stipa himalaica	Poaceae	India	Aptroot (1995)	
Montagnula opuntiae	Opuntia lindheimeri	Cactaceae	USA	Huhndorf (1992)	
Montagnula palmacea	Chamaerops humilis	Arecaceae	France	Aptroot (1995)	
	Cocos capitata	Arecaceae	Spain	Aptroot (1995)	
	Daviesia nudiflora	Fabaceae	Australia	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Egypt	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Greece	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Iraq	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Italy	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Pakistan	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Saudi Arabia	Aptroot (1995)	
	Phoenix dactylifera	Arecaceae	Tunisia	Aptroot (1995)	
	Phoenix sylvestris	Arecaceae Pakistan		Aptroot (1995)	
	Pitcairnia chrysantha	Bromeliaceae	Chile	Aptroot (1995)	
	Unknown leaves	NA	USA	Aptroot (1995)	
	Unknown petiole	NA	USA	Aptroot (1995)	
Montagnula perforans	Calamagrostis arenaria	Poaceae	France	Aptroot (2006)	
Montagnula phragmospora	Agave americana	Asparagaceae	Portugal	Checa (2004)	
	Agave americana	Asparagaceae	Spain	Checa (2004)	
	Agave hookeri	Asparagaceae	Portugal	Checa (2004)	
	Agave hookeri	Asparagaceae	Spain	Checa (2004)	
	Agave sp.	Asparagaceae	France	Barr (1990)	
	Agave sp.	Asparagaceae	Portugal	Checa (2004)	
	Agave sp.	Asparagaceae	Spain	Checa (2004)	

Species	Host species	Host family	Country	Reference	
Montagnula phragmospora	Yucca brevifolia	Asparagaceae	California	Barr (1990)	
	Yucca sp.	Asparagaceae	Portugal	Checa (2004)	
	Yucca sp.	Asparagaceae	Spain	Checa (2004)	
Montagnula puerensis	Dead wood	NA	China	Du et al. (2021)#	
Montagnula rhodophaea	Arundo donax	Poaceae	Italy	Leuchtmann (1984)	
	Phragmites communis	Poaceae	Switzerland	Leuchtmann (1984)	
Montagnula saikhuensis	Citrus sp.	Rutaceae	Thailand	Wanasinghe et al. (2016)#	
Montagnula scabiosae	Scabiosa sp.	Caprifoliaceae Italy		Hongsanan et al. (2015)#	
Montagnula shangrilana	Rhododendron sp.	Ericaceae China		This study#	
Montagnula sp.	Carex fuliginosa	Cyperaceae Austria		Scheuer (1988)	
Montagnula spartii	Aeluropus littoralis	Poaceae	Russia	Aptroot (1995)	
	Ammophila arenaria	Poaceae	Belgium	Aptroot (1995)	
	Ammophila arenaria	Poaceae	Denmark	Aptroot (1995)	
	Ammophila arenaria	Poaceae	Sweden	Aptroot (1995)	
	Ammophila arenaria	Poaceae	United Kingdom	Aptroot (1995)	
	Calamagrostis epigeios	Poaceae	Russia	Aptroot (1995)	
	Calycotome spinosa	Fabaceae	France	Aptroot (1995)	
	Calycotome spinosa	Fabaceae	Spain	Aptroot (1995)	
	Calycotome villosa	Fabaceae	Italy	Aptroot (1995)	
	Carex rostrata	Cyperaceae	Sweden	Aptroot (1995)	
	Chamaerops humilis	Arecaceae	Spain	Aptroot (1995)	
	Leymus arenarius	Poaceae	Russia	Aptroot (1995)	
	Ephedra ciliata	Ephedraceae	Unknown country in Asia	Aptroot (1995)	
	Ephedra sp.	Ephedraceae	Iran	Aptroot (1995)	
	Festuca arenaria	Poaceae	France	Aptroot (1995)	
	Festuca sulcata	Poaceae	Iran	Aptroot (1995)	
	Genista aspalathoides	Fabaceae	Italy	Aptroot (1995)	
	Gramineae	Gramineae	Austria	Aptroot (1995)	
	Koeleria cristata	Poaceae	Germany	Aptroot (1995)	
	Koeleria glauca	Poaceae	Denmark	Aptroot (1995)	
	Linum austriacum	Linaceae	Germany	Aptroot (1995)	
	Luzula spadicea	Juncaceae	Switzerland	Aptroot (1995)	
	Lygeum spartum	Poaceae	Spain	Aptroot (1995)	
	Melica ciliata	Poaceae	France	Aptroot (1995)	
	Nardus stricta	Poaceae	Austria	Aptroot (1995)	
	Puccinellia peisonis	Poaceae	Austria	Aptroot (1995)	
	Sarothamnus scoparius	Fabaceae	Poland	Mulenko et al. (2008)	
	Sarothamnus scoparius	Fabaceae	Switzerland	Aptroot (1995)	
	Sesleria caerulea	Poaceae	Italy	Aptroot (1995)	
Montagnula spartii	Spartium junceum	Fabaceae	Albania	Aptroot (1995)	
	Spartium junceum	Fabaceae	France	Aptroot (1995)	
	Spartium junceum	Fabaceae	Greece	Aptroot (1995)	
	Spartium junceum	Fabaceae	Turkey	Aptroot (1995)	
	Ulex sp.	Fabaceae	Spain	Aptroot (1995)	
Montagnula spinosella	Abelia triflora	Caprifoliaceae	Spain	Checa (2004)	
	Carex aterrima	Cyperaceae	Austria	Scheuer (1988)	

Species	Host species	Host family	Country	Reference	
Montagnula spinosella	Carex misandra	Cyperaceae	Norway	Holm and Holm (1993, 1994)	
	Colpodium vahlianum	Poaceae	Norway	Holm and Holm (1993, 1994)	
	Deschampsia caespitosa	Poaceae	Norway	Holm and Holm (1993, 1994)	
	Juncus maritimus	Juncaceae	Spain	Holm and Holm (1993), Checa (2004)	
	Luzula confusa	Juncaceae Norway		Holm and Holm (1993, 1994)	
Montagnula stromatosa	Phoenix hanceana	Arecaceae	China	Lu et al. (2000)	
	Phoenix sp.	Arecaceae	China	Zhuang (2001)	
	Trachycarpus fortunei	Arecaceae	China	Hyde et al. (1999)	
	Trachycarpus fortunei	Arecaceae	United Kingdom	Hyde et al. (1999)	
Montagnula subsuperficialis	Panicum grumosum	Poaceae	Argentina	Shoemaker (1989)	
Montagnula thailandica	Chromolaena odorata	Asteraceae	Thailand	Mapook et al. (2020)#	
	Hevea brasiliensis	Euphorbiaceae	Thailand	Senwanna et al. (2021)#	
	Coffea arabica var. catimor	Rubiaceae	China	Lu et al. (2022) [#]	
	Unidentified twig	NA	Thailand	Boonmee et al. (2021)#	
Montagnula thevetiae	Thevetia peruviana	Apocynaceae	China	This study#	
Montagnula thuemeniana	Yucca sp.	Asparagaceae	USA	Barr (1990)	
Montagnula triseti	Trisetum distichophyllum	Poaceae	Switzerland	Crivelli (1983)	
Montagnula vakrabeejae	Unidentified twig	NA	Andaman	Niranjan and Sarma (2018)	
Montagnula verniciae	Vernicia fordii	Euphorbiaceae	China	Li et al. (2023)#	
Montagnula yuccigena Yucca baccata		Asparagaceae	Asparagaceae Mexico Ramaley and		

"#" Denotes molecular data available in GenBank. "##" Denotes none plant based. NA represents not applicable.

Table 2. GenBank accession numbers of sequences used for the phylogenetic analyses.

Taxon	Strain number	GenBank accession numbers					Deferrer
		ITS	LSU	SSU	tef1-a	rpb2	Reference
Montagnula acaciae	MFLUCC 18-1636	ON117280	ON117298	ON117267	ON158093	NA	Tennakoon et al. (2022)
	NCYUCC 19-0087 [™]	ON117281	ON117299	ON117268	ON158094	NA	Tennakoon et al. (2022)
Montagnula aloes	CPC 19671	JX069863	JX069847	NA	NA	NA	Crous et al. (2012)
	CBS 132531 [⊤]	NR_111757	NG_042676	NA	NA	NA	Crous et al. (2012)
Montagnula appendiculata	CBS 109027 [⊤]	DQ435529	AY772016	NA	NA	NA	Wanasinghe et al. (2016)
Montagnula aquatica	MFLU 22-0171 [⊤]	OP605992	OP605986	OP600504	NA	NA	Sun et al. (2023)
Montagnula aquatica	KUNCC 23-14425	OR583097	OR583116	OR583135	OR588088	OR588107	This study
	KUNCC 23-14557	OR583099	OR583118	OR583137	OR588090	OR588109	This study
Montagnula aquilariae	KUNCC 22-10815 [⊤]	0P452927	OP482265	OP482268	OP426318	NA	Hyde et al. (2023)
	KUNCC 22-10816	OP554219	OP482266	OP482269	OP426319	NA	Hyde et al. (2023)
	KUNCC 22-10815 [™]	0P452927	OP482265	OP482268	OP426318	NA	Hyde et al. (2023)
	KUNCC 22-10816	0P554219	OP482266	OP482269	OP426319	NA	Hyde et al. (2023)
Montagnula aquilariae	KUNCC 23-14430	OR583100	OR583119	OR583138	OR588091	OR588110	This study
	KUNCC 23-14431	OR583101	OR583120	OR583139	OR588092	OR588111	This study
	KUNCC 23-14432	OR583102	OR583121	OR583140	OR588093	OR588112	This study
Montagnula bellevaliae	MFLUCC 14-0924 ^T	KT443906	KT443902	KT443904	NA	NA	Hongsanan et al. (2015)
Montagnula camporesii	MFLUCC 16-1369 ^T	MN401746	NG_070946	NG_068418	MN397908	MN397909	Hyde et al. (2020)
Montagnula chiangraiensis	MFLUCC 17-1420 ^T	NR_168864	NG_068707	NG_070155	NA	NA	Mapook et al. (2020)
Montagnula chromolaenae	MFLUCC 17-1435 ^T	NR_168865	NG_068708	NG_070156	NA	NA	Mapook et al. (2020)

-	o: · · · ·	GenBank accession numbers					
Taxon	Strain number	ITS	LSU	SSU	tef1-a	rpb2	Keterence
Montagnula chromolaenicola	MFLUCC 17-1469 ^T	NR_168866	NG_070948	NG_070157	MT235773	MT235809	Mapook et al. (2020)
Montagnula chromolaenicola	KUNCC 23-14426	OR583098	OR583117	OR583136	OR588089	OR588108	This study
	KUNCC 23-14427	OR583103	OR583122	OR583141	OR588094	OR588113	This study
	KUNCC 23-14558	OR583104	OR583123	OR583142	OR588095	OR588114	This study
Montagnula cirsii	MFLUCC 13-0680	KX274242	KX274249	KX274255	KX284707	NA	Hyde et al. (2016)
Montagnula cylindrospora	CBS 146572 [⊤]	LT796834	LN907351	NA	LT797074	LT796994	Crous et al. (2020)
Montagnula donacina	HFG07004	MF967419	MF183940	NA	NA	NA	Zhao et al. (2017)
	HVVV01	KJ628375	KJ628377	KJ628376	NA	NA	Pitt et al. (2014)
	HKAS 124552	OP605991	OP605987	NA	NA	NA	Sun et al. (2023)
	KUMCC 21-0653	OP058961	OP059052	OP059003	OP135938	NA	Ren et al. (2021)
	KUMCC 21-0579	OP058963	OP059054	OP059005	OP135940	NA	Ren et al. (2021)
	KUMCC 21-0631	OP058962	OP059053	OP059004	OP135939	NA	Ren et al. (2021)
	UESTCC 23.0030	OR253120	OR253279	OR253194	NA	NA	Unpublished
Montagnula donacina	KUNCC 23-14428	OR583105	OR583124	OR583143	OR588096	OR588115	This study
	KUNCC 23-14429	OR583106	OR583125	OR583144	OR588097	OR588116	This study
Montagnula graminicola	MFLUCC 13-0352 ^T	KM658314	KM658315	KM658316	NA	NA	Liu et al. (2015)
Montagnula guiyangensis	HKAS 124556 [⊤]	OP605989	OP600484	OP600500	NA	NA	Sun et al. (2023)
	GUCC 22-0817	OP605990	OP600485	OP600501	NA	NA	Sun et al. (2023)
Montagnula jonesii	MFLUCC 16-1448 ^T	KY313619	KY273276	KY313618	KY313620	NA	Tennakoon et al. (2016)
	MFLU 18-0084	ON117282	ON117300	ON117269	ON158095	NA	Tennakoon et al. (2022)
Montagnula krabiensis	MFLUCC 16-0250 ^T	NR168179	NG068826	NG068385	MH412776	NA	Tibpromma et al. (2018)
Montagnula lijiangensis	HKAS 126540	OR583107	OR583126	OR583145	OR588098	OR588117	This study
	HKAS 126541 ⁺	OR583108	OR583127	OR583146	OR588099	OR588118	This study
Montagnula menglaensis	KUNCC 23-14422	OR583109	OR583128	OR583147	OR588100	OR588119	This study
	KUNCC 23-14423	OR583110	OR583129	OR583148	OR588101	OR588120	This study
	KUNCC 23-14424 ^T	OR583111	OR583130	OR583149	OR588102	OR588121	This study
Montagnula puerensis	KUMCC 20-0225 [™]	MW567739	MW575866	MW575864	MW575859	NA	Du et al. (2021)
	KUMCC 20-0331	MW567740	MW575867	MW575865	MW575860	NA	Du et al. (2021)
Montagnula saikhuensis	MFLUCC 16-0315 ^T	KU743209	KU743210	KU743211	NA	NA	Wanasinghe et al. (2016)
Montagnula scabiosae	MFLUCC 14-0954 ⁺	KT443907	KT443903	KT443905	NA	NA	Hongsanan et al. (2015)
Montagnula shangrilana	KUNCC 23-14433	OR583112	OR583131	OR583150	OR588103	OR588122	This study
	KUNCC 23-14434'	OR583113	OR583132	OR583151	OR588104	OR588123	This study
Montagnula thailandica	MFLUCC 17-0363	OL/82142	OL/82059	OL/80525	OL8/5102	OL828754	Senwanna et al. (2021)
	MFLUCC 17-1508	MT214352	NG0/0949	NG0/0158	M1235774	M1235810	Mapook et al. (2020)
	MFLUCC 21-0075	0P297807	0P297777	0P297791	0P321576	NA	Lu et al. (2022)
	ZHKUCC 22-0206	0P297808	0P297778	0P297792	0P321577	NA	Lu et al. (2022)
	ZHKUCC 22-0207	MZ538515	MZ538549	NA ODE00150	MZ567092	NA	Boonmee et al. (2021)
wontagnula thevetlae	HKAS 126963	00503114	OR583133	OR583152	00500100	00588124	
Neekelmusiais=++	HKAS 126964	ON007040	ON007000	UK583153	ON000100	ON000107	
weokaimusia jonannuimel		011007043	01007039	011007048	010009133	010009137	Mortimer (2022)
Neokalmusia jonahhulmei	KUMCC 21-0819	ON007044	ON007040	ON007049	ON009134	ON009138	Wanasinghe and Mortimer (2022)

Ex-type strains are indicated with superscript "T", and newly generated sequence is shown in bold. NA represents sequences that are unavailable in GenBank. CBS: Culture Collection of the Westerdijk Fungal Biodiversity Institute, Netherlands; CPC: Personal collection of P.W. Crous, Netherlands; HFG: Personal collection of Zhen-Zhu Zhao; GUCC: Guizhou University Culture Collection (GUCC), Guiyang, China; HKAS/KUNCC: Kunming Institute of Botany Culture Collection, China; HVVV: Personal collection of Wayne Pitt from *Vitis vinifera*; MFLUCC/MFLU: Mae Fah Luang University Culture Collection, China; HXAS/KUNCC: National Chiayi University Culture Collection, Taiwan, China; UESTCC: University of Electronic Science and Technology Culture Collection; ZHKUCC: Zhongkai University of Agriculture and Engineering Culture Collection.



Figure 1. Phylogenetic analysis of SSU, LSU, ITS, *tef*1- α , and *rpb*2 of the *Montagnula*. Species names given in bold are ex-type, ex-epitype and ex-paratype strains. Species names highlighted in blue are generated from this study. Branch support of nodes \geq 75% ML BS and \geq 0.95 PP is indicated above the branches. The genus *Montagnula* is depicted within a pale gray box, with new species highlighted in white, and the outgroup indicated by a blue box.



Figure 2. Geographical distribution of *Montagnula* species with known ITS sequence data. **a** the map summarizes data from the GlobalFungi database (shown by circles). Each circle symbolizes a unique sample, with each color representing the specific biome from which it has been collected **b** the distribution of *Montagnula* sequences as a percentage of total abundance across different biomes **c** the distribution of *Montagnula* sequences as a percentage of total abundance across different continents. See Suppl. material 1 for primary data.

Results

Phylogenetic analyses

In order to examine the evolutionary relationships of our new strains within *Montagnula*, phylogenetic analyses were performed based on the combined SSU, LSU, ITS, *tef*1-a, and *rpb*2 DNA sequences of 56 representatives of the genus and two strains from *Neokalmusia jonahhulmei* (KUMCC 21-0818, KUMCC 21-0819) as the outgroup taxon. The full dataset consisted of 4,268 characters including gaps (18S = 1,023 characters, 28S = 896, ITS = 508, *tef*1-a = 885, *rpb*2 = 956). The RAxML analysis of the combined dataset yielded a best-scoring tree with a final ML optimization likelihood value of -14,343.052271. The matrix had 1004 distinct



Figure 3. The distribution of *Montagnula* occurrences across oceans, continents and various substrates, as documented in the existing literature. On the x-axis, the logarithmic abundance of each record for different sources is displayed.



Figure 4. The species richness of recorded Montagnula species across different plant families (Table 1).

alignment patterns, with 23.88% undetermined characters or gaps. Parameters for the GTR + I + G model of the combined amplicons were as follows: Estimated base frequencies; A = 0.244145, C = 0.256118, G = 0.269851, T = 0.229886; substitution rates AC = 1.815063, AG = 3.954334, AT = 1.414215, CG = 1.362941, CT = 10.779403, GT = 1.000; proportion of invariable sites I = 0.559204; and gamma distribution shape parameter α = 0.542439. The Bayesian analysis ran 1,675,000 generations before the average standard deviation for split frequencies reached below 0.01 (0.009994). The analyses generated 16,751 trees, from which we sampled 12,564 trees after discarding the first 25% as burn-in. The alignment contained a total of 1,005 unique site patterns. The BI and ML trees were not in conflict; the ML tree is shown in Fig. 1. Where applicable, the phylogenetic results obtained (Fig. 1) are discussed in the descriptive notes below.

We conducted a thorough study of a compilation of data derived from multiple metabarcoding studies, which documented the occurrence of Montagnula species worldwide, excluding Antarctica. Among the continents, the highest number of studies were recorded in Asia, Australia, Europe, and North America (Fig. 2). These studies encompassed a diverse range of 11 distinct sources, revealing that sediments and "other" sources yielded the highest number of sequences (Fig. 3). Across different continents, the sequences obtained from various sources exhibited moderate similarity. However, in regions such as Asia, Australia, Europe, and North America, studies revealed Montagnula species from a diverse array of sources, in contrast to other studies, which identified species from a more limited selection of sources. Furthermore, in culture-based investigations, the primary focus was on extracting Montagnula species from plant substrates originating from 45 distinct plant families (Fig. 4). Among these families, Poaceae yielded the most substantial number of isolated species, followed by Asparagaceae and Fabaceae. Additionally, two records were also detected in mushrooms and human skin samples.

Taxonomy

Pleosporales Luttr. ex M.E. Barr, Prodromus to class Loculoascomycetes: 67 (1987)

Didymosphaeriaceae Munk, Dansk botanisk Arkiv 15 (2): 128 (1953)

Montagnula Berl., Icones Fungorum. Pyrenomycetes 2: 68 (1896)

Notes. This study presents an updated and comprehensive phylogenetic classification of the genus *Montagnula*, incorporating SSU, LSU, ITS, *tef1-a*, and *rpb2* DNA sequence analyses. By combining morphological and phylogenetic considerations, we have identified four new species, *M. lijiangensis*, *M. men-glaensis*, *M. shangrilana* and *M. thevetiae* within the genus. Additionally, this research accounts for the existing species *viz.*, *M. aquatica*, *M. aquilariae*, *M. chromolaenicola* and *M. donacina*. The note sections of this publication provide detailed information on these taxonomic accounts, including additional discussion and supporting evidence. Each newly identified species adds to the known biodiversity within the genus, expanding our knowledge of the ecological and morphological characteristics exhibited by *Montagnula* taxa.

Montagnula aquatica Y.R. Sun, Yong Wang bis & K.D. Hyde, Plants 12 (4, no. 738): 2 (2023)

MycoBank No: 900129

Descriptions and illustrations. See Sun et al. (2023).

Habitat and distribution. This species is found in freshwater habitats of Chiang Rai, Thailand, terrestrial habitats of Yunnan, China, inhabiting dead wood of deciduous hosts (Sun et al. 2023, this study).

Material examined. CHINA, Yunnan Province, Honghe Hani and Yi Autonomous Prefecture, Honghe County, Dayangjiexiang (23.389965°N, 102.225552°E, 1194 m), on dead woody litter of an unidentified plant, 13 March 2023, D.N. Wanasinghe, DWHH23-51 (HKAS 130322), new country and habitat record, living culture KUNCC 23-14425. *ibid*. 23.388966°N, 102.224786°E, 1215 m, DWHH23-51-2 (HKAS 130323), living culture KUNCC 23-14557.

Notes. Based on our phylogenetic analyses, we have determined that the newly collected strains (i.e. KUNCC 23-14425 and KUNCC 23-14557) are monophyletic with the ex-type strain of Montagnula aguatica (MFLU 22-0171). Further morphological investigations comparing our isolate with the type species have revealed similarities in the size range of the ascomata, asci, and ascospores, as well as the ascospore septation (Sun et al. 2023). Therefore, we document KUNCC 23-14425 and KUNCC 23-14557 as new records of Montagnula aquatica in China, accompanied by protein sequence data (tef1- α and rpb2) for this species. It is worth noting that the holotype of Montagnula aquatica was previously reported on submerged decaying wood in a freshwater habitat in Thailand, while our collection was made from a terrestrial habitat in China. This observation suggests that this fungus exhibits adaptability to a wide range of habitats, although its exploration in diverse geographic locations remains limited. The inclusion of Montagnula aquatica as a new record in China expands our understanding of the distribution and ecological preferences of this species in both terrestrial and aquatic habitats. Additionally, the protein sequence data obtained for this strain contributes valuable information to the existing knowledge on Montagnula aquatica. Further studies exploring the ecological aspects of this fungus in different geographic locations will provide deeper insights into its adaptability and potential ecological roles.

Montagnula aquilariae **T.Y. Du & Tibpromma, Mycosphere 14 (1): 705 (2023)** MycoBank No: 846332 Fig. 5

Description. Saprobic on dead woody litter of an unknown deciduous host. **Teleomorph** Ascomata 450–600 µm high × 480–550 µm diam., immersed to semi-erumpent, gregarious or rarely clustered, globose to subglobose, ostiolate. Ostiole $120-220 \times 70-110 \mu m$ ($\bar{x} = 139 \times 89 \mu m$, n = 5), papillate, central, straight, dark brown to black, filled with hyaline cells, periphyses are lacking. *Peridium* 20–40 µm thick on the sides and can reach up to 60 µm near the apex, with an outer layer consisting of heavily pigmented cells that have thick walls and exhibit a **textura angularis** to **textura globulosa** texture at the apex, **textura angularis** texture at the sides and base; the innermost layer consists of narrow, hyaline compressed rows of cells that merge with pseudoparaphyses. Hamath-



Figure 5. *Montagnula aquilariae* (HKAS 126542) **a**, **b** ascomata on natural wood surface **c** vertical section through an ascoma **d** ostiolar neck **e** peridium cells at the apex **f** peridium cells at the side **g** pseudoparaphyses **h**–**l** asci **m**–**r** ascospores (see vertuculose feature of the ascospore in r) **s**, **t** culture characters on PDA (s = above, t = reverse). Scale bars: 100 μ m (**c**, **d**); 50 μ m (**e**–**g**, **m**–**r**); 20 μ m (**h**–**l**).

ecium of 2–4 µm broad, dense, narrow, branched, cellular pseudoparaphyses. **Asci** 100–120 × 16–22 µm (\bar{x} = 110.8 × 18.4 µm, n = 20), bitunicate, fissitunicate, cylindrical-clavate to clavate, pedicel 30–50 µm long, 8-spored, biseriate, with a minute ocular chamber best seen in immature ascus. **Ascospores** 20–25 × 8.5–11 µm (\bar{x} = 21.8 × 9.6 µm, n = 30), ellipsoidal to narrowly oblong, straight or somewhat curved, ends conically rounded, golden-brown to dark brown, 1-septate, constricted at the septum, large guttules in each cell, verruculose, with a thin mucilaginous sheath. **Anamorph** Undetermined.

Habitat and distribution. This species is found in terrestrial habitats of Yunnan, China, specifically inhabiting dead woody twigs of deciduous hosts, including *Aquilaria sinensis* (Hyde et al. 2023, this study).

Material examined. CHINA, Yunnan Province, Kunming City, Kunming Institute of Botany (25.141723°N, 102.750013°E, 1970 m), on dead woody litter of an unidentified plant, 24 April 2022, L. Qinxian, KIB22-17-1 (HKAS 126542), living culture KUNCC 23-14430; *ibid*. 25.141487°N, 102.748863°E, 1982 m, K2B22-17-3 (HKAS 126543), living culture KUNCC 23-14431; *ibid*. K2B22-17-4 (HKAS 126544), living culture KUNCC 23-14432.

Notes. *Montagnula aquilariae* was recently introduced by Hyde et al. (2023) based on samples obtained from *Aquilaria sinensis* in Xishuangbanna, Yunnan Province. In our new collections, three strains (KUNCC 23-14430, KUNCC 23-14431, KUNCC 23-14432) exhibited a monophyletic relationship with the previously known strains of *Montagnula aquilariae* (KUNCC 22-10815 [ex-type] and KUNCC 22-10816). Through further morphological, ecological, and nucleotide (SSU, LSU, ITS, *tef*1- α) comparisons, we have confirmed that these new strains indeed belong to *Montagnula aquilariae*. Our research also provides additional insights into the characteristics of *Montagnula aquilariae*. Specifically, we report the verruculose feature of the ascospores and present *rpb*2 sequence data for this fungus, advancing our knowledge of its morphological and genetic attributes.

Montagnula chromolaenicola Mapook & K.D. Hyde, Fungal Diversity 101: 35 (2020)

MycoBank No: 557298

Descriptions and illustrations. See Mapook et al. (2020).

Habitat and distribution. This species was observed in terrestrial habitats in Mae Hong Son, Thailand, specifically on dead stems of *Chromolaena odorata* (Mapook et al. 2020). Additionally, it has also been found in terrestrial habitats in Yunnan, China, where it inhabits dead wood of deciduous hosts (this study).

Material examined. CHINA, Yunnan Province, Honghe County, Honghe Hani and Yi Autonomous Prefecture, Dayangjiexiang (23.389965°N, 102.225552°E, 1201 m), on a dead woody climber of an unidentified host, 13 March 2023, D.N. Wanasinghe, DWHH23-17A (HKAS 130321), living culture KUNCC 23-14426. *ibid.* 23.389295°N, 102.224780°E, 1200 m, on dead twigs of *Lagerstroemia* sp. DWHH23-33-2 (HKAS 126543), living culture KUNCC 23-14427; *ibid.* DWHH23-33-3 (HKAS 130320), living culture KUNCC 23-14558.

Notes. Through our phylogenetic analyses, we have determined that the newly isolated strains HH33 and HH17A exhibit a monophyletic relationship with the ex-type strain of *Montagnula chromolaenicola* (MFLUCC 17-1469). Upon conducting further investigations and morphological comparison of our collection with the type species, we have discovered several similarities. These include the size range of the ascomata, asci, and ascospores, as well as the ascospore septation (Mapook et al. 2020). Consequently, we hereby document our new collections (i.e. HKAS 130321, HKAS 126543 and HKAS 130320) as

new records of *Montagnula chromolaenicola* in China. In a recent study by Sun et al. (2023), *Montagnula chromolaenicola*, *M. puerensis*, *M. saikhuensis*, and *M. thailandica* were synonymized under the name *M. donacina* due to the absence of obvious branches in their phylogenetic tree and the close morphological resemblance between these species. However, it is important to note that most of these strains lack informative sequence data for *tef*1-a or *rpb*2. Our observations, on the other hand, have revealed that the inclusion of protein data in this group leads to the formation of distinct branches and independent lineages. Therefore, we propose retaining the older names for these species, except for *Montagnula thailandica*, until further research resolves this group using all available sequence data.

Montagnula donacina (Niessl) Wanas., E.B.G. Jones & K.D. Hyde, Index Fungorum 319: 1 (2017) MycoBank No: 552762

Descriptions and illustrations. See Pitt et al. (2014).

Habitat and distribution. This species has been reported worldwide on various hosts within terrestrial habitats (see Table 2). Specifically, it has been documented in Australia (*Calamus australis, Vitis vinifera*), Brazil (*Bambusoideae, Saccharum officinarum*), Central African Republic (*Coffea robusta*), China (*Althaea rosea, Craterellus odoratus, Trachycarpus fortunei*), Colombia (unknown plant), France (*Pseudosasa japonica*), Georgia (*Zea mays*), India (*Acacia sp., Adhatoda vasica, Ailanthus altissima, Annona squamosa, Cajanus cajan, Careya arborea, Citrus aurantiifolia, Clerodendrum infortunatum, C. multiflorum, Duranta repens, Ficus glomerata, Hibiscus sp., Ipomoea carnea, Mallotus philippinensis, Morus alba, Nerium odorum, Pistacia indica, Tectona grandis, Terminalia tomentosa*), Japan (*Phyllostachys bambusoides*), Myanmar (*Nephelium litchi*), Namibia (*Acacia reficiens*), Papua New Guinea (*Bambusoideae*), Paraguay (*Coffea arabica*), Philippines (*Premna cumingiana*), Portugal (*Arundo donax*), Sierra Leone (*Funtumia africana*), Thailand (dead wood) and the USA (*Platanus sp., Wikstroemia sp.*).

Material examined. CHINA, Yunnan Province, Honghe (23.424892°N, 102.231417°E, 600 m), on dead woody litter of an unidentified plant, 14 August 2022, D.N. Wanasinghe, DWHH22-23-1 (HKAS 126545), living culture KUNCC 23-14428. *ibid*. DWHH22-23-2 (HKAS 126546), living culture KUNCC 23-14429.

Notes. Wanasinghe et al. (2016) regarded *Munkovalsaria* as a synonym of *Montagnula* and established *Montagnula donacina* (=*Munkovalsaria donacina*). So far, *Montagnula donacina* stands as the most extensively distributed species within the genus. Despite its global presence, there is a scarcity of molecular data available for *Montagnula donacina*. A preliminary analysis revealed only 20 sequence data entries when searching for "*Montagnula donacina*" in the NCBI database, originating from only seven strains: HFG07004, HKAS 124552, HVVV01, KUMCC 21-0579, KUMCC 21-0631, KUMCC 21-0653, and UESTCC:23.0030. Our phylogenetic analysis demonstrated a close relationship between two strains designated as *Montagnula donacina* (HVVV01 and HFG07004) and the type strain of *Montagnula chromolaenicola* (MFLUCC

17-1469). Additionally, we observed that the strains of *Montagnula thailandica* formed a monophyletic group alongside the remaining *Montagnula donacina* strains (HKAS 124552, KUMCC 21-0579, KUMCC 21-0631, KUMCC 21-0653, and UESTCC:23.0030). Furthermore, two newly generated sequences, KUNCC 23-14428 and KUNCC 23-14429, were also clustered with the strains of *Montagnula donacina*. We hereby introduce these two strains as belonging to *Montagnula donacina* and provide *rpb2* sequence data for this species for the first time.

Montagnula lijiangensis Wanas., sp. nov.

MycoBank No: 850093 Fig. 6

Etymology. The specific epithet "lijiangensis" refers to Lijiang, Yunnan Province, where the holotype was collected.

Holotype. HKAS 126541.

Description. Saprobic on dead woody litter of Quercus sp. Teleomorph Ascomata 500-700 µm high × 500-600 µm diam., immersed, gregarious or rarely clustered, globose to subglobose, ostiolate. Ostiole 100-140 × $80-120 \ \mu m$ ($\overline{x} = 125 \times 96 \ \mu m$, n = 5), apapillate, central, straight, filled with hyaline cells. *Peridium* 20–30 μ m thin on the sides and can reach up to 70 μ m near the apex, with an outer layer consisting of heavily pigmented cells that have thick walls and exhibit a textura angularis texture at the apex, textura angularis texture at the sides and base; the innermost layer consists of narrow, hyaline compressed rows of cells. Hamathecium of 3-7.5 µm broad, dense, narrow, branched, cellular pseudoparaphyses that are swollen at the base. Asci $130-160 \times 20-26 \ \mu m$ (x = 152.8 × 23.9 \ \mu m, n = 20), bitunicate, fissitunicate, cylindrical-clavate to clavate, pedicel 30-60 µm long, 8-spored, uni to biseriate, with a minute ocular chamber best seen in immature ascus. Ascospores 22-26 \times 10–14 µm (\overline{x} = 24.8 \times 11.8 µm, n = 30), ellipsoidal to narrowly oblong, mostly straight, with conically rounded ends at the immature stage that become rounded when mature, golden-brown to dark brown, 1-septate and constricted at the septum, with large guttules in each cell, verruculose, surrounded by a thick mucilaginous sheath. Anamorph Undetermined.

Habitat and distribution. This species is found in terrestrial habitats of Yunnan, China, inhabiting dead woody twigs of deciduous hosts (this study).

Material examined. CHINA, Yunnan Province, Lijiang, Yulong County (26.86389°N, 99.824738°E, 2725 m), on dead woody litter of *Quercus* sp. (Fagaceae), 17 August 2021, L. Qinxian, STX09-03-1 (*holotype*, HKAS 126541, *ibid*. 26.863484°N, 99.824548°E, 2706 m, STX09-03-3 (HKAS 126540).

Notes. The analysis of two newly generated sequences revealed a monophyletic clade in our phylogenetic analysis (Fig. 1), demonstrating a close phylogenetic relationship to *Montagnula aquilariae*. This relationship is further supported by morphological features such as asci and ascospores. However, a comparison of nucleotide differences (without gaps) between these two clades (KUNCC 22-10815 and KUNCC 23-14430 vs HKAS 126541) showed 12/508 (2.3%) differences in the ITS region, 15/885 (1.7%) differences in the tef1- α region, and 19/956 (2%) differences in the *rpb2* region.



Figure 6. Montagnula lijiangensis (HKAS 126541, holotype) **a**, **b** ascomata on natural wood surface **c** vertical section through an ascoma **d** ostiolar neck and peridium cells at the apex **e** pseudoparaphyses \mathbf{f} -i asci \mathbf{j} -o ascospores (see verruculose feature of the ascospore in **k**). Scale bars: 100 µm (**c**); 20 µm (**d**, \mathbf{f} -i); 10 µm (**e**-o).

Montagnula menglaensis Wanas., sp. nov.

MycoBank No: 850094 Fig. 7

Etymology. The specific epithet "menglaensis" refers to Mengla County, Yunnan Province, where the holotype was collected.

Holotype. HKAS 130318.

Description. Saprobic on dead culms of *Indocalamus tessellatus* (Munro) Keng f. Teleomorph Ascomata 200–300 μ m high × 240–320 μ m diam.,



Figure 7. *Montagnula menglaensis* (HKAS 130318, holotype) **a**–**c** ascomata on natural wood surface **d**, **e** vertical section through ascomata **f**, **g** pseudoparaphyses **h** peridium **i**–**k** asci **l**, **m** ascospores (see vertuculose feature of the ascospore in **n**) **o**, **p** culture characters on PDA (o = above, p = reverse) **q**, **r** conidiomata **s** pycnidial wall **t** conidia. Scale bars: 100 μ m (**d**, **e**); 10 μ m (**f**–**h**, **I**–**n**, **s**, **t**); 20 μ m (**i**–**k**).

immersed, gregarious or rarely clustered, globose to subglobose. Peridium 10-25 µm thin with an outer layer consisting of heavily pigmented cells that have thick walls and exhibit a textura angularis texture at the sides and base; the innermost layer consists of narrow, hyaline compressed rows of cells. Hamathecium of 3-7.5 µm broad, dense, branched, cellular pseudoparaphyses that are swollen at some septa. Asci 60-80 \times 9-11 µm (\overline{x} = 71 \times 9.8 µm, n = 15), bitunicate, fissitunicate, cylindrical-clavate, pedicel 15-30 µm long, 8-spored, uni to biseriate, with a minute ocular chamber best seen in immature ascus. Ascospores 10.5-14 × 4.5-5.5 µm (x = 12.6 × 5.1 µm, n = 20), ellipsoidal, mostly straight, with conically rounded ends, golden-brown to dark brown, 1-septate and constricted at the septum, upper cell wider than the lower cell, with large guttules in each cell, verruculose, and surrounded by a thin mucilaginous sheath which is thicker at both ends. Anamorph Coelomycetous on PDA. Conidiomata pycnidial, gregarious, immersed to superficial, globose to subglobose, dark brown to black. Pycnidial wall thin, composed of brown cells of textura angularis. Conidiogenous cells did not observed. Conidia 2.3-3.3 × $1.4-2 \mu m$ ($\bar{x} = 3 \times 1.7 \mu m$, n = 30), hyaline, aseptate, round to oblong or ellipsoidal, with small guttules.

Culture characteristics. Ascospores germinated on PDA within 24 h. Following a two-week incubation period at 25 °C, the colonies on PDA medium reached a diameter of 5 cm. These colonies exhibited an undulate margin, initially appearing creamy whitish and transitioning to orange, raised in the center. The colonies were orange at the center and a creamy orange towards the periphery when observed from the reverse side.

Habitat and distribution. This species is found in terrestrial habitats of Yunnan, China, inhabiting dead woody twigs of deciduous hosts (this study).

Material examined. CHINA, Yunnan Province, Xishuangbanna, Mengla County (21.588394°N, 101.435042°E, 776 m), on dead culms of *Indocalamus tessellatus*, 29 January 2022, L. Qinxian, ML23-7-3 (holotype, HKAS 130318), ex-type KUNCC 23-14424; *ibid*. 21.589178°N, 101.435752°E, 782 m, ML23-7-2 (HKAS 130316), living culture KUNCC 23-14422; *ibid*. ML23-7-5 HKAS 130317), living culture KUNCC 23-14423.

Notes. *Montagnula menglaensis* is described as a novel species based on its holomorph. The anamorph of *Montagnula* is rarely encountered; however, Crous et al. (2020) recently reported *Montagnula cylindrospora* based on its anamorphic features. The conidia of *Montagnula menglaensis* resemble to those of *M. cylindrospora*, although the latter fungus exhibits a more cylindrical shape.

Montagnula shangrilana Wanas., sp. nov.

MycoBank No: 850095 Fig. 8

Etymology. The specific epithet "shangrilana" refers to Shangri-La, Yunnan Province, where the holotype was collected.

Holotype. HKAS 126539.

Description. Saprobic on dead woody litter of *Rhododendron* sp. **Teleomorph** Ascomata 120–180 µm high × 150–210 µm diam., immersed to semi-erumpent,



Figure 8. *Montagnula shangrilana* (HKAS 126541, holotype) **a** ascomata on natural wood surface **b** vertical section through an ascoma **c** pseudoparaphyses **d** peridium cells **e**–**h** asci **i**–**o** ascospores (see vertuculose feature of the ascospore in **o**). Scale bars: 100 μm (**b**); 10 μm (**c**, **d**, **j**–**o**); 20 μm (**e**–**h**).

gregarious or rarely clustered, globose to subglobose, ostiolate. **Ostiole** $80-110 \times 50-80 \ \mu m$ ($\overline{x} = 100 \times 64 \ \mu m$, n = 6), papillate, central, straight, filled with hyaline cells. **Peridium** $10-20 \ \mu m$ thin on the sides and can reach up to 40 μm near

the apex, with an outer layer consisting of heavily pigmented cells that have thick walls and exhibit a **textura angularis** arrangement at the apex, **textura angularis** texture at the sides; the innermost layer consists of hyaline compressed rows of cells. **Hamathecium** of 2–4.5 µm broad, dense, branched, cellular pseudoparaphyses. **Asci** 90–140 × 20–30 µm ($\bar{x} = 116.2 \times 24 \mu m$, n = 10), bitunicate, fissitunicate, cylindrical-clavate, pedicel 25–40 µm long, 8-spored, uni to biseriate, with a minute ocular chamber best seen in immature ascus. **Ascospores** 48–60 × 17–22 µm ($\bar{x} = 55.8 \times 19.3 \mu m$, n = 20), ellipsoidal to narrowly oblong, mostly straight, with conically rounded ends at the immature stage that become rounded when mature, golden-brown to dark brown, 3-septate, with large guttules in each cell, verruculose, surrounded by a thick mucilaginous sheath. **Anamorph** Undetermined.

Culture characteristics. Ascospores germinated on PDA within 24 h. Following a two-week incubation period at 25 °C, the colonies on PDA medium reached a diameter of 5 cm. These colonies exhibited a filiform margin, initially appearing whitish and transitioning to greenish gray, raised in the center. The colonies were grey at the center and a greenish gray towards the periphery and radiated when observed from the reverse side.

Habitat and distribution. This species is found in terrestrial habitats of Yunnan, China, inhabiting dead woody twigs of deciduous hosts, in a subalpine environment (this study).

Material examined. CHINA, Yunnan Province, Diqing Tibetan Autonomous Prefecture, Shangri-La (27.289707°N, 100.034477°E, 2744 m), on dead woody litter of *Rhododendron* sp. (Ericaceae), 22 August 2021, L. Qinxian, WTS8-2-2 (holotype, HKAS 126539), ex-type KUNCC 23-14434; *ibid*. (27.290007°N, 100.035233°E, 2833 m, WTS8-2 (HKAS 126538), living culture KUNCC 23-14433.

Notes. In the combined SSU, LSU, ITS, *tef*1- α , and *rpb*2 phylogenetic analysis, two strains of *Montagnula shangrilana* (HKAS 126538, HKAS 126539) formed a monophyletic clade closely related to *M. camporesii* (MFLUCC 16-1369), *M. cirsii* (MFLUCC 13-0680), and *M. scabiosae* (MFLUCC 14-0954). While there were slight variations in size, shape, and color, all four species shared the common characteristic of 3-transversely septate ascospores. The sequence data of *Montagnula camporesii*, *M. cirsii*, and *M. scabiosae* showed no significant differences in their base pair comparisons, suggesting that they may be conspecific. Morphologically, these three species exhibited clavate asci and ellipsoid to fusiform, brown, overlapping, 3-septate ascospores. In contrast, our newly discovered species differed from these three species by 10/508 (1.96%) differences in the ITS region, 13/885 (1.5%) differences in the *tef*1- α region, and 15/956 (1.56%) differences in the *rpb*2 region (only *M. camporesii* possesses *rpb*2).

Montagnula thevetiae Wanas., sp. nov.

MycoBank No: 850096 Fig. 9

Etymology. The specific epithet "thevetiae" refers to the host *Thevetia peruviana* from which the holotype was isolated. **Holotype.** HKAS 126964.



Figure 9. Montagnula thevetiae (HKAS 126564, holotype). **a**, **b** ascomata on natural wood surface **c** vertical section through an ascoma **d** closeup of ostiole **e** pseudoparaphyses **f**-**h** asci **j**-**l** ascospores **m**, **n** culture characteristics on PDA (m = above, n = reverse). Scale bars: 100 μ m (c); 50 μ m (d, f-h); 10 μ m (e, i-l).

Description. Saprobic on dead twigs of *Thevetia peruviana*. Teleomorph Ascomata 140–160 µm high × 150–190 µm diam., immersed, gregarious or rarely clustered, globose to subglobose, ostiolate. Ostiole 40–65 × 50–90 µm ($\bar{x} = 50 \times 78$ µm, n = 6), papillate, central, straight, filled with hyaline to brown cells. Peridium 10–20 µm thin on the sides and can reach up to 30 µm near the apex, with an outer layer consisting of heavily pigmented cells that have thick walls and textura angularis arrangement, the inner layer consists of hyaline compressed rows of cells. Hamathecium of 2–3.5 µm broad, dense, branched, cellular pseudoparaphyses. Asci 110–160 × 25–35 µm ($\bar{x} = 126.4 \times 30.3$ µm, n = 12), bitunicate, fissitunicate, cylindrical-clavate, pedicel 25–35 µm long, 8-spored, uni to biseriate, with a minute ocular chamber best seen in immature ascus. **Ascospores** $30-40 \times 11.5-14 \ \mu m$ ($\overline{x} = 37.3 \times 12.8 \ \mu m$, n = 20), ellipsoidal to narrowly oblong, straight to curved, with conically rounded ends, brown to dark brown, 1-septate, constricted at the septum, with large guttules in each cell, verruculose, surrounded by a thin mucilaginous sheath. **Anamorph** Undetermined.

Culture characteristics. Ascospores germinated on PDA within 24 h. Following a two-week incubation period at 25 °C, the colonies on PDA medium reached a diameter of 4 cm. These colonies exhibit an irregular, flattened to slightly raised morphology and display various color sectors ranging from white, creamy orange to pale brown. The reverse side of the colonies appears creamy orange, with occasional dark patches that can be observed.

Habitat and distribution. This species is found in terrestrial habitats of Yunnan, China, inhabiting dead woody twigs of *Thevetia peruviana* (this study).

Material examined. CHINA, Yunnan Province, Kunming city, Kunming Institute of Botany (25.142238°N, 102.750354°E, 1971 m), on dead twigs of *Thevetia peruviana*, 24 April 2022, L. Qinxian, K2B22-26-2 (holotype, HKAS 126964), *ibid*. (25.140859°N, 102.749045°E, 1968 m, K2B22-26 (HKAS 126963).

Notes. *Montagnula thevetiae* is isolated from the dead twigs of *Thevetia peruviana*. The newly obtained sequences of this fungus formed a monophyletic clade closely related to *Montagnula menglaensis*. Morphologically, they share similarities in having 1-septate ascospores, although *Montagnula thevetiae* exhibits a darker pigmentation. On the other hand, *Montagnula thevetiae* differs from *M. menglaensis* by 15/1023 (1.46%) differences in the SSU region, 19/895 (2.12%) differences in the LSU region, 32/508 (6.3%) differences in the ITS region, 27/885 (3%) differences in the *tef*1- α region, and 86/956 (9%) differences in the *rpb*2 region.

Discussion

Montagnula species in Yunnan Province

The study of lignicolous microfungi in Yunnan Province resulted in the collection of eight Montagnula species, including four novel species. This study contributes to our understanding of the diversity and distribution of Montagnula species and provides insight into the ecological roles played by these fungi in their respective habitats. Montagnula aquatica was previously documented as occurring on submerged decaying wood within a freshwater habitat in Thailand (Sun et al. 2023). However, our recent collection of this species was obtained from a terrestrial habitat in China. The holotype was collected in the Bandu District of the Chiang Rai Province, situated at an approximate elevation of 400-450 m and characterized by a tropical climate. The collection site was near to a waterfall (Sun et al. 2023). In contrast, our new collections were made in the Honghe region of Yunnan Province, which possesses an elevation of approximately 1200 m. The local environment in this region is characterized by poor, eroded soils, steep valleys, and a subtropical climate. This observation suggests that Montagnula aquatica may possess an adaptable nature, enabling it to thrive in a wide range of habitats across diverse geographic locations. Montagnula aquilariae, another species within the genus, has been identified in the terrestrial habitats of Yunnan, China. It specifically colonizes dead woody twigs of deciduous hosts, including Aquilaria sinensis (Hyde et al. 2023). The holotype of this species was collected from a hilly area in Nanmo, Menghai and Xishuangbanna, situated at an elevation of ~1100 m and characterized by a tropical climate. Additional collections were made from Kunming, located within the same province but at an elevation of ~2000 m, and characterized by a warm and temperate climate. Montagnula chromolaenicola has been observed in terrestrial habitats in Thailand, particularly on dead stems of Chromolaena odorata (Mapook et al. 2020). The holotype of this species was collected from the Mae Yen mountainous area of Mae Hong Son Province, at an elevation of ~900 m. The local environment of this area exhibits a tropical savanna climate. In our study, we collected this fungus from a terrestrial habitat within the steep valleys of subtropical Honghe, Yunnan, China. In this region, Montagnula chromolaenicola was found to inhabit the dead woody litter of deciduous hosts. Montagnula donacina has been reported across various terrestrial habitats worldwide, with the majority of records originating from India (Table 1). This species primarily associates with hosts from the Poaceae family. In our study, we collected Montagnula donacina from the subtropical Honghe region in China, specifically on decaying woody litter at an elevation of ~600 m. Montagnula lijiangensis was collected from terrestrial habitats at a high elevation of ~2725 m. This species was found on dead woody litter of Quercus sp. within an environment characterized by a mild subtropical highland climate. Montagnula menglaensis was discovered in the terrestrial habitats of Mengla County, Yunnan, China. It was observed colonizing dead culms of *Indocalamus tessellatus*. The local environment of this region exhibits a tropical savanna climate, with an elevation of ~800 m. Montagnula shangrilana was found in the terrestrial habitats of Shangri-La, Yunnan, China, where it inhabits dead woody twigs of Rhododendron sp. This species has also been observed at higher elevations, reaching ~2800 m, within an environment characterized by a humid continental climate. Montagnula thevetiae was discovered within the terrestrial habitat of the botanical garden at the Kunming Institute of Botany in Yunnan, China. This species was found colonizing dead woody twigs of Thevetia peruviana. The collection site is situated at an elevation of ~2000 m and experiences a warm and temperate climate.

Taxonomic reassessment and phylogenetic analysis of *Montagnula* species

In a recent study conducted by Sun et al. (2023), *Montagnula chromolaenicola*, *M. puerensis*, *M. saikhuensis*, and *M. thailandica* were regarded as the synonyms of *M. donacina* (Wanasinghe et al. 2016). This decision was based on the absence of clear branches in their phylogenetic tree and the close morphological resemblance between these species. However, upon further examination, it was observed in this study that only *Montagnula donacina* and *M. thailandica* appear to be conspecific, based on combined gene analyses (Fig. 1). When informative sequence data such as *tef1-a* or *rpb2* were added to the analysis for *Montagnula chromolaenicola*, *M. puerensis*, *M. saikhuensis*, and *M. thailandica*, distinct branches and independent lineages were observed (Fig. 1). This suggests that these species are separate entities. Notably, two sequences of *M. donacina* (HVVV01 and HFG07004) were found to be monophyletic with the type strain of *Montagnula chromolaenicola* (MFLUCC 17-1469), indicating that they belong

to the latter species. In the case of Montagnula camporesii (MFLUCC 16-1369), M. cirsii (MFLUCC 13-0680), and M. scabiosae (MFLUCC 14-0954), the type strains formed a monophyletic lineage as a single species. Nucleotide base pair comparison of LSU, SSU, and ITS between these three strains did not reveal any differences. Therefore, it is suggested that Montagnula camporesii and M. cirsii should be synonymized under M. scabiosae, as it is the oldest name. However, it is important to note that this taxonomic clarification was not within the scope of our study, and future studies should compare the morphology of the holomorphs to resolve any remaining taxonomic confusion. Apart from these two clades, all other species formed distinct lineages in the multi-gene phylogenetic analysis. Out of the accepted 54 species in this genus, sequence data are currently available for only 28 species, including the four new species introduced in this study. This leaves approximately 48% of the species in need of phylogenetic sorting. Hence, future studies based on taxonomy should prioritize obtaining DNA sequence data for the remaining species. They should aim to acquire informative sequence data, such as tef1-a and rpb2, for all strains, and focus on revising the taxonomy of all species within the genus Montagnula.

Morphological characterization of Montagnula species

The genus Montagnula exhibits rare reporting of its anamorphic features, with only one species, M. cylindrospora, described from its anamorph in addition to our study (Crous et al. 2020). This finding has helped confirm its phylogenetic placement within the genus. The teleomorph, rather than the anamorph, appears to be more commonly observed in the natural environment. The majority of Montagnula species produce immersed or semi-immersed ascomata, which are globose to subglobose in shape and possess a central papillate ostiole. However, there are a few exceptions, such as M. camporesii, M. cirsii, and M. Iongipes, which have been reported to have superficial ascomata. Upon closer examination, it becomes apparent that Montagnula camporesii and M. cirsii actually have semi-immersed ascomata, as illustrated in Hyde et al. (2016, 2020). It is worth mentioning that Aptroot (1995) did not illustrate the ascomata, and their orientation remains unclear. Additionally, only one species, Montagnula bellevaliae, has been reported to possess an eccentric ostiole (Hongsanan et al. 2015). The peridium cells of Montagnula species commonly exhibit a thickwalled arrangement with a textura angularis pattern. Notably, the cells near the apex are often thicker compared to those on the sides and base walls. A distinguishing characteristic for species within this genus is the presence of swollen cells in pseudoparaphyses. The asci, typically exhibit a cylindrical to clavate shape with a prominent pedicel. Ascospores in Montagnula are predominantly described as ellipsoidal to fusiform, pigmented, and septate. The majority of species (>15) have ascospores with a single septum, while some species, including M. dasylirionis, M. dura, M. infernalis, M. mohavensis, M. phragmospora, M. spinosella, and M. yuccigena, have been reported to possess muriform spores (Du et al. 2023). The remaining species have ascospores with either 3 or 5 septa. A distinct characteristic within the genus is the verruculose surface texture of the ascospores which is neglected by most of the studies. Only Montagnula appendiculata, M. chiangraiensis, and M. chromolaenae have been documented to possess polar appendages (Aptroot 2004; Mapook et al. 2020).

Ecological preferences and worldwide distribution of *Montagnula* species through culture-dependent studies

The information we gathered from our culture-based investigations revealed that Montagnula species were found on 105 genera in 45 distinct plant families, in 55 countries (Table 1). This highlights the wide ecological range and adaptability of Montagnula species across different hosts and geographic regions. Among the plant families, Poaceae emerged as the most significant contributor, yielding the highest number of isolated Montagnula species (Fig. 4). This finding suggests a potential association between Montagnula species and grasses, indicating the ecological importance of the Poaceae family in the life cycle and development of Montagnula species. Furthermore, Montagnula species were also detected in other plant families, such as Asparagaceae and Fabaceae, indicating their potential interactions with a diverse range of host plants. Among the more than 100 plant genera associated with Montagnula species, Agave (Asparagaceae), Opuntia (Cactaceae), Phoenix (Arecaceae), Ammophila (Poaceae), and Yucca (Asparagaceae) were found to have the greatest number of species, collectively representing 25% of the total count. This highlights the potential preference of Montagnula species for these specific plant genera within their respective families. The analysis of country-wise distribution revealed that India had the highest number of Montagnula entries (Table 1). The majority of these entries were attributed to Montagnula donacina, indicating a wide distribution of this species in India. Among the countries where Montagnula species were reported, China exhibited the highest diversity with nine different species, followed by Italy and the USA with seven different species each. This suggests regional variations in the diversity and distribution of Montagnula species. Interestingly, our study also detected Montagnula species in mushrooms and human skin samples, indicating their presence in alternative sources and potential interactions with other organisms. This highlights the need for further investigation into the ecological roles and potential impacts of Montagnula species in these non-traditional habitats. Except for Antarctica, Montagnula donacina has been reported from various countries across all six continents. Additionally, it has been identified in 25 different plant families. Investigating the reasons behind its wide distribution and adaptation to diverse ecological conditions would be intriguing. Future studies should focus on the morphological features, secondary metabolites, and gene data-based analyses of the species. To date, only six studies, including this one, have provided entries featuring both morphology and DNA-based sequence data evidence (Pitt et al. 2014; Zhao et al. 2018; Ren et al. 2022a; Li et al. 2023; Sun et al. 2023).

These findings elucidate the global distribution and ecological preferences of *Montagnula* species, highlighting the significance of different sources and plant families in their occurrence and potential ecological interactions. The wide range of sources from which species were identified suggests their adaptability and potential ecological roles in various ecosystems. The study also has important implications for our understanding of the ecology and biology of *Montagnula* fungi. All of the new species described in this study were found to be associated with dead wood, indicating the role that these fungi play in the decomposition of organic matter in forest ecosystems. We suggest that future studies could investigate the functional roles played by *Montagnula* fungi in ecosystem processes, such as carbon and nutrient cycling.

Global biogeography and ecological versatility of *Montagnula* based on metabarcoding data through culture-independent studies (NGS)

In addition to the taxonomic novelties, this study utilized metabarcoding data from the GlobalFungi database (Větrovský et al. 2020) to gain insights into the global diversity and distribution of Montagnula. Metabarcoding is a valuable tool that allows for the rapid identification of multiple species from complex environmental samples, providing confirmation of their presence in specific habitats. The analysis of multiple metabarcoding studies provided comprehensive information on the occurrence and distribution patterns of Montagnula species worldwide. The distribution of Montagnula across diverse biomes underscores their remarkable ecological adaptability and diversity. Forests, constituting 61% of their habitats, emerge as the predominant biome, indicating a strong preference or adaptation of the genus to forest ecosystems. Grasslands, accounting for 18%, also represent a significant habitat, suggesting the versatility in adapting to open and semi-open landscapes of them. Croplands (6%) and shrublands (7%) further exemplify the adaptability of Montagnula, thriving in both cultivated areas and natural, low-vegetation environments. Notably, woodlands and anthropogenic areas, representing 2% and 1% respectively, highlight the ability to exist in moderately wooded areas and regions significantly influenced by human activity. Additionally, their presence in aquatic environments, deserts, and wetlands, each accounting for 1% of their habitats, along with a notable 3% in mangroves, reflects the broad ecological niche of them. The marginal occurrence in tundras (0.1%) suggests a limited but notable ability to survive in extreme cold climates. The presence of Montagnula in such varied biomes underscores its ecological versatility and the importance of diverse habitats in understanding its biogeography.

The presence of *Montagnula* species has been documented in various regions of Africa, Arctic Ocean, Asia, Australia, Europe, Indian Ocean, North America, Pacific Ocean and South America indicating their widespread occurrence and ecological significance in these areas. In Asia, *Montagnula* species have been observed in multiple countries, including China, India, Indonesia, Iran, Japan, Malaysia, South Korea, Thailand and others (Suppl. material 1). The diverse range of habitats in these regions, such as freshwater habitats, terrestrial environments, and mountainous areas, offer suitable ecological niches for *Montagnula* colonization and growth. The detection of *Montagnula* species in different ecological contexts within Asia suggests their ability to adapt to various local conditions and substrates, contributing to their wide distribution across the continent. For example, in China, *Montagnula* species have been found in diverse habitats ranging from aquatic environments to forests and grasslands (Suppl. material 1), indicating their adaptability to different ecosystems. This adaptability may be attributed to their ability to utilize a wide range of organic materials as substrates, including decaying plant remains.

Australia also exhibits a notable presence of *Montagnula* species, indicating their occurrence in diverse habitats throughout the continent (Bissett et al. 2016; Luis et al. 2019; Turner et al. 2019; Gui et al. 2023). The unique ecosystems in Australia, including deserts, rainforests and grasslands, provide opportunities for *Montagnula* to establish themselves in different ecological niches. The metabarcoding studies were used for various biomes i.e. anthropogenic, aquatic, cropland, desert, forest, grassland, mangrove, shrubland, wetland and woodland (Fig. 3). This highlights the higher presence and distribution of *Montagnula* in different

habitats within Australia. In Europe, Montagnula species have been recorded in several countries, including Austria, Belgium, Czech Republic (highest), Estonia, France, Germany, Italy, Netherlands Slovenia, Sweden Switzerland and Spain (Suppl. material 1). The presence of Montagnula in Europe suggests their ability to adapt to different climates and ecological conditions. This broad distribution across Europe indicates the need for further investigation into the ecological preferences and potential impacts of Montagnula species in this region. For instance, studies in Europe have identified Montagnula species in different habitats, such as anthropogenic, aquatic, cropland, desert, forest, grassland, shrubland, tundra, wetland and woodland (Suppl. material 1). Africa and North America also demonstrates a diverse distribution of Montagnula species, with the majority of records coming from the South Africa, Namibia, Botswana, Zambia, Mozambique, Kenya, Kenya and Ivory Coast in Africa respectively. United States was having the highest number of sampling locations in North America. Comparatively, the occurrences of Montagnula species using metabarcoding data in China, the USA, and European countries are relatively well-documented. However, the rest of the world remains a mystery in terms of Montagnula distribution. For example, the majority of Asia, including India and Russia, lacks metabarcoding data for Montagnula species. This emphasizes the need for more extensive research and data collection to better understand the global distribution of Montagnula and its ecological roles.

Conclusion

Our study on Montagnula species has provided valuable insights into their ecological preferences and global distribution patterns. The findings indicate that these fungi exhibit a wide range of climatic distribution, suggesting their adaptability to different temperature ranges and potentially reducing their vulnerability to climate change. The ability of Montagnula species to utilize a diverse range of organic materials as substrates, including decaying plant remains, contributes to their widespread distribution across various habitats. Our analysis revealed a diverse range of sources from which Montagnula species were detected, including freshwater and terrestrial habitats, further highlighting their ecological versatility. Sediments were found to be particularly rich in Montagnula sequences, suggesting their potential as suitable habitats for colonization and growth. Although moderate sequence similarity was observed across different sources and continents, regional variations in ecological preferences and distribution patterns were evident. The diverse host range observed in our field collections aligns with global meta-barcoding sources, emphasizing the ability of Montagnula species to thrive in various ecosystems. The ecological adaptability and versatility of Montagnula species underscore their success in colonizing diverse habitats. Further research and investigation into their biogeography will contribute to our understanding of their global distribution, ecological roles, and potential impacts on ecosystems. This knowledge is crucial for effective conservation efforts, understanding ecosystem dynamics, and managing ecological balance in different regions.

Acknowledgments

We gratefully thank the Chinese Academy of Sciences for providing molecular laboratory facilities.

Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

Funding

Dhanushka Wanasinghe thanks CAS President's International Fellowship Initiative (number 2021FYB0005), the National Science Foundation of China (NSFC) under the project code 32150410362, Smart Yunnan Project (Young Scientists) under project code E13K281261 and the Postdoctoral Fund from Human Resources and Social Security Bureau of Yunnan Province. Thilina Nimalrathna expresses gratitude for the support provided by the Belt and Road Chinese Government Scholarship and The Alliance of International Science Organizations (ANSO) Ph.D. scholarship. We also extend our appreciation to the Researchers Supporting Project at King Saud University, Riyadh, Saudi Arabia, for funding this research project (Fund no. RSP2024R784). Jianchu Xu thanks National Natural Science Foundation of China (grant number: 31861143002), the Yunnan Provincial Science and Technology Department (grant number: 202101AS070045), Yunnan Provincial Sciences and Technology Oppartment (grant number: 202205AM070007) and Yunnan Department of Sciences and Technology of China (grant number: 202302AE090023).

Author contributions

Conceptualization: DNW. Data curation: LQX, DNW. Formal analysis: TKF, DNW, TSN. Investigation: TSN, DNW. Methodology: TSN, DNW. Project administration: PEM, JX. Resources: JX. Supervision: JX, PEM. Writing – original draft: TSN, DNW. Writing – review and editing: PEM, TKF.

Author ORCIDs

Dhanushka N. Wanasinghe [©] https://orcid.org/0000-0003-1759-3933 Thilina S. Nimalrathna [©] https://orcid.org/0000-0002-2368-042X Li Qin Xian [©] https://orcid.org/0009-0006-4936-9409 Turki KH. Faraj [©] https://orcid.org/0000-0002-6012-8474 Jianchu Xu [©] https://orcid.org/0000-0002-2485-2254 Peter E. Mortimer [©] https://orcid.org/0000-0003-3188-9327

Data availability

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

References

- Abeywickrama PD, Wanasinghe DN, Karunarathna SC, Jayawardena RS, Hyde KD, Zhang W, Li X, Yan J (2020) A new host report of *Diaporthe manihotia* (Diaporthales, Ascomycota) from *Camellia* sp. in Yunnan province, China. Asian Journal of Mycology 3(1): 473–489. https://doi.org/10.5943/ajom/3/1/17
- Aluthmuhandiram JVS, Wanasinghe DN, Chethana KWT, Gafforov Y, Saichana N, Hong LX, Jiye Y, Mamarakhimov O (2022) Lophiostomataceae (Dothideomycetes): Intro-

ducing *Lophiostoma khanzada-kirgizbaeva* sp. nov. and *Paucispora xishanensis* sp. nov. Phytotaxa 559(3): 247–262. https://doi.org/10.11646/phytotaxa.559.3.3

- Aptroot A (1995) Redisposition of some species excluded from *Didymosphaeria* (Ascomycotina). Nova Hedwigia 60: 325–379.
- Aptroot (2004) Two new ascomycetes with long gelatinous appendages collected from monocots in the tropics. Studies in Mycology 50: 307–311.
- Aptroot (2006) *Mycosphaerella* and its anamorphs: 2. Conspectus of *Mycosphaerella*. CBS Biodiversity Series 5: 1–231.
- Ariyawansa H, Tanaka K, Thambugala KM, Phookamsak R, Camporesi E, Hongsanan S, Monkai J, Wanasinghe DN, Tian Q, Mapook A, Chukeatirote E, Kang J-C, Xu J-C, McKenzie EHC, Jones EBG, Hyde KD (2014) A molecular phylogenetic reappraisal of the Didymosphaeriaceae (=Montagnulaceae). Fungal Diversity 68(1): 69–104. https:// doi.org/10.1007/s13225-014-0305-6
- Bánki O, Roskov Y, Döring M, Ower G, Vandepitte L, Hobern D, Remsen D, Schalk P, De Walt RE, Keping M (2023) Catalogue of Life Checklist (Version 2023-01-12). Catalogue of Life. https://doi.org/10.48580/dfqz
- Barr ME (1990) Some dictyosporous genera and species of Pleosporales in North America. Memoirs of the New York Botanical Garden 62: 1–92.
- Bissett A, Fitzgerald A, Meintjes T, Mele PM, Reith F, Dennis PG, Breed MF, Brown B, Brown MV, Brugger J, Byrne M, Caddy-Retalic S, Carmody B, Coates DJ, Correa C, Ferrari BC, Gupta VVSR, Hamonts K, Haslem A, Hugenholtz P, Karan M, Koval J, Lowe AJ, Macdonald S, McGrath L, Martin D, Morgan M, North KI, Paungfoo-Lonhienne C, Pendall E, Phillips L, Pirzl R, Powell JR, Ragan MA, Schmidt S, Seymour N, Snape I, Stephen JR, Stevens M, Tinning M, Williams K, Yeoh YK, Zammit CM, Young A (2016) Introducing BASE: The Biomes of Australian Soil Environments soil microbial diversity database. GigaScience 5(1): 1–21. https://doi.org/10.1186/s13742-016-0126-5
- Bivand R, Keitt T, Rowlingson B (2022) rgdal: Bindings for the 'Geospatial' Data Abstraction Library. R package version 1.6–2. https://CRAN.R-project.org/package=rgdal
- Boonmee S, Wanasinghe DN, Calabon MS, Huanraluek N, Chandrasiri SKU, Jones GEB, Rossi W, Leonardi M, Singh SK, Rana S, Singh PN, Maurya DK, Lagashetti AC, Choudhary D, Dai YC, Zhao CL, Mu YH, Yuan HS, He SH, Phookamsak R, Jiang HB, Martín MP, Dueñas M, Telleria MT, Kałucka IL, Jagodziński AM, Liimatainen K, Pereira DS, Phillips AJL, Suwannarach N, Kumla J, Khuna S, Lumyong S, Potter TB, Shivas RG, Sparks AH, Vaghefi N, Abdel-Wahab MA, Abdel-Aziz FA, Li GJ, Lin WF, Singh U, Bhatt RP, Lee HB, Nguyen TTT, Kirk PM, Dutta AK, Acharya K, Sarma VV, Niranjan M, Rajeshkumar KC, Ashtekar N, Lad S, Wijayawardene NN, Bhat DJ, Xu RJ, Wijesinghe SN, Shen HW, Luo ZL, Zhang JY, Sysouphanthong P, Thongklang N, Bao DF, Aluthmuhandiram JVS, Abdollahzadeh J, Javadi A, Dovana F, Usman M, Khalid AN, Dissanayake AJ, Telagathoti A, Probst M, Peintner U, Garrido-Benavent I, Bóna L, Merényi Z, Boros L, Zoltán B, Stielow JB, Jiang N, Tian CM, Shams E, Dehghanizadeh F, Pordel A, Javan-Nikkhah M, Denchev TT, Denchev CM, Kemler M, Begerow D, Deng CY, Harrower E, Bozorov T, Kholmuradova T, Gafforov Y, Abdurazakov A, Xu JC, Mortimer PE, Ren GC, Jeewon R, Maharachchikumbura SSN, Phukhamsakda C, Mapook A, Hyde KD (2021) Fungal diversity notes 1387-1511: Taxonomic and phylogenetic contributions on genera and species of fungal taxa. Fungal Diversity 111(1): 1-335. https://doi.org/10.1007/ s13225-021-00489-3
- Bundhun D, Wanasinghe DN, Maharachchikumbura SSN, Bhat DJ, Huang S-K, Lumyong S, Mortimer PE, Hyde KD (2021) *Yuxiensis granularis* gen. et sp. nov., a novel quellkörper-bearing fungal taxon added to Scortechiniaceae and inclusion of Parasympodiella-

ceae in Coronophorales based on phylogenetic evidence. Life 11(10): e1011. https://doi.org/10.3390/life11101011

Checa J (2004) Dictyosporic Dothideales. Flora Mycologica Iberica 6: 1–162.

- Crivelli PG (1983) Über die heterogene Ascomycetengattung *Pleospora* Rabh.; Vorschlag für eine Aufteilung. Diss. ETH Zürich 7318: 1–213. https://doi.org/10.3929/ethz-a-000300166
- Crous PW, Summerell BA, Shivas RG, Burgess TI, Decock CA, Dreyer LL, Granke LL, Guest DI, St J, Hardy GE, Hausbeck MK, Huberli D, Jung T, Koukol O, Lennox CL, Liew ECY, Lombard L, McTaggart AR, Pryke JS, Roets F, Saude C, Shuttleworth LA, Stukely MJC, Vanky K, Webster BJ, Windstam ST, Groenewald JZ (2012) Fungal planet description sheets: 107–127. Persoonia 28(1): 138–182. https://doi. org/10.3767/003158512X652633
- Crous PW, Wingfield MJ, Chooi YH, Gilchrist CLM, Lacey E, Pitt JI, Roets F, Swart WJ, Cano-Lira JF, Valenzuela-Lopez N, Hubka V, Shivas RG, Stchigel AM, Holdom DG, Jurjević Ž, Kachalkin AV, Lebel T, Lock C, Martín MP, Tan YP, Tomashevskaya MA, Vitelli JS, Baseia IG, Bhatt VK, Brandrud TE, De Souza JT, Dima B, Lacey HJ, Lombard L, Johnston PR, Morte A, Papp V, Rodríguez A, Rodríguez-Andrade E, Semwal KC, Tegart L, Abad ZG, Akulov A, Alvarado P, Alves A, Andrade JP, Arenas F, Asenjo C, Ballarà J, Barrett MD, Berná LM, Berraf-Tebbal A, Bianchinotti MV, Bransgrove K, Burgess TI, Carmo FS, Chávez R, Čmoková A, Dearnaley JDW, Santiago ALCMA, Freitas-Neto JF, Denman S, Douglas B, Dovana F, Eichmeier A, Esteve-Raventós F, Farid A, Fedosova AG, Ferisin G, Ferreira RJ, Ferrer A, Figueiredo CN, Figueiredo YF, Reinoso-Fuentealba CG, Garrido-Benavent I, Cañete-Gibas CF, Gil-Durán C, Glushakova AM, Gonçalves MFM, González M, Gorczak M, Gorton C, Guard FE, Guarnizo AL, Guarro J, Gutiérrez M, Hamal P, Hien LT, Hocking AD, Houbraken J, Hunter GC, Inácio CA, Jourdan M, Kapitonov VI, Kelly L, Khanh TN, Kisło K, Kiss L, Kiyashko A, Kolařík M, Kruse J, Kubátová A, Kučera V, Kučerová I, Kušan I, Lee HB, Levicán G, Lewis A, Liem NV, Liimatainen K, Lim HJ, Lyons MN, Maciá-Vicente JG, Magaña-Dueñas V, Mahiques R, Malysheva EF, Marbach PAS, Marinho P, Matočec N, McTaggart AR, Mešić A, Morin L, Muñoz-Mohedano JM, Navarro-Ródenas A, Nicolli CP, Oliveira RL, Otsing E, Ovrebo CL, Pankratov TA, Paños A, Paz-Conde A, Pérez-Sierra A, Phosri C, Pintos Á, Pošta A, Prencipe S, Rubio E, Saitta A, Sales LS, Sanhueza L, Shuttleworth LA, Smith J, Smith ME, Spadaro D, Spetik M, Sochor M, Sochorová Z, Sousa JO, Suwannasai N, Tedersoo L, Thanh HM, Thao LD, Tkalčec Z, Vaghefi N, Venzhik AS, Verbeken A, Vizzini A, Voyron S, Wainhouse M, Whalley AJS, Wrzosek M, Zapata M, Zeil-Rolfe I, Groenewald JZ (2020) Fungal planet description sheets: 1042-1111. Persoonia 44(1): 301-459. https://doi.org/10.3767/persoonia.2020.44.11
- Dissanayake LS, Maharachchikumbura SSN, Mortimer PE, Hyde KD, Kang JC (2021) Acrocordiella yunnanensis sp. nov. (Requienellaceae, Xylariales) from Yunnan, China. Phytotaxa 487(2): 103–113. https://doi.org/10.11646/phytotaxa.487.2.1
- Du T, Hyde KD, Mapook A, Mortimer PE, Xu JC, Karunarathna SC, Tibpromma S (2021) Morphology and phylogenetic analyses reveal *Montagnula puerensis* sp. nov. (Didymosphaeriaceae, Pleosporales) from southwest China. Phytotaxa 514(1): 1–25. https://doi.org/10.11646/phytotaxa.514.1.1
- Eriksson OE (1992) The Non-Lichenized Pyrenomycetes of Sweden. Btjtryck Lund, Sweden, 208 pp.

Fakirova VI (2004) New record of Bulgarian ascomycetes. Mycologia Balcanica 1: 41–43.
Felsenstein J (1985) Confidence limits on phylogenies: An approach using the bootstrap. Evolution; International Journal of Organic Evolution 39(4): 783–791. https://doi.org/10.2307/2408678

- French AM (1989) California Plant Disease Host Index. California Department of Food and Agriculture, Sacramento, 394 pp.
- Gao Y, Monkai J, Gentekaki E, Ren G-C, Wanasinghe DN, Xu J, Gui H (2021) *Dothidea kunmingensis*, a novel asexual species of Dothideaceae on *Jasminum nudiflorum* (winter jasmine) from Southwestern China. Phytotaxa 529(1): 43–56. https://doi. org/10.11646/phytotaxa.529.1.3
- Gui H, Breed M, Li Y, Xu Q, Yang J, Wanasinghe DN, Li Y, Xu J, Mortimer PM (2023) Continental-scale insights into the soil microbial co-occurrence networks of Australia and their environmental drivers. Soil Biology & Biochemistry 186: e109177. https://doi. org/10.1016/j.soilbio.2023.109177
- 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.
- Hijmans R (2023) raster: Geographic Data Analysis and Modeling. R package version 3.6–26. https://CRAN.R-project.org/package=raster

Holm L, Holm K (1993) The genus Pleospora s.1. from Svalbard. Sydowia 45: 167-187.

- Holm L, Holm K (1994) Svalbard Pyrenomycetes. An annotated checklist. Karstenia 34(2): 65–78. https://doi.org/10.29203/ka.1994.308
- Hongsanan S, Hyde KD, Bahkali AH, Camporesi E, Chomnunti P, Ekanayaka H, Gomes AAM, Hofstetter V, Jones EBG, Pinho DB, Pereira OL, Tian Q, Wanasinghe DN, Xu J-C, Buyck B (2015) Fungal biodiversity profiles 11–20. Cryptogamie. Mycologie 36(3): 355–380. https://doi.org/10.7872/crym/v36.iss3.2015.355
- Huhndorf SM (1992) Studies in *Leptosphaeria*. Transfer of *Leptosphaeria opuntiae* to *Mon-tagnula* (Ascomycetes). Brittonia 44(2): 208–212. https://doi.org/10.2307/2806834
- Hyde KD, Aptroot A, Frohlich J, Taylor JE (1999) Fungi from palms. XLII. *Didymosphaeria* and similar ascomycetes from palms. Nova Hedwigia 69(3–4): 449–471. https://doi. org/10.1127/nova.hedwigia/69/1999/449
- Hyde KD, Hongsanan S, Jeewon R, Bhat DJ, McKenzie EHC, Jones EBG, Phookamsak R, Ariyawansa HA, Boonmee S, Zhao Q, Abdel-Aziz FA, Abdel-Wahab MA, Banmai S, Chomnunti P, Cui BK, Daranagama DA, Das K, Dayarathne MC, de Silva NI, Dissanayake AJ, Doilom M, Ekanayaka AH, Gibertoni TB, Go'es-Neto A, Huang SK, Jayasiri SC, Jayawardena RS, Konta S, Lee HB, Li WJ, Lin CG, Liu JK, Lu YZ, Luo ZL, Manawasinghe IS, Manimohan P, Mapook A, Niskanen T, Norphanphoun C, Papizadeh M, Perera RH, Phukhamsakda C, Richter C, de Santiago ALCMA, Drechsler-Santos ER, Senanayake IC, Tanaka K, Tennakoon TMDS, Thambugala KM, Tian Q, Tibpromma S, Thongbai B, Vizzini A, Wanasinghe DN, Wijayawardene NN, Wu HX, Yang J, Zeng XY, Zhang H, Zhang JF, Bulgakov TS, Camporesi E, Bahkali AH, Amoozegar AM, Araujo-Neta LS, Ammirati JF, Baghela A, Bhatt RP, Bojantchev S, Buyck B, da Silva GA, de Lima CLF, de Oliveira RJV, de Souza CAF, Dai YC, Dima B, Duong TT, Ercole E, Mafalda-Freire F, Ghosh A, Hashimoto A, Kamolhan S, Kang JC, Karunarathna SC, Kirk PM, Kyto"vuori I, Lantieri A, Liimatainen K, Liu ZY, Liu XZ, Lu¨cking R, Medardi G, Mortimer PE, Nguyen TTT, Promputtha I, Raj KNA, Reck MA, Lumyong S, Shahzadeh-Fazeli SA, Stadler M, Soudi MR, Su HY, Takahashi T, Tangthirasunun N, Uniyal P, Wang Y, Wen TC, Xu JC, Zhang ZK, Zhao YC, Zhou JZ, Zhu L (2016) Fungal diversity notes 367-490: Taxonomic and phylogenetic contributions to fungal taxa. Fungal Diversity 80: 1-270. https://doi.org/10.1007/s13225-016-0373-x
- Hyde KD, Dong Y, Phookamsak R, Jeewon R, Bhat DJ, Jones EBG, Liu NG, Abeywickrama PD, Mapook A, Wei D, Perera RH, Manawasinghe IS, Pem D, Bundhun D, Karunarathna A, Ekanayaka AH, Bao DF, Li JF, Samarakoon MC, Chaiwan N, Lin CG, Phutthacharoen K, Zhang SN, Senanayake IC, Goonasekara ID, Thambugala KM, Phukhamsakda C, Tennakoon DS, Jiang HB, Yang J, Zeng M, Huanraluek N, Liu JK, Wijesinghe SN,

Tian Q, Tibpromma S, Brahmanage RS, Boonmee S, Huang SK, Thiyagaraja V, Lu YZ, Jayawardena RS, Dong W, Yang EF, Singh SK, Singh SM, Rana S, Lad SS, Anand G, Devadatha B, Niranjan M, Sarma VV, Liimatainen K, Hudson BA, Niskanen T, Overall A, Alvarenga RLM, Gibertoni TB, Pfliegler WP, Horváth E, Imre A, Alves AL, da Silva Santos AC, Tiago PV, Bulgakov TS, Wanasinghe DN, Bahkali AH, Doilom M, Elgorban AM, Maharachchikumbura SSN, Rajeshkumar KC, Haelewaters D, Mortimer PE, Zhao Q, Lumyong S, Xu JC, Sheng J (2020) Fungal diversity notes 1151–1276: Taxonomic and phylogenetic contributions on genera and species of fungal taxa. Fungal Diversity 100(1): 5–277. https://doi.org/10.1007/s13225-020-00439-5

- Hyde KD, Norphanphoun C, Ma J, Yang HD, Zhang JY, Du TY, Gao Y, Gomes de Farias AR, He SC, He YK, Li CJY, Li JY, Liu XF, Lu L, Su HL, Tang X, Tian XG, Wang SY, Wei DP, Xu RF, Xu RJ, Yang YY, Zhang F, Zhang Q, Bahkali AH, Boonmee S, Chethana KWT, Jayawardena RS, Lu YZ, Karunarathna SC, Tibpromma S, Wang Y, Zhao Q (2023) Mycosphere notes 387–412 – novel species of fungal taxa from around the world. Mycosphere : Journal of Fungal Biology 14(1): 663–744. https://doi.org/10.5943/ mycosphere/14/1/8
- Katoh K, Rozewicki J, Yamada KD (2019) MAFFT online service: Multiple sequence alignment, interactive sequence choice and visualization. Briefings in Bioinformatics 20(4): 1160–1166. https://doi.org/10.1093/bib/bbx108
- Leuchtmann A (1984) Uber *Phaeosphaeria* Miyake und andere bitunicate Ascomyceten mit mehrfach querseptierten Ascosporen. Sydowia 37: 75–194.
- Li WL, Liang RR, Dissanayake AJ, Liu JK (2023) Mycosphere Notes 413–448: Dothideomycetes associated with woody oil plants in China. Mycosphere: Journal of Fungal Biology 14(1): 1436–1529. https://doi.org/10.5943/mycosphere/14/1/16
- Liu YJ, Whelen S, Hall BD (1999) Phylogenetic relationships among ascomycetes: Evidence from an RNA polymerse II subunit. Molecular Biology and Evolution 16(12): 1799–1808. https://doi.org/10.1093/oxfordjournals.molbev.a026092
- Liu JK, Hyde KD, Gareth EBG, Ariyawansa HA, Bhat DJ, Boonmee S, Maharachchikumbura SS, Mckenzie EH, Phookamsak R, Phukhamsakda C, Shenoy BD, AbdelWahab MA, Buyck B, Chen J, Chethana KWT, Singtripop C, Dai DQ, Dai YC, Daranagama DA, Dissanayake AJ, Doilom M, D'souza MJ, Fan XL, Goonasekara ID, Hirayama K, Hongsanan S, Jayasiri SC, Jayawardena RS, Karunarathna SC, Li WJ, Mapook A, Norphanphoun C, Pang KL, Perera RH, Peršoh D, Pinruan U, Senanayake IC, Somrithipol S, Suetrong S, Tanaka K, Thambugala KM, Tian Q, Tibpromma S, Udayanga D, Wijayawardene NN, Wanasinghe D, Wisitrassameewong K, Zeng XY, Abdel-Aziz FA, Adamcík S, Bahkali AH, Boonyuen N, Bulgakov T, Callac P, Chomnunti P, Greiner K, Hashimoto A, Hofstetter V, Kang JC, Xing DL, Li H, Liu XZ, Liu ZY, Matsumura M, Mortimer PE, Rambold G, Randrianjohany E, Sato G, Sri-Indrasutdhi V, Tian CM, Verbeken A, Brackel W, Wang Y, Wen TC, Xu JC, Yan JY, Zhao RL, Camporesi E (2015) Fungal diversity notes 1–110: Taxonomic and phylogenetic contributions to fungal species. Fungal Diversity 72(1): 1–197. https://doi.org/10.1007/s13225-015-0324-y
- Lu B, Hyde KD, Ho WH, Tsui KM, Taylor JE, Wong KM Yanna, Zhou D (2000) Checklist of Hong Kong Fungi. Fungal Diversity Press, Hong Kong, 207 pp.
- Lu L, Karunarathna SC, Dai D-Q, Xiong Y-R, Suwannarach N, Stephenson SL, Elgorban AM, Al-Rejaie S, Jayawardena RS, Tibpromma S (2022) Description of four novel species in Pleosporales associated with coffee in Yunnan, China. Journal of Fungi 8(10): e1113. https://doi.org/10.3390/jof8101113
- Luis P, Saint-Genis G, Vallon L, Bourgeois C, Bruto M, Marchand C, Record E, Hugoni M (2019) Contrasted ecological niches shape fungal and prokaryotic community

structure in mangroves sediments. Environmental Microbiology 21(4): 1407–1424. https://doi.org/10.1111/1462-2920.14571

- Maharachchikumbura SSN, Chen Y, Ariyawansa HA, Hyde KD, Haelewaters D, Perera RH, Samarakoon MC, Wanasinghe DN, Bustamante DE, Liu J-K, Lawrence DP, Cheewangkoon R, Stadler M (2021a) Integrative approaches for species delimitation in Ascomycota. Fungal Diversity 109(1): 155–179. https://doi.org/10.1007/s13225-021-00486-6
- Maharachchikumbura SSN, Wanasinghe DN, Cheewangkoon R, Al-Sadi AM (2021b) Uncovering the hidden taxonomic diversity of fungi in Oman. Fungal Diversity 106(1): 229–268. https://doi.org/10.1007/s13225-020-00467-1
- Maharachchikumbura SSN, Wanasinghe DN, Elgorban AM, Al-Rejaie SS, Kazerooni EA, Cheewangkoon R (2022) *Brunneosporopsis yunnanensis* gen. et sp. nov. and *Alloc-ryptovalsa xishuangbanica* sp. nov., new terrestrial Sordariomycetes from Southwest China. Life 12(5): e635. https://doi.org/10.3390/life12050635
- Mapook A, Hyde KD, McKenzie EHC, Gareth Jones EB, Bhat DJ, Jeewon R, Stadler M, Samarakoon MC, Malaithong M, Tanunchai B, Buscot F, Wubet T, Purahong W (2020) Taxonomic and phylogenetic contributions to fungi associated with the invasive weed *Chromolaena odorata* (Siam weed). Fungal Diversity 101(1): 1–175. https://doi. org/10.1007/s13225-020-00444-8
- Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for Inference of large Phylogenetic Trees. Proceedings of the Gateway Computing Environments Workshop (GCE). New Orleans, 8 pp. https://doi.org/10.1109/GCE.2010.5676129
- Monkai J, Wanasinghe DN, Jeewon R, Promputtha I, Phookamsak R (2021) Morphological and phylogenetic characterization of fungi within Bambusicolaceae: Introducing two new species from the Greater Mekong Subregion. Mycological Progress 20(5): 721–732. https://doi.org/10.1007/s11557-021-01694-9
- Mortimer PE, Jeewon R, Xu JC, Lumyong S, Wanasinghe DN (2021) Morpho-phylo taxonomy of novel dothideomycetous fungi associated with dead woody twigs in Yunnan Province, China. Frontiers in Microbiology 12: e654683. https://doi.org/10.3389/ fmicb.2021.654683
- Mulenko W, Majewski T, Ruszkiewicz-Michalska M (2008) A preliminary checklist of Micromycetes in Poland. W. Szafer Institute of Botany Polish Academy of Sciences 9: e752.
- Niranjan M, Sarma VV (2018) Twelve new species of ascomycetous from Andaman Islands, India. Kavaka 50: 84–97.
- Pebesma E, Bivand R (2023) Spatial Data Science: With Applications in R. Chapman and Hall/CRC, 352 pp. https://doi.org/10.1201/9780429459016
- Phookamsak R, Wanasinghe DN, Hongsanan S, Phukhamsakda C, Huang S, Tennakoon DS, Norphanphoun C, Camporesi E, Bulgakov TS, Promputtha I, Mortimer PE, Xu J, Hyde KD (2017) Towards a natural classification of *Ophiobolus* and ophiobolus-like taxa; introducing three novel genera *Ophiobolopsis*, *Paraophiobolus* and *Pseudoophiobolus* in Phaeosphaeriaceae (Pleosporales). Fungal Diversity 87(1): 299–339. https://doi.org/10.1007/s13225-017-0393-1
- Pitt WM, Úrbez-Torres JR, Trouillas FP (2014) *Munkovalsaria donacina* from grapevines and Desert Ash in Australia. Mycosphere 5(5): 656–661. https://doi.org/10.5943/mycosphere/5/5/6
- R Core Team (2022) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna. https://www.R-project.org/
- Ramaley AW, Barr ME (1995) New dictyosporous species from leaves of Agavaceae. Mycotaxon 54: 75–90.

Rambaut A (2012) FigTree version 1.4.0. http://tree.bio.ed.ac.uk/software/figtree

- 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(1): 84–98. https://doi.org/10.3852/mycologia.97.1.84
- Rehner SA, Samuels GJ (1994) Taxonomy and phylogeny of *Gliocladium* analysed from nuclear large subunit ribosomal DNA sequences. Mycological Research 98(6): 625–634. https://doi.org/10.1016/S0953-7562(09)80409-7
- Ren GC, Wanasinghe DN, Monkai J, Mortimer PE, Hyde KD, Xu J-C, Pang A, Gui H (2021a) Novel saprobic *Hermatomyces* species (Hermatomycetaceae, Pleosporales) from China (Yunnan Province) and Thailand. MycoKeys 82: 57–79. https://doi. org/10.3897/mycokeys.82.67973
- Ren GC, Wanasinghe DN, Monkai J, Hyde KD, Mortimer PE, Xu J, Pang A, Gui H (2021b) Introduction of *Neolophiotrema xiaokongense* gen. et sp. nov. to the poorly represented Anteagloniaceae (Pleosporales, Dothideomycetes). Phytotaxa 482(1): 25–35. https://doi.org/10.11646/phytotaxa.482.1.3
- Ren GC, Wanasinghe DN, De Farias ARG, Hyde KD, Yasanthika E, Xu JC, Balasuriya A, Chethana KWT, Gui H (2022a) Taxonomic novelties of woody litter fungi (Didymosphaeriaceae, Pleosporales) from the Greater Mekong Subregion. Biology 11(11): e1660. https://doi.org/10.3390/biology11111660
- Ren GC, Wanasinghe DN, Jeewon R, Monkai J, Mortimer PE, Hyde KD, Xu J-C, Gui H (2022b) Taxonomy and phylogeny of the novel rhytidhysteron-like collections in the Greater Mekong Subregion. MycoKeys 86: 65–85. https://doi.org/10.3897/mycokeys.86.70668
- Ronquist F, Huelsenbeck J, Teslenko M (2011) MrBayes Version 3.2 Manual: Tutorials and Model Summaries. https://brahms.biology.rochester.edu/software.html [Accessed on 10 June 2023]
- Scheuer C (1988) Ascomyceten auf Cyperaceen und Juncaceen im Ostalpenraum. Bibliotheca Mycologica 123: 1–274.
- Senwanna C, Mapook A, Samarakoon MC, Karunarathna A, Wang Y, Tang AMC, Haituk S, Suwannarach N, Hyde KD, Cheewangkoon R (2021) Ascomycetes on Para rubber (*Hevea brasiliensis*). Mycosphere 12(1): 1334–1512. https://doi.org/10.5943/myco-sphere/12/1/18
- Shoemaker RA, Babcock CE (1989) Phaeosphaeria. Canadian Journal of Botany 67(5): 1500–1599. https://doi.org/10.1139/b89-199
- Stamatakis A (2014) RAxML Version 8: A tool for Phylogenetic Analysis and Post-Analysis of Large Phylogenies. Bioinformatics 30(9): 1312–1313. https://doi.org/10.1093/ bioinformatics/btu033
- Sun Y-R, Zhang J-Y, Hyde KD, Wang Y, Jayawardena RS (2023) Morphology and phylogeny reveal three *Montagnula* species from China and Thailand. Plants 12(4): e738. https://doi.org/10.3390/plants12040738
- Sung GH, Sung JM, Hywel-Jones NL, Spatafora JW (2007) A multi-gene phylogeny of Clavicipitaceae (Ascomycota, Fungi): Identification of localized incongruence using a combinational bootstrap approach. Molecular Phylogenetics and Evolution 44(3): 1204–1223. https://doi.org/10.1016/j.ympev.2007.03.011
- Tedersoo L, Bahram M, Põlme S, Kõljalg U, Yorou NS, Wijesundera R, Villarreal Ruiz L, Vasco-Palacios AM, Thu PQ, Suija A, Smith ME, Sharp C, Saluveer E, Saitta A, Rosas M, Riit T, Ratkowsky D, Pritsch K, Põldmaa K, Piepenbring M, Phosri C, Peterson M, Parts K, Pärtel K, Otsing E, Nouhra E, Njouonkou AL, Nilsson RH, Morgado LN, Mayor J, May TW, Majuakim L, Lodge DJ, Lee SS, Larsson KH, Kohout P, Hosaka K, Hiiesalu I, Henkel TW, Harend H, Guo LD, Greslebin A, Grelet G, Geml J, Gates G, Dunstan W, Dunk C, Drenkhan

R, Dearnaley J, De Kesel A, Dang T, Chen X, Buegger F, Brearley FQ, Bonito G, Anslan S, Abell S, Abarenkov K (2014) Fungal biogeography. Global diversity and geography of soil fungi. Science 346(6213): e1256688. https://doi.org/10.1126/science.1256688

- Tennakoon DS, Hyde KD, Wanasinghe DN, Bahkali AH, Camporesi E, Khan S, Phookamsak R (2016) Taxonomy and phylogenetic appraisal of *Montagnula jonesii* sp. nov. (Didymosphaeriaceae, Pleosporales). Mycosphere 7(9): 1346–1356. https://doi. org/10.5943/mycosphere/7/9/8
- Tennakoon DS, Thambugala KM, de Silva NI, Suwannarach N, Lumyong S (2022) A taxonomic assessment of novel and remarkable fungal species in Didymosphaeriaceae (Pleosporales, Dothideomycetes) from plant litter. Frontiers in Microbiology 13: e1016285. https://doi.org/10.3389/fmicb.2022.1016285
- Thaung MM (2008) Pathologic and taxonomic analysis of leaf spot and tar spot diseases in a tropical dry to wet monsoon ecosystem of lowland Burma. Australasian Plant Pathology 37(2): 180–197. https://doi.org/10.1071/AP08007
- Thiyagaraja V, Senanayake IC, Wanasinghe DN, Karunarathna SC, Worthy FR, To-anun C (2019) Phylogenetic and morphological appraisal of *Diatrype lijiangensis* sp. nov. (Diatrypaceae, Xylariales) from China. Asian Journal of Mycology 2(1): 198–208. https://doi.org/10.5943/ajom/2/1/10
- Thiyagaraja V, Hyde KD, Wanasinghe DN, Worthy FR, Karunarathna SC (2020) Addition to Melanommataceae: A new geographical record of *Alpinaria rhododendri* from Shangri La, China. Asian Journal of Mycology 3(1): 335–344. https://doi.org/10.5943/ajom/3/1/8
- Thiyagaraja V, Wanasinghe DN, Karunarathna SC, Tennakoon DS, Hyde KD, To-Anun C, Cheewangkoon R (2021) *Alloleptosphaeria shangrilana* sp. nov. and first report of the genus (Leptosphaeriaceae, Dothideomycetes) from China. Phytotaxa 491(1): 12–22. https://doi.org/10.11646/phytotaxa.491.1.2
- Tibpromma S, Hyde KD, McKenzie EH, Bhat DJ, Phillips AJ, Wanasinghe DN, Samarakoon MC, Jayawardena RS, Dissanayake AJ, Tennakoon DS, Doilom M, Phookamsak R, Tang AM, Xu J, Mortimer PE, Promputtha I, Maharachchikumbura SSN, Khan S, Karunarathna SC (2018) Fungal diversity notes 840–928: Micro-fungi associated with Pandanaceae. Fungal Diversity 93(1): 1–160. https://doi.org/10.1007/s13225-018-0408-6
- Trifinopoulos J, Nguyen LT, von Haeseler A, Minh BQ (2016) W-IQ-TREE: A fast online phylogenetic tool for maximum likelihood analysis. Nucleic Acids Research 44(W1): W232–W235. https://doi.org/10.1093/nar/gkw256
- Turner BL, Zemunik G, Laliberté E, Drake JJ, Jones FA, Saltonstall K (2019) Contrasting patterns of plant and microbial diversity during long-term ecosystem development. Journal of Ecology 107(2): 606–621. https://doi.org/10.1111/1365-2745.13127
- Větrovský T, Morais D, Kohout P, Lepinay C, Algora C, Awokunle Hollá S, Bahnmann BD, Bílohnědá K, Brabcová V, D'Alò F, Human ZR, Jomura M, Kolařík M, Kvasničková J, Lladó S, López-Mondéjar R, Martinović T, Mašínová T, Meszárošová L, Michalčíková L, Michalová T, Mundra S, Navrátilová D, Odriozola I, Piché-Choquette S, Štursová M, Švec K, Tláskal V, Urbanová M, Vlk L, Voříšková J, Žifčáková L, Baldrian P (2020) GlobalFungi, a global database of fungal occurrences from high-throughput-sequencing metabarcoding studies. Scientific Data 7(1): e228. https://doi.org/10.1038/s41597-020-0567-7
- Vilgalys R, Hester M (1990) Rapid genetic identification and mapping of enzymatically amplified ribosomal DNA from several *Cryptococcus* species. Journal of Bacteriology 172(8): 4238–4246. https://doi.org/10.1128/jb.172.8.4238-4246.1990
- Wanasinghe DN, Mortimer PE (2022) Taxonomic and phylogenetic insights into novel Ascomycota from forest woody litter. Biology 11(6): e889. https://doi.org/10.3390/ biology11060889

- Wanasinghe DN, Jones EB, Camporesi E, Dissanayake AJ, Kamolhan S, Mortimer PE, Xu J, Abd-Elsalam KA, Hyde KD (2016) Taxonomy and phylogeny of *Laburnicola* gen. nov. and *Paramassariosphaeria* gen. nov. (Didymosphaeriaceae, Massarineae, Pleosporales). Fungal Biology 120(11): 1354–1373. https://doi.org/10.1016/j.funbio.2016.06.006
- Wanasinghe DN, Jeewon R, Tibpromma S, Jones EBG, Hyde KD (2017) Saprobic Dothideomycetes in Thailand: *Muritestudina* gen. et sp. nov. (Testudinaceae) a new terrestrial pleosporalean ascomycete, with hyaline and muriform ascospores. Studies in Fungi 2(1): 219–234. https://doi.org/10.5943/sif/2/1/26
- Wanasinghe DN, Jeewon R, Peršoh D, Jones EBG, Camporesi E, Bulgakov TS, Gafforov YS, Hyde KD (2018a) Taxonomic circumscription and phylogenetics of novel didymellaceous taxa with brown muriform spores. Studies in Fungi 3(1): 152–175. https:// doi.org/10.5943/sif/3/1/17
- Wanasinghe DN, Phukhamsakda C, Hyde KD, Jeewon R, Lee HB, Jones EBG, Tibpromma S, Tennakoon DS, Dissanayake AJ, Jayasiri SC, Gafforov Y, Camporesi E, Bulgakov TS, Ekanayake AH, Perera RH, Samarakoon MC, Goonasekara ID, Mapook A, Li W, Senanayake IC, Li J, Norphanphoun C, Doilom M, Bahkali AH, Xu J, Mortimer PE, Tibell L, Tibell S, Karunarathna SC (2018b) Fungal diversity notes 709–839: Taxonomic and phylogenetic contributions to fungal taxa with an emphasis on fungi on Rosaceae. Fungal Diversity 89(1): 1–236. https://doi.org/10.1007/s13225-018-0395-7
- Wanasinghe DN, Wijayawardene NN, Xu J, Cheewangkoon R, Mortimer PE (2020) Taxonomic novelties in *Magnolia*-associated pleosporalean fungi in the Kunming Botanical Gardens (Yunnan, China). PLOS ONE 15(7): e0235855. https://doi.org/10.1371/ journal.pone.0235855
- Wanasinghe DN, Mortimer PE, Xu J (2021) Insight into the systematics of microfungi colonizing dead woody twigs of *Dodonaea viscosa* in Honghe (China). Journal of Fungi 7(3): e180. https://doi.org/10.3390/jof7030180
- Wanasinghe DN, Mortimer PE, Bezerra JDP (2022a) Editorial: Fungal Systematics and Biogeography. Frontiers in Microbiology 12: e827725. https://doi.org/10.3389/ fmicb.2021.827725
- Wanasinghe DN, Ren GC, Xu JC, Cheewangkoon R, Mortimer PE (2022b) Insight into the taxonomic resolution of the pleosporalean species associated with dead woody litter in natural forests from Yunnan, China. Journal of Fungi 8(4): e375. https://doi. org/10.3390/jof8040375
- Wanasinghe DN, Bezerra JDP, Xu J, Mortimer PE (2023) *Honghemyces pterolobii*, gen. et sp. nov. (Bezerromycetaceae, Tubeufiales), a new ascomycetous fungus from *Pterolobium macropterum* in Honghe, China. Sydowia 75: 13–21. https://doi. org/10.12905/0380.sydowia75-2022-0013
- Wang YZ, Aptroot A, Hyde KD (2004) Revision of the Ascomycete Genus *Amphisphaeria*. Fungal Diversity Press, Hong Kong, 168 pp.
- White TJ, Bruns T, Lee S, Taylor JW (1990) Amplification and Direct Sequencing of Fungal Ribosomal RNA Genes for Phylogenetics. PCR Protocols: A Guide to Methods and Applications. Academic Press, San Diego, 315–322. https://doi.org/10.1016/B978-0-12-372180-8.50042-1
- Wickham H (2011) ggplot2. Wiley Interdisciplinary Reviews: Computational Statistics 3(2): 180–185. https://doi.org/10.1002/wics.147
- Wijayawardene NN, Phillips AJL, Pereira DS, Dai DQ, Aptroot A, Monteiro JS, Druzhinina IS, Cai F, Fan X, Selbmann L, Coleine C, Castañeda-Ruiz RF, Kukwa M, Flakus A, Fiuza PO, Kirk PM, Kumar KCR, Arachchi ISL, Suwannarach N, Tang LZ, Boekhout T, Tan

CS, Jayasinghe RPPK, Thines M (2022) Forecasting the number of species of asexually reproducing fungi (Ascomycota and Basidiomycota). Fungal Diversity 114(1): 463–490. https://doi.org/10.1007/s13225-022-00500-5

- Yasanthika E, Dissanayake LS, Wanasinghe DN, Karunarathna SC, Mortimer PE, Samarakoon BC, Monkai J, Hyde KD (2020) *Lonicericola fuyuanensis* (Parabambusicolaceae) a new terrestrial pleosporalean ascomycete from Yunnan Province China. Phytotaxa 449(2): 103–113. https://doi.org/10.11646/phytotaxa.446.2.3
- Zhao ZZ, Zhao K, Chen HP, Bai X, Zhang L, Liu JK (2018) Terpenoids from the mushroom-associated fungus *Montagnula donacina*. Phytochemistry 147: 21–29. https:// doi.org/10.1016/j.phytochem.2017.12.015

Zhuang W-Y (2001) Higher Fungi of Tropical China. Mycotaxon Ltd, Ithaca, 485 pp.

Supplementary material 1

The biogeography, substrate and habitat affinity of *Montagnula* inferred from the GlobalFungi database

Authors: Dhanushka N. Wanasinghe, Thilina S. Nimalrathna, Li Qin Xian, Turki KH. Faraj, Jianchu Xu, Peter E. Mortimer

Data type: xlsx

Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: https://doi.org/10.3897/mycokeys.101.113259.suppl1