Morpho-molecular taxonomic studies reveal a high number of endophytic fungi from *Magnolia candolli* and *M. garrettii* in China and Thailand

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Abstract

Endophytic fungi are internal inhabitants of plant tissues that do not apparently cause harm to the host. Ecologically they provide a number of benefits to the plants by decreasing herbivory, increasing drought and disease resistance and enhancing the growth of plants. Endophytes have emerged as an exciting research topic as they have the potential to provide numerous metabolites with different biological activities. This study is focused on taxonomic novelties and new host or geographical records of endophytic fungi associated with *Magnolia candolli* collected from Yunnan Province, China and *M. garrettii* from Chiang Mai Province, Thailand. *Magnolia* plants are economically important and used in furniture, ornamental plants in gardens, temple trees, flowers for decorations and valuable medicine in China. In this study, 56 fungal endophytic isolates were obtained from *Magnolia* species, of which 54 belong to ascomycetes and two to basidiomycetes. The 56 endophytic fungal isolates were identified in 31 taxa that are distributed in eight orders, ten
families, 13 genera, including eight new species (*Colletorrichum chiangmaiense, C. xishuangbananense, Coprinellus magnolia, Diaporthe chinensis, Epicoccum endophyticum, Letendraea magnoliae, Nigrospora magnoliae and Pestalotiopsis endophytica*) and 23 new host and or geographical records. The results indicate that members of Sordariomycetes are dominant groups of endophytic fungi in Magnolia candolli and M. garrettii. Considering the total fungal endophytic isolates from Magnolia candolli and M. garrettii, Sordariomycetes comprises the the highest number of isolates of 82%, following Dothideomycetes 14% and Agaricomycetes 4%. Detailed morphological descriptions, micrographs and phylogenetic analyses are provided to show the placement of the novel taxa.

**Key words** – Ascomycota – Basidiomycota – Dothideomycetes – Multi-locus sequence analysis – Sordariomycetes – Taxonomy

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Introduction

Over the past decade, endophytes have emerged as a compelling research topic as they are considered as the hidden component of fungal diversity that have the potential to produce metabolites with different kinds of biological activities (Huang et al. 2009, Ek-Ramos et al. 2013, Zheng et al. 2016, de Silva et al. 2019a, Rashmi et al. 2019). De Bary (1886) introduced the term ‘endophytes’ as any organism inhabiting a living plant tissue. Petrini (1991) stated “endophytes” as “all organisms inhabiting plant organs at some time in their life that can colonize internal plant tissues without causing apparent harm to the host”. The universal presence of endophytes in diverse flora is facilitated by their asymptomatic colonization of various above and below ground tissues of liverworts, mosses, ferns and spermatophytes (Potshangbam et al. 2017, de Silva et al. 2019a, Rashmi et al. 2019). Fungal endophytes are symbiont in natural flora (Rodriguez et al. 2009, Rashmi et al. 2019). These mycobionts ecologically benefit their host plants by increasing drought and disease resistance and enhancing growth (Rodriguez et al. 2009, de Silva et al. 2019a, Rashmi et al. 2019). Arnold et al. (2003) showed that endophytic fungi (*Colletotrichum* sp., *Fusarium* sp., *Nectria* sp. and *Xylaria* sp.) significantly decreased both leaf necrosis and leaf mortality caused by a major pathogen, *Phytophthora* in host plant *Theobroma cacao*. This could be due to creating a barrier by endophytes that prevents pathogenic microorganisms from colonizing the same host plant (Bamisile et al. 2018). In another study by Zhang et al. (2017), greenhouse experiments were conducted to assess the ability of drought tolerance of dark septate endophytes in sorghum seedlings (*Sorghum bicolor*). Endophytic *Exophiala pisciphila* isolated from the roots of sorghum was used as the inoculum. *Exophiala pisciphila* inoculated sorghum seedlings showed a greater growth performance, photosynthetic capacity and secondary metabolism as compared to non-inoculated seedlings under drought-stressed conditions (Zhang et al. 2017). These critical findings can be beneficial to optimize the harvest of crops and utilize fungal endophytes for pest and disease management programs (Bamisile et al. 2018). Endophytes can also serve as storehouse of bioactive secondary metabolites, such as alkaloids, saponins, tannins, phenolic acids, steroids, quinones and terpenoids (Gouda et al. 2016, Bamisile et al. 2018) and consist of important properties such as insect antagonist, antimicrobial and anticancer properties (Bamisile et al. 2018). Endophytes can be referred to as biofertilizers because they serve as plant growth promoters that facilitate nutrient uptake through plant roots and also insect-derived transformation of nitrogen to plants (Bamisile et al. 2018).

The number of fungal endophytes has been estimated at 7% of the 1.5 million species of total fungi (Hawksworth 2000, Chowdhary & Kaushik 2015, Tibpromma et al. 2018a). Hawksworth & Lucking (2017) replaced previous estimates of global fungal species richness of 1.5 million to an updated range of 2.2 to 3.8 million. Hyde et al. (2020a) have stated that the estimated fungal species number can be ranging from 0.5 to 13.2 million. This recent estimate might increase the number of endophytic fungi associated with plants. It is suggested that a single tropical leaf may harbour approximately 90 endophytic species, and 50 distinct genera in a grassland species (Porras-Alfaro et al. 2008, Rashmi et al. 2019). The diversity of endophytes inhabiting approximately 300,000 plant species on earth is roughly estimated as about one million taxa based on 1:4 or 1:5 fungi per host (Rashmi et al. 2019). The diversity of endophytes can also be influenced by the sample size and geographical location (Walther & Moore 2005, Rashmi et al. 2019). Sample size can be determined by the number of asymptomatic leaves (or twigs or any relevant substrate).
selected for the isolation of endophytes. In general, sample size can be increased by selecting leaves from different individual plants from the same host in the same geographical location and or different geographical location. Endophytic fungal diversity in various geographical locations had revealed a correlation with environmental parameters (temperature and rainfall) (Zimmerman & Vitousek 2012, Rashmi et al. 2019). Small sample size from a particular host plant species in one geographical location might result in low endophytic diversity compared to a larger sample size of the same host plant species from different geographical locations (Rashmi et al. 2019). A comprehensive study by López-González et al. (2017) evaluated the effect of leaf age on the colonization frequency, species richness and diversity of the endophytic community of foliar fungal endophytes associated with lima bean plants. They showed that the richness and diversity of the foliar endophytic fungal community increased with the age of the leaves of lima bean under natural conditions. These facts reported that the diversity of endophytic fungal communities could be varied with distinct geographic locations, hosts and different sampling strategies.

In general, this diverse and polyphyletic group of endophytic microorganisms are classified into two groups, the clavicipitaceous (C) and the non-clavicipitaceous (NC) based on different criteria including, evolutionary relatedness, taxonomy, host plant range and ecological functions of fungi (Rodriguez et al. 2009, Santangelo et al. 2015, de Silva et al. 2019a). Clavicipitaceous endophytes (family Clavicipitaceae), i.e. Atkinsonella, Balansia, Balansiospis, Echinodothis, Epichloe, Myriogenospora, Neotyphodium and Parepichloe species are mutualists that rely on plants in family Poaceae throughout their fungal life cycle (Rodriguez et al. 2009, Hardoim et al. 2015, de Silva et al. 2016, 2019a). Non-clavicipitaceous endophytes, i.e. Colletotrichum sp., Fusarium sp., Phomopsis sp., and Xylaria sp. occur in most terrestrial plants and might not inhabit for their entire life cycle inside the host (Petrini et al. 1992, Promputtha et al. 2005, 2007, Rodriguez et al. 2009, Jayawardena et al. 2016, de Silva et al. 2016, 2019a). Non-clavicipitaceous endophytes might be able to switch their life mode between pathogenic and saprobic when environmental conditions become unfavourable to the host (Promputtha et al. 2005, 2007, Delaye et al. 2013, de Silva et al. 2016, 2019a).

Identification of endophytes is mainly based on cultural procedures through different *in vitro* techniques (Promputtha et al. 2005, Wang et al. 2005, Ko et al. 2011, de Silva et al. 2019a). Most of the studies have conducted surface sterilization of fresh, asymptomatic plant tissues using ethanol, NaClO solution and sterile distilled water. At the same time, isolation of fungi was done on different artificial media (water agar (WA), potato dextrose agar (PDA)) (Promputtha et al. 2005, de Silva et al. 2019a). Molecular techniques advance the identification and classification of endophytes that were initially based on the cultural approach (Lacap et al. 2003, Promputtha et al. 2005, Ko et al. 2011, Doilom et al. 2017). DNA sequence data is a reliable molecular tool to identify sporulating and non-sporulating endophytes up to species level (Lacap et al. 2003, Promputtha et al. 2005, 2007, Doilom et al. 2017, Tibpromma et al. 2018a).

Magnoliaceae is an ancient and important group of flowering plants that is widely distributed in temperate and tropical South East and East Asia and One of the largest genera in Magnoliaceae is *Magnolia* that has early divergent flowering plants, bearing a number of large odoriferous flowers with chamber blossoms (Nootboon & Chalermglin 2009, Wang et al. 2017a). *Magnolia* plant species are important for furniture making (e.g. wood of *Magnolia champaca*) and as a valuable drug in China (e.g. *Magnolia officinalis*) (Nootboon & Chalermglin 2009). Besides, many *Magnolia* species and their hybrids are cultivated as ornaments in gardens, as temple trees, and the flowers are used for decorations (Nootboon & Chalermglin 2009). Nevertheless, *Magnolia* habitats have been badly degraded and fragmented due to heavy habitat destruction in the past, resulting in most (sub) populations being isolated from each other (Wang et al. 2017a). Therefore, it is important to explore the diversity of endophytes from tropical florals that are facing deforestation before their values could be permanently disappeared from nature (Bamisile et al. 2018). Few previous studies have investigated the diversity of endophytic communities from fresh leaves of *Magnolia garrettii* in Thailand based on ITS sequence data (Promputtha et al. 2005, 2007,
In this study, we present taxonomic novelties of fungal endophytes on *Magnolia candolli* Yunnan Province, China and *M. garrettii* in Chiang Mai Province, Thailand.

**Materials & Methods**

Fresh leaves of *Magnolia candolli* were collected in Xishuangbanna Tropical Botanical Garden, Yunnan Province, China in April 2017 while fresh leaves of *M. garrettii* were collected in Chiang Mai, Thailand in September 2017. The leaves were kept at 4°C in sterile polyethylene bags until they are processed in the laboratory. Isolation of endophytes was done according to the methods described by Promputtha et al. (2007) with some modifications. Leaves were first washed using tap water and cut into small pieces (5 × 5 mm²). The cut pieces were soaked in distilled water for 1 minute and surface sterilized by dipping in 70% alcohol, followed by 2% NaOCl for 30 s. The pieces of leaves were washed thoroughly with sterile distilled water, air dried and plated on PDA. The PDA plates were incubated in ambient light at 25°C. Growing hyphal tips from the leaf pieces were aseptically transferred to fresh PDA plates and incubated in ambient light at 25°C. Finally, the fungi were isolated into pure culture and grouped according to their culture morphology.

One month old cultures on PDA and or WA were used to prepare dry fungal cultures. A solution was prepared by adding 2 g of agar powder into 200 ml of distilled water containing 10 ml of Glycerol. The solution was heated until the agar melted and kept for 5 to 10 minutes to cool. The solution was then poured on the fungal culture (removed from the original Petri plate and placed on an aseptic surface) and kept to air dry at room temperature (25°C).

The specimens (dried cultures) cited in this paper were deposited at the Mae Fah Luang University Herbarium (Herb. MFLU), Chiang Rai, Thailand. The living fungal cultures recovered in this study were deposited at Mae Fah Luang University Culture Collection (MFLUCC) and Kunming Institute of Botany Culture Collection (KUMCC). Faces of Fungi numbers and Index Fungorum numbers were registered as described in Jayasiri et al. (2015) and Index Fungorum (2021), respectively.

**DNA extraction and PCR amplification**

One-week old pure cultures on PDA were used for DNA extraction (Dissanayake et al. 2020). The mycelia was scraped off from pure cultures and genomic DNA was extracted using a Biospin fungus genomic DNA kit (BioFlux®, P.R. China) following the manufacturer’s protocol. DNA was kept at 4 °C for DNA amplification and maintained at −20°C for long term storage.

Polymerase chain reaction (PCR) was used to amplify partial gene regions of Internal Transcribed Spacers (ITS) and 28S ribosomal RNA (LSU), 18S ribosomal RNA (SSU), RNA polymerase II second largest subunit (RPB2), β-tubulin (*tub2*), Actin (ACT), Glyceraldehyde-3-Phosphate Dehydrogenase (GADPH), Chitin synthase 1 (CHS–1), Calmodulin (CAL) and Translation Elongation Factor 1-alpha (*tef1*) where appropriate using primers as shown in Table 1. The final volume of the PCR reaction was 25 μl, containing 1 μl of DNA template, 1 μl of each forward and reward primers, 12.5 μl of 2×Easy Taq PCR SuperMix (a mixture of EasyTaq TM DNA Polymerase, dNTPs, and optimized buffer, Beijing TransGen Biotech Co., Ltd., Beijing, P.R. China) and 9.5 μl of ddH₂O. Amplification of PCR reactions were performed following Li et al. (2020) for ITS, LSU, SSU, *tef1*, RPB2, *tub2*, Gomes et al. (2013) for CAL and Weir et al. (2012) for ACT, GADPH, CHS–1. PCR purification and sequencing of amplified PCR products were carried out at Shanghai Sangon Biological Engineering Technology & Services Co., Ltd, P.R. China.

**Table 1** Details of genes/loci with PCR primers and references

<table>
<thead>
<tr>
<th>Loci</th>
<th>Primer pairs (Forward/Reverse)</th>
<th>References</th>
</tr>
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<tbody>
<tr>
<td>LSU</td>
<td>LROR 5′–ACCCGCTGAACCTAAGC–3′</td>
<td>Vilgalys &amp; Hester (1990)</td>
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<td></td>
<td>LR5 5′–ATCTTGAGGGAAACTTC–3′</td>
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<tr>
<td>ITS</td>
<td>ITS5 5′–GGAAGTAAAGTGTAGAACAAGG–3′</td>
<td>White et al. (1990)</td>
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<td></td>
<td>ITS4 5′–TCCTCCGCTTATTGATATGC–3′</td>
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Table 1 Continued.

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<th>Loci</th>
<th>Primer pairs (Forward/Reverse)</th>
<th>References</th>
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<tbody>
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<td>SSU</td>
<td>NS1 5′–GTAGTCATATGCTTGCTC–3′</td>
<td>White et al. (1990)</td>
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<td></td>
<td>NS4 5′–CTCTCGTCAATTCTTTAAAG–3′</td>
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<tr>
<td>tefl</td>
<td>983F 5′–GCTCCGATGAC RCAGG AGRTGTTG–3′</td>
<td>Rehner (2001)</td>
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<tr>
<td></td>
<td>728F 5′–TCTTGCCTGGC AGTTGA AAGGC–3′</td>
<td>Carbone &amp; Kohn (1999)</td>
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<td></td>
<td>986R 5′–TAC TGTGGATGAA CCC TTA C–3′</td>
<td>Rehner (2001)</td>
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<td></td>
<td>526F 5′–GTCGCTGCGGTTCGTTG–3′</td>
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<tr>
<td></td>
<td>1567R 5′–GCCGTCGCTGCGGTTCGTTG–3′</td>
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<tr>
<td>RPB2</td>
<td>fRPB2–5f 5′–GGG GWG AYC AGA AGA AGG C</td>
<td>Liu et al. (1999)</td>
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<tr>
<td></td>
<td>fRPB2–7cR 5′–GGG GWG AYC AGA AGA AGG C</td>
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<td>CAL</td>
<td>228F 5′–GAG TTC AAG GAG GCC TTC TCC C–3′</td>
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<td></td>
<td>737R 5′–CAT CTT TCT GGC CAT C CAT GG–3′</td>
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<tr>
<td>tub2</td>
<td>BT2a 5′–GGTAACCAAATCGGTGCTGCTTT–3′</td>
<td>Glass &amp; Donaldson (1995)</td>
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<td>BT2b 5′–ACCCCTCAGTGCTCTGCCTTTCC–3′</td>
<td>O’Donnell &amp; Cigelnik (1997)</td>
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<td>T1 5′–AAC ATG CGT GAG ATT GTA AGT–3′</td>
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<td>T2 5′–TAG TGA CCC TGG GCC CAG TGG–3′</td>
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<td>ACT</td>
<td>512F 5′–ATG TGC AAG GCC GGT TTC GC–3′</td>
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<td>783R 5′–TAC GAG TCC TTC TGG CCC AT–3′</td>
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<td>CHS–1</td>
<td>79F 5′–TGG GCC AAG GAT GCT TGG AAG AAG–3′</td>
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<td>354R 5′–TGG AAC AAG CAT CTG TGA GAG TTG–3′</td>
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<td>GADPH</td>
<td>Gpd1 5′–ATT GGC CGC ATC GTC TTC–3′</td>
<td>Myllys et al. (2002)</td>
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<td>Gpd2 5′–CCC ACT CGT TGT CGT ACC–3′</td>
<td>Templeton et al. (1992)</td>
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<td>GDF 5′–GCC GTC AAC GAC CCC TTC ATT GA–3′</td>
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<td></td>
<td>GDR 5′–GGG TGG AGT CGT ACT TGA GCA TGT–3′</td>
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Newly generated nucleotide sequences were deposited in GenBank and accession numbers were mentioned in relevant entries. Sequences of the individual loci were aligned with MAFFT v. 7 online version (Yamada et al. 2016) using default settings. BioEdit v. 7.0.5.2 (Hall 1999) software was used to refine the alignments manually where necessary and to exclude incomplete portions at the ends of the sequences before the analyses.

Phylogenetic analyses

Maximum likelihood analysis was performed with RAxML GUI v. 1.3 (Silvestro & Michalak 2012) and maximum parsimony analysis was done with PAUP (Phylogenetic Analysis Using Parsimony) v. 4.0b10 (Swofford 2002). Evolutionary models for phylogenetic analyses were selected independently for each locus using MrModeltest v. 3.7 (Posada & Crandall 1998) under the Akaike Information Criterion (AIC). Parameters for maximum likelihood were set to rapid bootstrapping and the analysis carried out using 1000 replicates using the GTR + GAMMA model of nucleotide substitution. Bayesian analysis was conducted with MrBayes v. 3.1.2 (Huelsenbeck & Ronquist 2001). Parameters of Bayesian analysis in MrBayes v. 3.2; markov chains were run for 1 000 000 generations and trees were sampled every 100th generations (printfreq = 100) and 10 000 trees were obtained. Initial trees were discarded (20% burn-in value) and remaining trees were used to evaluate posterior probabilities (PP) in the majority rule consensus tree. Parameters for maximum likelihood were set to rapid bootstrapping and the analysis carried out using 1000 replicates using the GTR + GAMMA model of nucleotide substitution. Maximum parsimony was run with the heuristic search option, random taxon addition, tree bisection-reconnection (TBR) for the branch swapping algorithm and 1000 random sequence additions, with maxtrees set at 1000. Gaps were treated as missing data. Tree Length [TL], Consistency Index [CI], Retention Index [RI], Relative Consistency Index [RC] and Homoplasy Index [HI] were calculated for the most parsimonious tree. Phylogenograms were visualized with FigTree v1.4.0 (Rambaut 2012) and annotated in Microsoft PowerPoint (2010). We conducted above two or 3 different analyses to obtain phylogenetic support and discussed results in respective entries.
Genealogical concordance phylogenetic species recognition analysis

Two new species of *Colletotrichum* and their most closely related species were analysed using the GCPSR model. A pairwise homoplasy index (PHI) (Philippe & Bryant 2006) test was performed in SplitsTree4 (Huson 1998, Huson & Bryant 2006) as described by Quaedvlieg et al. (2014), in order to determine the recombination level within phylogenetically closely related species using a five-locus concatenated dataset for Two new species of *Colletotrichum*. If the pairwise homoplasy index is below a 0.05 threshold (Φw < 0.05), it indicated that there is significant recombination present in the dataset. The relationships between closely related species were visualised by constructing a split graph, using both the LogDet transformation and splits decomposition options (Figs 23, 26).

Results

**Phylum Ascomycota** Caval. Sm.

**Class Dothideomycetes** O.E. Erikss. & Winka

**Subclass Dothideomycetidae** P.M. Kirk

**Cladosporiales** Abdollahz. & Crous

**Cladosporiaceae** Nann.

The family includes saprobic, pathogenic or endophytic and pathogenic species that are involved in plant, human and animal diseases (Hyde et al. 2013). Cladosporiaceae comprises nine genera (Hongsanan et al. 2020). The asexual morph of *Cladosporium* is characterized by solitary to fasciculate conidiophores containing unbranched or branched acropetal conidial chains (Sandoval-Denis et al. 2016). The sexual morph is characterized by pseudothecial ascomata, 8-spored obovoid to subcylindrical asci, and hyaline, obovoid to ellipsoid ascospores (Schubert et al. 2007, Sandoval-Denis et al. 2016).

**Cladosporium** Link

*Cladosporium* consists 170 species that are recognized as true *Cladosporium* in a monographic treatment (Bensch et al. 2018), with a worldwide distribution and isolated from a wide range of substrates (Bensch et al. 2012, 2015, Crous et al. 2014, Sandoval-Denis et al. 2016). Index Fungorum (2021) lists 855 epithets of *Cladosporium*. Species of *Cladosporium* are saprobes, endophytes and pathogens causing leaf spots on *Capsicum annuum*, *Populus tremuloides*, *Quercus robur*, *Yucca elephantipes*, or leaf lesions on *Iris* sp. as well as they are hyperparasites on other fungi (Bensch et al. 2012, Sandoval-Denis et al. 2016). There are three species complexes in *Cladosporium* mainly based on morphology; *Cladosporium cladosporioides*, *C. herbarum* and *C. sphaerospermum* species complexes (Bensch et al. 2010, 2018, Sandoval-Denis et al. 2016). Schubert et al. (2007) have performed comprehensive molecular analyses for the *C. herbarum* complex based on five genes, viz., rDNA ITS, ACT, CAL, *tef1* and histone. However, recent studies have used multigene DNA analysis employing three gene regions (ITS, ACT, *tef1*) and morphological studies to establish the identity and clarify of the taxonomic status of *Cladosporium* (Bensch et al. 2010, Sandoval-Denis et al. 2016). The *Cladosporium cladosporioides* complex includes a large group of species characterized by unbranched or branched, almost cylindrical conidiophores, bearing ovoid to ellipsoidal intercalary and terminal conidia, smooth or rarely showing fine ornamentation (Bensch et al. 2012, Sandoval-Denis et al. 2016). In this study, we identify *Cladosporium anthropophilum* and *C. subuliforme* as endophytes for the first time from healthy leaves of *Magnolia candoll*i in Yunnan Province, China.

Index Fungorum number: IF815334; Facesoffungi number: FoF06275

morph: Hyphomycetous sporulated on PDA. *Conidiophores* 20–60 × 2–4 μm (x = 30 × 3 μm), hyaline, erect, cylindrical, septate with a thickened wall. *Conidiogenous cells* inconspicuous. *Conidia* 4–10 × 3–5 μm (x = 7 × 4 μm), hyaline, ellipsoidal to subcylindrical, solitary, asceptate, smooth and tapering towards both ends of conidia.

Culture characteristics – Colonies on PDA reaching 30 mm diameter after 7 days at 25°C, colonies circular, margin wavy, fluffy appearance in the middle and on pine needles, colony from above: greyish white and; reverse: dark grey centre and grey margin.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, MT11, living culture, KUMCC 17–0217.


Notes – A new fungal isolate was identified as *Cladosporium anthropophilum* that clustered with the ex-type strain of *C. anthropophilum* (CBS 140685) and sister to *C. anthropophilum* (UTHSC DI 13–226) in the combined ITS and tef1 phylogenetic analysis with 100% ML, 100% MP and 1.00 BPPY support (Fig. 1). The type of *C. anthropophilum* was isolated from human bronchoalveolar lavage fluid in the USA and has also been recorded from an animal abscess, human cerebrospinal fluid, human foot skin, a human hand and human pleural fluid in the USA (Sandoval-Denis et al. 2016). To our knowledge, this is the first report of *C. anthropophilum* occurring on fresh leaves of *Magnolia candolli* (Sandoval-Denis et al. 2016). *Cladosporium anthropophilum* was reported on seed coat of *Pinus armandii* as saprobe or weak pathogen in Yunnan Province, China (Tibpromma et al. 2019). Therefore, we report *C. anthropophilum* as the first record from *Magnolia candolli* in Yunnan Province, China.

*Cladosporium subuliforme* Bensch, Crous & U. Braun, Studies in Mycology 67: 77 (2010)  Fig. 3

Index Fungorum number: IF517090; Facesoffungi number: FoF09439.

See description in Bensch et al. (2010)

Culture characteristics – Colonies on PDA reaching 28 mm diameter after 5 days at 25°C, colonies circular, margin wavy, flat, velvety appearance with greyish aerial mycelia in the middle, colony from above: greyish brown and; reverse: dark brown centre and brown margin.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, MT10 (MFLU 190216) = KUMCC 17–0216.


Notes – According to the combined ITS and tef1 phylogeny, this new isolate (MT10) clustered with *Cladosporium subuliforme* (CBS 126500) with 64% ML, 72% MP and 0.85 BPPY support (Fig. 1). We were unable to observe asexual or sexual morph characters in the culture. *Cladosporium subuliforme* was previously recorded from *Chamaedorea metallica* (Arecalesaceae) in Chiang Mai, Thailand (Bensch et al. 2010). In addition, *C. subuliforme* was recorded as a pathogen causing yellow leafspots on *Capsicum annuum* (Solanaceae) in Cuba (Ramos et al. 2016) and leafspots on *Carya illinoinensis* (Juglandaceae) (Walker et al. 2018). *Cladosporium subuliforme* was also recorded from indoor air in China (Bensch et al. 2018). Here we record the endophytic lifestyle of *C. subuliforme* from leaves of *Magnolia candolli* Yunnan Province, China.

**Subclass Pleosporomycetidae** C.L. Schoch et al.

**Pleosporales** Lutt. ex M.E. Barr

**Corynesporascaceae** Sivan.

*Corynesporascaceae* was introduced by Sivanesan (1996) with *Corynesporasca* as type genus and *C. caryotae* the type species. *Corynesporasca* was considered as the sexual morph of *Corynespora* (Sivanesan 1996). Rossman et al. (2015) recommended using the generic name *Corynespora* over *Corynesporasca* due to the extensive use of this name as plant pathogenic fungi. *Corynespora* was typified by *C. mazei*, a synonym of *C. cassincola* (Rossman et al. 2015). Hongsanan et al. (2020) accepted *Corynesporasca* and *Corynespora* as distinct genera in Corynesporascaceae until molecular data of the type species are available.
Figure 1 – Phylogram generated from maximum likelihood analysis based on combined ITS and tefl sequence data. Related sequences of Cladosporium cladosporioides complex were obtained from Sandoval-Denis et al. (2016). Seventy-three strains are included in the combined gene analyses comprising 2501 characters after alignment (850 characters for ITS and 634 characters for tefl). Cladosporium basiinflatum (CBS 822.64) is used as the outgroup taxon from C. herbarum.
complex. The best RAxML tree with a final likelihood value of -7888.749966 is presented. The matrix had 424 distinct alignment patterns, with 16.96% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.237147$, $C = 0.277351$, $G = 0.257682$, $T = 0.227821$; substitution rates $AC = 1.988191$, $AG = 3.176027$, $AT = 2.194815$, $CG = 0.863163$, $CT = 7.079129$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.633049$. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal to or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold.

**Figure 2** – *Cladosporium anthropophilum* (KUMCC 17–0217, new host record). a Colonies on PDA. b Mycelia masses on pine needle. c Mycelia masses. d Conidiophores and developing conidia. e Secondary ramoconidia and conidia. f, g Conidia. Scale bars: c, d = 20 μm, e–g = 5 μm.

**Figure 3** – *Cladosporium subuliforme* (MFLUCC 19–0051, new geographical and host record). a Mycelia on PDA. b, c Mycelia masses with hyphal coils. Scale bars: b = 50 μm, c = 30 μm.

*Corynespora* Güßow

Species of *Corynespora* are pathogens occurring on a broad range of substrates and causing diseases on leaves to twigs of forest plants in tropics and subtropics regions (Kumar & Singh 2016, Crous et al. 2018). *Corynespora* includes approximately 200 species names (Rossman et al. 2015, Voglmayr & Jaklitsch 2017). There are 208 epithets listed in Index Fungorum (2021). Some species of *Corynespora* are reported as endophytes and saprobes (Pujade-Renaud et al. 2019). Pathogenic *C. cassiicola* is the causal agent of the economically important *Corynespora* leaf fall.
Corynespora cassiicola has been recorded as not only as a pathogen but also as an endophyte in asymptomatic rubber leaves collected in Brazil (Pujade-Renaud et al. 2019). In this study, we report the endophytic lifestyle of C. cassiicola from Magnolia candollii in Yunnan Province, China.

Corynespora cassiicola (Berk. & M.A. Curtis) C.T. Wei, Mycological Papers 34: 5 (1950)  
Index Fungorum number: IF296024; Facesoffungi number: FoF06664  
Endophytic in fresh leaves of Magnolia candollii. Colonies on PDA white aerial mycelia and brown mycelia on pine needles. Mycelia superficial and immersed composed of septate, branched, 2–4 μm wide, hyaline and brown, with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: Hyphomycetous sporulated on PDA. Conidiophores 22–50 × 5–6 μm (x = 35 × 5.4 μm), hyaline, simple, erect or slightly curved. Conidiogenous cells not observed. Conidia 18–30 × 5–8 μm (x = 26 × 6 μm), hyaline to light brown, oblong, sometimes tapering towards the base of the conidia, solitary or in false chains with 0–2 septa.

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, feathery appearance with white aerial mycelia and brown mycelia on pine needles, colony from above: white and brown on pine needles and; reverse: grey centre and dark grey margin.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candollii (Magnoliaceae), 26 April 2017, N. I. de Silva, MT21 (MFLU 19–0520, dried culture); living culture, MFLUCC 19–0054 = KUMCC 17–0224.


Notes – A new isolate (MFLUCC 19–0054) was recovered from healthy leaves of Magnolia candollii in Yunnan, China. The isolate has a close phylogenetic affinity to Corynespora cassiicola (CBS 161.60) in combined ITS, LSU, SSU and tef1 sequence data analyses (Fig. 4). Corynespora cassiicola causes corynespora leaf fall in rubber plantations in Asia and Africa and responsible for yield loss in rubber (Pujade-Renaud et al. 2019). Corynespora cassiicola has been recorded from many plant species viz. Acanthus ilicifolius, Bambusa sp., Calotropis gigantean, Cassia sp. and Eucalyptus sp. (Farr & Rossman 2021). Previously, C. cassiicola was recorded from Magnolia garrettii as endophytes and saprobes in Thailand (Promputtha et al. 2010). In this study, we provide the first endophytic association of this species with M. candollii as a new host record.

Didymellaceae Gruyter et al.

De Gruyter et al. (2009) established the family with the type Didymella Sacc (emend. Q. Chen & L. Cai). Members of this family are cosmopolitan and distributed across a wide range of host plant families (such as Caprifoliaceae, Orchidaceae, Poaceae, Tamaricaceae) (Chen et al. 2015, 2017, Hyde et al. 2018, Jayasiri et al. 2019, Raza et al. 2019). Species of Didymellaceae are plant pathogens on a wide range of hosts, mainly causing leaf and stem lesions (Chen et al. 2017), endophytic, saprobic, fungicolous and lichenicolous (Aveskamp et al. 2010), as well as a few human pathogens (de Hoog et al. 2011). The family comprises 35 genera (Hongsanan et al. 2020).

Epicoccum Link

Epicoccum was introduced by Link (1815) and emended by Chen et al. (2015). The genus comprises plant pathogens (Raza et al. 2019), saprobes (Jayasiri et al. 2017) and endophytes (Dzoyem et al. 2017). Epicoccum nigrum is one of the frequently isolated endophytes of sugarcane plants (Favaro et al. 2012). Favaro et al. (2012) reported that Epicoccum nigrum enhance the root system biomass and inhibit the in vitro growth of sugarcane pathogens, Ceratocystis paradoxa, Colletotrichum falcatum, Fusarium verticillioides and Xanthomonas albilineans. These findings suggest that endophytic Epicoccum nigrum could be used as a natural antagonist for plant pathogens in sugarcane. Epicoccum has hyphomycetous and coelomycetous synanamorphs (Jayasiri et al. 2017, Thambugala et al. 2017). The hyphomycetous anamorph is characterized in
having dark sporodochia with branched conidiophores and mono- to polyblastic, colourless conidiogenous cells that produce coloured, sometimes verruculose, dictyoconidia (Seifert et al. 2011). The coelomycetous synanamorph is characterized by the formation of conidia in pycnidial conidiomata (Chen et al. 2015).

Figure 4 – Phylogram generated from maximum likelihood analysis based on combined ITS, LSU, SSU and tef1 sequence data. Related sequences of Corynespora were obtained from Hyde et al. (2020c). Twenty three strains are included in the combined gene analyses comprising 3280 characters after alignment (580 characters for ITS, 850 characters for LSU, 1010 characters for SSU and 840 characters for tef1). Two strains of Cyclothyriella rubronotata (TR and TR9) are used as outgroup taxa. The best RAxML tree with a final likelihood value of -10047.444970 is presented. The matrix had 591 distinct alignment patterns, with 27.28% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.241833, C = 0.255051, G = 0.272435, T = 0.230681; substitution rates AC = 1.847808, AG = 2.414015, AT = 1.809939, CG = 1.229651, CT = 8.652446, GT = 1.000000; gamma distribution shape parameter α = 0.731416. Bootstrap values
for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequence is indicated in red. Type and ex-type strains are in bold.

Figure 5 – Corynespora cassiicola (MT21/MFLUCC 19–0054, new host record). a, b Brown mycelial masses on a pine needle. c Brown and hyaline mycelia. d, e Conidiophores bearing conidia. f Conidial chain. g, h Conidia. Scale bars: c = 50 μm, d–h = 10 μm.

Epicoccum endophyticum N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.

Index Fungorum number: IF556133; Facesoffungi number: FoF09491

Etymology – The epithet “endophyticum” refers to the endophytic lifestyle of this fungus.

Holotype – MFLU 20–0584


Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, velvety appearance with cream and white mycelia, colony from above: white and light brown; reverse: brown.
Figure 6 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, tub2, and RPB2 sequence data. Related sequences of Epicoccum were obtained from Raza et al. (2019). Forty-five strains are included in the combined gene analyses comprising 3050 characters after alignment (1300 characters for LSU, 500 characters for ITS, 350 characters for tub2 and 900 characters for RPB2). Neoccucurbitaria aquatica (CBS 297.74) is used as outgroup taxon. The best RAxML tree with a final likelihood value of -12694.220358 is presented. The matrix had 741 distinct alignment patterns, with 25.52% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.241547, C = 0.244332, G = 0.273655, T = 0.240465; substitution rates AC = 1.588918, AG = 5.144466, AT = 1.850477, CG = 1.096518, CT = 12.554057, GT = 1.000000; gamma distribution shape parameter α = 0.492764. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian...
Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, S3 (MFLU 20–0584, holotype, dried culture); ex–type living culture, MFLUCC 19–0097 = KUMCC 17–0229.

Additional materials – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva MT5 (MFLU 19–0510, dried culture); living culture, MFLUCC 19–0047 = KUMCC 17–0211.


Notes – According to the phylogeny, *Epicoccum endophyticum* is closely related to the ex-neotype of *E. oryzae* (CBS 173.34) with 56% ML and 0.80 BYPP support (Fig. 6). A comparison of the RPB2 gene region of *E. endophyticum* and ex-neotype of *E. oryzae* (CBS 173.34) reveals 15 base pair differences (2.5%) across 600 nucleotides. Ito & Iwadare (1934) introduced *E. oryzae* from rice grains causing “red blotch” disease in Hokkaido, Japan. Hou et al. (2020) proposed the ex-neotype (CBS 173.34) of *E. oryzae* that were deposited in CBS collection by the original author of the species. *Epicoccum oryzae* produced sporodochia consist of globose, subglobose, or pyriform, granular, 9.9–23.1 × 6.6–16.5 μm conidia. *Epicoccum endophyticum* differs from *E. oryzae* in having ellipsoidal, smaller conidia (10–15 × 8–10 μm), whereas *E. oryzae* has subglobose-pyriform, larger conidia (9.9–23.1 × 6.6–16.5 μm) (Ito & Iwadare 1934). In addition, *Epicoccum oryzae* has been recorded as a pathogen on rice grains in Japan, while *E. endophyticum* was recorded as an endophyte from healthy leaves of *Magnolia candolli* in China. Therefore, we introduce *Epicoccum endophyticum* as a new species from China based on both morphology and phylogeny.

Figure 7 – *Epicoccum endophyticum* (MFLU 20–0584, holotype). a, b Sporodochia on PDA. c Section of sporodochium. d Conidia with mycelia. e Conidiogenous cells with attached conidium. f, g Conidia. Scale bars: c = 50 μm, d–h = 10 μm.

*Leptosphaerulina* McAlpine

*Leptosphaerulina* was introduced by McAlpine (1902) to accommodate *L. australis* as the type. *Leptosphaerulina* species have a cosmopolitan distribution and they have been recorded from various plant families of monocotyledons and dicotyledons in temperate and tropical countries.
(Phookamsak et al. 2013, Chen et al. 2017, Tennakoon et al. 2019). The genus is characterized by small, immersed ascomata, obpyriform asci with a large ocular chamber and an apical ring, muriform, ascospores which may be hyaline or pigmented (Zhang et al. 2012, Hyde et al. 2013, Phookamsak et al. 2013, Tennakoon et al. 2019). There are 64 Leptosphaerulina epithets in Index Fungorum (2021), but few species have molecular data. Species of Leptosphaerulina are reported as saprobic or parasitic on leaves or stems of various plants, including important crops (Phookamsak et al. 2013, Tennakoon et al. 2019). This is the first endophytic record of Leptosphaerulina from fresh leaves of Magnolia candolli in China.

**Figure 8** – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU, ITS, tefl and RPB2 sequence data. Related sequences of Leptosphaerulina were obtained from Tennakoon et al. (2019). Fifteen strains are included in the combined gene analyses comprising 3825 characters after alignment (875 characters for LSU, 1010 characters for SSU, 490 characters for ITS, 865 characters for tefl and 595 characters for RPB2). Two strains of Nothophoma quercina (CBS 633.92 and MFLUCC 16–1392) are used as outgroup taxa. The best RAxML tree with a final likelihood value of -6985.333624 is presented. The matrix had 209 distinct alignment patterns, with 30.39% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.245841, C = 0.236297, G = 0.270335, T = 0.247527; substitution rates AC = 1.811712, AG = 5.570845, AT = 1.370567, CG = 0.734941, CT = 19.045859, GT = 1.000000; gamma distribution shape parameter α = 1.250398. Bootstrap values
for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequence is indicated in red. Type and ex-type strains are in bold.

Index Fungorum number: IF556240, Facesoffungi number: FoF05820  
See description in Tennakoon et al. (2019)  
Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, velvety appearance with white aerial mycelia and black dots like small structures, colony from above: white and brown; reverse: dark brown. Not sporulated on PDA even after two months.  
Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, MT7 (MFLU 19–0512, dried culture); living culture, MFLUCC 19–0049 = KUMCC 17–0213.  
Notes – In this study, we obtained an isolate of *Leptosphaerulina longiflori* from fresh leaves of *Magnolia candolli* that has a close phylogenetic affinity to the ex-type strain of *L. longiflori* (MFLUCC 19–0148) in the present multi-gene phylogenetic analyses (Fig. 8). The sexual morph of the type *L. longiflori* was isolated from dead leaves of *Lilium longiflorum* in Taiwan region (Tennakoon et al. 2019). By considering the molecular data and the host recorded, we consider our collection as a new host record of endophytic *L. longiflori* from *Magnolia candolli*.

**Figure 9** – *Leptosphaerulina longiflori* (MFLUCC 19–0049, new host record). a Mycelia on PDA. b, c Mycelia masses and chlamydospores. Scale bars: b, c = 10 μm.

**Didymosphaeriaceae** Munk  
Munk (1953) introduced Didymosphaeriaceae and typified by *Didymosphaeria*. The family contains many saprobes, while other taxa are endophytes or pathogens in terrestrial and aquatic environments (Barr 2001, Zhang et al. 2012, Ariyawansa et al. 2014, Wanasinghe et al. 2016). Many studies have been conducted on the family and Ariyawansa et al. (2014) synonymized Montagnulaceae under Didymosphaeriaceae based on the priority of the oldest name. The family comprises 32 genera (Hongsan et al. 2020).

**Letendraea** Sacc.  
Saccardo (1880) introduced *Letendraea* with the type of *L. eurotioides*. Species of *Letendraea* are saprobic in terrestrial habitats, pathogenic associated with leaf spot disease in *Cordyline* sp. (Ariyawansa et al. 2014) and also recorded from marine environments (Huang et al. 2019). There are 20 epithets recorded in Index Fungorum (2021) of which four have molecular data.
Figure 10 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU, ITS and tef1 sequence data. Related sequences of *Letendrea* were obtained from Ariyawansa et al. (2014), Wanasinghe et al. (2016). Fifteen strains are included in the combined gene analyses comprising 3135 characters after alignment (850 characters for LSU, 975 characters for SSU, 550 characters for ITS and 760 characters for *tef1*). *Spegazzinia tessartha* (SH 287) and *S. radermacherae* (C264) are used as outgroup taxa. The best RAxML tree with a final likelihood value of -7478.513639 is presented. The matrix had 406 distinct alignment patterns, with 34.13% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.242352, C = 0.240745, G = 0.275677, T = 0.241227; substitution rates AC = 1.299979, AG = 1.543467, AT = 1.104724, CG = 0.913835, CT = 5.527514, GT = 1.000000; gamma distribution shape parameter α = 0.641405. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold.
Letendraea helminthicola (Berk. & Broome) Weese, Transactions of the British Mycological Society 21(3–4): 277 (1938) [1937]

Index Fungorum number: IF252540; Facesoffungi number: FoF09440

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA flat to slightly raised, light brown with velvety appearance. Mycelia superficial and immersed composed of septate, branched, 1–2 μm wide, hyaline, with smooth and thick-walled hyphae. Chlamydospores hyaline, subglobose or irregular shape.

Culture characteristics – Colonies on PDA reaching 23 mm diameter after 7 days at 25°C, colonies circular, margin entire to slightly undulate, flat to slightly raised, dense, velvety appearance, light brown mycelia and colony from above: cream and light brown; reverse: brown.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, MT22, (MFLU 19–0521, dried culture); living culture, MFLUCC 19–0055 = KUMCC 17–0225.


Notes – Sexual morph of Letendraea helminthicola was introduced as hyper-pathogenic fungi on Helminthosporium appendiculatum by Petch (1938) from the UK. Our new isolate (MFLU 19–0521) is phylogenetically related to L. helminthicola (CBS 884.85) with 81% ML, 50% MP and 0.95 BYPP support (Fig. 10). Therefore, based on phylogenetic affinity, we identify our strain as L. helminthicola and it is reported here as a new host and geographical record.

Figure 11 – Letendraea helminthicola (MFLUCC 19–0055, new host and geographical record).

a Colonies on PDA. b, c Mycelia masses and chlamydospores. Scale bars: b = 10 μm, c = 5 μm.

Letendraea magnoliae N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.

Index Fungorum number: IF556134; Facesoffungi number: FoF09492

Etymology – Name reflects the host genus Magnolia, from which the new species was isolated.

Holotype – MFLU 19–0517

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA flat, light brown with velvety appearance. Mycelia superficial and immersed composed of septate, branched, 1–2 μm wide, hyaline, with smooth and thick-walled hyphae. Sexual morph: sporulated on PDA. Ascomata 150–250 μm diam., brown, semi-immersed to superficial covered with mycelia, globose to subglobose, solitary or aggregated. Hamathecium not observed. Peridium 10–15 μm wide, composed of 7–8 layers, thin-walled with equal thickness, small, flattened pseudoparenchymatous, brown cells of textura angularis. Asci (immature asci) 40–50 × 4–7 μm (x = 46 × 6 μm), 8–spored, bitunicate, cylindrical-clavate, slightly curved, sessile. Ascospores 14–16 × 2–4 μm (x = 15 × 3 μm), olivaceous brown, ellipsoidal to fusiform, straight or slightly curved, one seporate, acute at both ends, thick-walled with guttules. Asexual morph: Undetermined.

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat to slightly raised, dense, velvety appearance, orange mycelia and colony from above: orange; reverse: orangish brown.
Figure 12 – Letendraea magnoliae (MFLU 19–0517, holotype). a, b Colonies on PDA (upper view and lower view). c Ascomata on a pine needle. d, e Sections through ascoma. f Peridium. g, h Immature asci stained in congo red. i–l Ascospores. Scale bars: d, e = 50 μm, f–h = 10 μm, i–l = 5 μm.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, MT16 (MFLU 19–0517, holotype, dried culture); ex-type living culture, MFLUCC 19–0052 = KUMCC 17–0221.


Notes – Letendraea magnoliae obtained from healthy leaves, clustered distinctly and basal to other Letendraea species in combined LSU, SSU, ITS and tef1 phylogenetic analysis with 100%
ML, 100% MP and 1.00 BYPP support (Fig. 10). Ascospores of *L. brasiliensis* and *L. cordylinicola* differ from *L. magnoliae* in having mucilaginous sheath while *L. magnoliae* lacks mucilaginous sheath. *Letendraea cordylinicola* has distinct appendage at both ends whereas *L. magnoliae* does not have appendages. *Letendraea magnoliae* differs from *L. eurotioides* and *L. helminthicola* in having ascospores with acute ends. *Letendraea eurotioides* and *L. helminthicola* have ascospores with rounded ends.

*Letendraea eurotioides* is saprobic fungus introduced from the USA on dead branches of *Rubi fruticosi*, *L. cordylinicola* is a saprobic or parasitic fungus from Thailand on leaves of *Cordyline* sp. (Ariyawansa et al. 2014) and *L. helminthicola* is pathogenic fungus on *Helminthosporium appendiculatum* (Petch 1938).

In the current study, we introduce *L. magnoliae* as a new species from China associated with healthy leaves of *Magnolia candolli* based on morphology and phylogeny.

Class **Sordariomycetes** O.E. Erikss. & Winka  
Subclass **Diaporthomycetidae** Senan. et al.  
**Diaporthales** Nannf.  
**Diaporthaceae** Höhn. ex Wehm.

Diaporthaceae was introduced by von Höhnel (1917) and verified by Wehmeyer (1975) to accommodate *Diaporthe* and *Mazzantia*. Castlebury et al. (2002) confirmed Diaporthaceae comprised only *Diaporthe* and *Mazzantia* based on phylogenetic analyses of LSU rDNA sequence data. Lamprecht et al. (2011), Voglmayr & Jaklitsch (2014) subsequently studied Diaporthaceae by adding new genera and Maharachchikumbura et al. (2015) revised the family placement in Diaporthales based on morphology and phylogeny. Wijayawardene et al. (2020) accepted following 15 genera: Apioporthella, Apiosphaeria, Chaetoconis, Chiangraiomyces, Diaporthe, Hyaliappendispora, Leucodiaporthe, Massariothea, Mazzantia, Ophiodiaporthe, Paradiaporthe, Phaeocytostroma, Phaeodiaporthe, Pustulomyces and Stenocarpella.

**Diaporthe** Nitschke

*Diaporthe* species are important plant pathogens, endophytes or saprobes and is typified by *D. eres* Nitschke (Udayanga et al. 2011, 2014a, Gomes et al. 2013, Senanayake et al. 2017, 2018). Pathogenic species are responsible for severe diebacks, cankers, leaf-spots, blights, decay or wilts on a broad range of economically important agricultural crops (Udayanga et al. 2011, Gomes et al. 2013, Gao et al. 2017). Previously, *Diaporthe* species were largely identified based on morphological characters and host association (Dissanayake et al. 2017a, b, Gao et al. 2017). Recently, multi-locus DNA data together with morphology and ecology have been used for *Diaporthe* species delimitation (Dissanayake et al. 2017a, b, Gao et al. 2017, Marin-Felix et al. 2019). Few species complexes have been identified in *Diaporthe*, such as *D. arecae*, *D. eres* and *D. sojae* based on phylogenetic analyses (Udayanga et al. 2014b, 2015, Marin-Felix et al. 2019).

Index Fungorum number: IF811217; Facesoffungi number: FoF08403  
See description in Gao et al. (2016)  
Culture characteristics – Colonies on PDA reaching 23 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, velvety appearance with a thick texture, dark brown and yellowish mycelia on pine needles, colony from above: initially white, became brown and yellowish; reverse: dark brown. The culture did not produce any spores even on prolonged incubation.  
Figure 13 – Phylogram generated from maximum likelihood analysis based on combined ITS, \(tub2\), \(tef1\) and CAL sequence data (with additional strains that closely related to newly generated sequences and removed some distantly related sequences). Related sequences of \textit{Diaporthe} were obtained from Manawasinghe et al. (2019). One hundred and seven strains are included in the combined gene analyses comprising 1700 characters after alignment (530 characters for ITS, 410...
characters for \( \text{tub2} \), 320 characters for \( \text{tef1} \) and 440 characters for \( \text{CAL} \). \textit{Diaporthella corylina} (CBS 121124) is used as outgroup taxon. The best RAxML tree with a final likelihood value of -27814.096708 is presented. The matrix had 1264 distinct alignment patterns, with 31.70% undetermined characters or gaps. Estimated base frequencies were as follows: \( A = 0.217400, C = 0.315962, G = 0.240127, T = 0.226511 \); substitution rates \( AC = 1.078130, AG = 3.025956, AT = 1.253018, CG = 0.883162, CT = 4.360330, GT = 1.000000 \); gamma distribution shape parameter \( \alpha = 0.970320 \). Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% are placed above the branches. The newly generated sequences are indicated in \textcolor{red}{red}. Type and ex-type strains are in \textbf{bold}.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure13}
\caption{Continued.}
\end{figure}
Notes – *Diaporthe apiculata* was introduced by Gao et al. (2016) from healthy leaves of *Camellia sinensis* in Jiangxi Province, China and was also recorded as a pathogen on diseased leaves of *C. sinensis*. Two new isolates of GMT16 and GMT23 share a close phylogenetic affinity to ex-type of *D. apiculata* (LC 3418) in the present combined ITS, *tub2*, *tef1* and CAL gene data analyses. This species has not been reported from the plant family Magnoliaceae (Farr & Rossman 2021) and here we provide the first association of this species with plant host *Magnolia*.


Index Fungorum number: IF810578; Facesoffungi number: FoF08406

See description in Huang et al. (2015)

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, velvety appearance, white and light brown mycelia, elevated mycelia masses (pycnidia like structures), colony from above: white and light brown; reverse: dark brown. The culture did not produce any spores even on prolonged incubation.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, S13 (MFLU 20–0590, dried culture); living culture, MFLUCC 19–0104 = KUMCC 17–0238.


Notes – A new isolate (S13) clustered with *Diaporthe biconispora* (ex-type; ZJUD62 and two other strains; ZJUD60 and ZJUD61) with 100% ML and 100% MP support (Fig. 13). Huang et al. (2015) introduced endophytic *D. biconispora* from the symptomless branch of *Citrus grandis* in Fujian Province, China. And also, *D. biconispora* recorded from symptomless branch of *Fortunella margarita* in Guangxi Province, China (Huang et al. 2015). The current collection of endophytes revealed *D. biconispora* associated with healthy leaves of *Magnolia candolli* in Yunnan Province, China and on *M. garrettii* Chiang Mai Province, Thailand.

Figure 14 – *Diaporthe apiculata* (MFLUCC 18–0938, new host record). a Upper view of culture on PDA. b Mycelia on PDA. c Mycelia masses with chlamydospores. Scale bars: c = 20 μm.

Figure 15 – *Diaporthe biconispora* (MFLUCC 19–0104, new host record). a Upper view of culture on PDA. b Mycelia on PDA. c Mycelia masses with chlamydospores. Scale bars: c = 10 μm.
Diaporthe chinensis N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.

Index Fungorum number: IF556135; Facesoffungi number: FoF09493

Etymology – Referring to the country where the specimen was collected, China.

Holotype – MFLU 20–0587

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA flat, white, felted appearance. Mycelia superficial and immersed composed of septate, branched, 2–3 μm wide, hyaline, with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: conidial masses with mycelia in culture on PDA. Conidiophores 12–15× 2–3 μm (x = 13 × 2.6 μm), hyaline, cylindrical, straight or slightly curved. Conidiogenous cells not observed. Alpha conidia 10–14 × 3–6 μm (x = 12 × 5 μm), hyaline, fusiform, both ends obtuse with single guttulate. Beta conidia not observed.

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, felted appearance, white mycelia, colony from above: white; reverse: cream.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva S10 (MFLU 20–0587, holotype, dried culture); living culture, MFLUCC 19–0101 = KUMCC 17–0235.

Additional materials – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, S16, living culture, MFLUCC 19–0106 = KUMCC 17–0241.

GenBank numbers – (S10); ITS: MW187324, tub2: MW245013, tef1: MW205017, CAL: MW294199, (S16); ITS: MW187325, tub2: MW245014, tef1: MW219603, CAL: MW294200.

Notes – Based on combined multi-gene phylogenetic analysis, Diaporthe chinensis clustered separately, sister to D. yunnanensis with 92% ML and 95% MP support (Fig. 13). Alpha conidia of D. chinensis (10–14 × 3–6 μm) are larger than those of D. yunnanensis (3–6.5 × 1–2.5 μm) (Gao et al. 2017). Diaporthe chinensis can also be distinguished from the ex-type of D. yunnanensis (LC 6168) considering base pair differences of ITS, tub2, tef1 and CAL; 13/490 = 2.65% in ITS, 14/397 = 3.52% in tub2, 13/293 = 4.43% in tef1, 12/410 = 2.92% in CAL. Diaporthe yunnanensis was

Figure 16 – Diaporthe chinensis (MFLU 20–0587). a Upper view of culture on PDA. b, c Mycelia on PDA. d, e Mycelia masses and chlamydospores. f Conidia with mycelium. Scale bars: d–f = 20 μm.
isolated from healthy leaves of Coffea sp. in Xishuangbanna, Yunnan Province, China (Gao et al. 2017). Diaporthe chinensis found to associate with healthy leaves of Magnolia candolli in Xishuangbanna, Yunnan Province, China. Therefore, herein we introduce D. chinensis as a new species from China associated with healthy leaves of M. candolli based on morphology and phylogeny.


- Index Fungorum number: IF820684; Facesoffungi number: FoF09441
- See description in Gao et al. (2017)
- Culture characteristics – Colonies on PDA reaching 22 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, felted appearance, white and yellowish mycelia, colony from above: white and yellowish; reverse: brown. The culture did not produce any spores even on prolonged incubation.
- Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, MT8 (MFLU 19–0513, dried culture); living culture, MFLUCC 19–0050 = KUMCC 17–0214.
- Notes – Pathogenic Diaporthe velutina was introduced from diseased leaves of Neolitsea sp. in Jiangxi Province, China (Gao et al. 2017). Gao et al. (2017) additionally reported pathogenic D. velutina from diseased leaves of Callerya cinerea and Camellia sinensis in China. An endophytic strain also recovered from healthy leaves of C. sinensis in Xishuangbanna, Yunnan Province, China (Gao et al. 2017). A new endophytic strain, MT8 clustered with D. velutina with 100% ML and 100% MP support in the combined ITS, tub2, tef1 and CAL phylogeny (Fig. 13). Therefore, we consider this is the first record of endophytic D. velutina from healthy leaves of Magnolia candolli.

**Diaporthe sp. 1**

*Endophytic* in fresh leaves of Magnolia candolli. Colonies on PDA flat, brown felted appearance. Mycelia superficial and immersed composed of septate, branched, 2–3 µm wide, hyaline and light brown with smooth and thick-walled hyphae. Conidia were not observed. Some thick-walled, light brown, ovoid, 0–2-septate, chlamydospore-like structures, embedded in hyphae were observed.

- Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, velvety appearance, light brown mycelia and white aerial mycelia and colony from above: light brown; reverse: dark brown.
- Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, S11 (MFLU 20–0588, dried culture); living culture, MFLUCC 19–0102 = KUMCC 17–0236.

**Figure 17** – Diaporthe velutina (MFLUCC 19–0050, new host record). a Upper view of culture on PDA. b Mycelia on PDA. c Mycelia masses. Scale bars: c = 20 µm.

**Figure 18**

Diaporthe sp. 1
Notes – Based on combined multi-gene phylogenetic analysis, the new isolate (S11) clustered separately, sister to *Diaporthe hongkongensis* and *D. lithocarpus* with 96% ML and 100% MP support (Fig. 13). This new isolate did not sporulate in culture. Therefore it was not possible to observe conidial characters. The ex-type of *D. hongkongensis* (CBS 115448) was isolated on fruits of *Dichroa febrifuga* in Hong Kong (Gomes et al. 2013). The ex-type of *D. lithocarpus* (CGMCC 3.15175) was isolated from leaf spots of *Lithocarpus glabra* in Zhejiang Province, China (Gao et al. 2014). The new isolate (S11) found to associate with healthy leaves of *Magnolia candolli* in Xishuangbanna, Yunnan Province, China. The new isolate (S11) might be a distinct taxon based on our phylogenetic analyses and host association. However, we do not propose a new species of *Diaporthe* here, as we were unable to observe complete morphology of the fungus.

Figure 18 – *Diaporthe* sp. 1 (MFLU 20–0588). a Upper view of culture on PDA. b, c Mycelium on PDA. d–f Mycelia masses and chlamydospore-like structures. Scale bars: d = 20 μm, e, f = 10 μm.

**Subclass Hypocreomycetidae** O.E. Erikss. & Winka  
**Glomerellales** Chade. ex Re´blova´ et al.  
**Glomerellaceae** Locq.

*Glomerellaceae* (Glomerellales, Sordariomycetes), consists of the sole member of *Colletotrichum* (Jayawardena et al. 2016, Hyde et al. 2020b). Species of this genus are important pathogens; some are endophytes as well as saprobes (Hyde et al. 2014, Jayawardena et al. 2016).

**Colletotrichum** Corda  
*Colletotrichum* is one of the major economically important plant pathogenic genera with worldwide distribution and studies revealed that these pathogens may exhibit as saprophytic or endophytic lifestyles in nature (Damm et al. 2012, Gautam 2014, Jayawardena et al. 2016). Pathogenic species infect plants using different methods for invading host tissue, ranging from intracellular hemibiotrophic to subcuticular intramural necrotrophy (Gautam 2014). Index Fungorum (2021) lists 928 species epithets for *Colletotrichum*.

Multigene phylogenetic analyses detected new species and several species complexes in this genus (Cannon et al. 2012, Jayawardena et al. 2016, 2020). Recent studies used phylogenetic analysis based on concatenated genes for species delimitation within the genus (Jayawardena et al. 2016). Jayawardena et al. (2020) identified 14 species complexes in *Colletotrichum* and 20 singleton species. The gloeosporioides species complex mainly consists of plant pathogens (Weir et

Figure 19 – Phylogram generated from maximum parsimony analysis based on combined ITS, GAPDH, CHS, ACT and tub2 sequence data. Related sequences of Colletotrichum (C. gloeosporioides complex) were obtained from Jayawardena et al. (2020). Seventy eight strains are included in the combined gene analyses comprising 1760 characters after alignment (500
Characters for ITS, 250 characters for GAPDH, 280 characters for CHS, 250 characters for ACT and 480 characters for tub2. Colletotrichum boninense (CBS 123755) and C. brasiliense (CBS 128501) are used as outgroup taxa. The best RAxML tree with a final likelihood value of -13102.99262 is presented. The matrix had 1042 distinct alignment patterns, with 20.67% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.229409, C = 0.295382, G = 0.243801, T = 0.231407; substitution rates AC = 1.015583, AG = 2.923416, AT = 1.312931, CG = 0.864260, CT = 4.263907, GT = 1.000000; gamma distribution shape parameter α = 1.566948. Bootstrap values for maximum parsimony, maximum likelihood equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold.

**Colletotrichum chiangmaiense** N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.  
Fig. 20

Index Fungorum number: IF556379; Facesoffungi number: FoF09494

Etymology – Referring to the location where the fungus was discovered, Chiang Mai Province, Thailand.

Holotype – MFLU 20–0604

Endophytic in fresh leaves of Magnolia garrettii. Colonies on water agar (WA), flat, white, velvety appearance. Mycelia superficial and immersed composed of septate, branched, 2–3 µm wide, hyaline and light brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: acervulus with conidial masses and mycelia in culture on WA. Conidiophores not observed. Conidiogenous cells 2–4 × 1–2 µm (μ = 3 × 1.4 µm), hyaline, short, cylindrical or ampulliform, formed directly from vegetative hyphae. Conidia 4–7 × 2–3.5 µm (μ = 5 × 2 µm), hyaline, fusiform, ellipsoidal to cylindrical with one end slightly acute when immature state, both ends obtuse when mature.

Culture characteristics – Colonies on WA reaching 20 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, velvety appearance, white aerial mycelia and colony from above: white; reverse: cream.

Material examined – THAILAND, Chiang Mai Province, healthy leaves of Magnolia garrettii (Magnoliaceae), 14 August 2017, N. I. de Silva, GMT30 (MFLU 20–0604), holotype, dried culture; ex-type living culture, MFLUCC 18–0945.


Notes – Colletotrichum chiangmaiense (GMT30) has a close phylogenetic affinity to C. henanense and C. yulongense and clustered separately to those two sister taxa with 65% MP, 78% ML and 0.79 BYPP support (Fig. 19). Conidia of C. chiangmaiense (4–7 × 2–3.5 µm) is smaller than those of C. henanense (8–17 × 3–5.5 µm) (Liu et al. 2015a) and C. yulongense (11–13 × 4–5 µm) (Wang et al. 2019).

In a BLASTn search of GenBank, the ITS sequence had 100% similarity to C. siamense (bl20), the GAPDH sequence had 98% similarity to C. camelliae (MZ 12), the CHS had 100% similarity to C. camelliae (FWY41) and the ACT sequence had 93% similarity to C. aotearoa (BM2). Comparison of nucleotides differences between C. chiangmaiense and C. henanense reveals 3/480 = 0.62% in ITS, 6/220 = 2.72% in GAPDH, 10/220 = 4.54% in ACT. CHS sequence data is not available for C. henanense. Comparison of nucleotides differences between C. chiangmaiense and C. yulongense reveals 5/480 = 1.04% in ITS, 11/200 = 5.5% in GAPDH, 6/222 = 2.7% in CHS, 11/198 = 5.5% in ACT. A PHI test revealed no significant recombination event among C. chiangmaiense and its closely related taxa (Fig. 21).

Colletotrichum henanense was isolated from Camellia sinensis in China (Liu et al. 2015a) and C. yulongense from healthy leaves of Vaccinium dunalianum var. urophyllum in China (Wang et al. 2019). There is a Colletotrichum species isolated from leaves of Magnolia grandiflora in Portugal named as C. magnoliae. Conidia of C. magnoliae (15–18 × 5–6 µm) (Saccardo 1931) are larger than those of C. chiangmaiense. In this study, C. chiangmaiense is introduced as a novel taxon based on morphology and phylogeny.
Figure 20 – *Colletotrichum chiangmaiense* (MFLU 20–0604, holotype). a, b Mycelia on WA. c Mycelia. d Developing conidia. e–g Conidia. Scale bars: c–g = 5 μm.

**Figure 21** – The splits graph from the pairwise homoplasy index (PHI) test generated from the concatenated gene set of ITS, GAPDH, CHS, ACT and *tub2* sequence data of closely related species of *Colletotrichum* using both LogDet transformation and splits decomposition. PHI test result (Φw) < 0.05 indicates significant recombination within the dataset. The strain determined in this study is in red.


Index Fungorum number: IF515409; Facesoffungi number: FoF06767

Conidiogenous cells 6–8 × 2–3 μm (\(\bar{x} = 7 \times 2.3 \mu m\)), hyaline, cylindrical, formed directly from vegetative hyphae. Conidia 10–13 × 4–6 μm (\(\bar{x} = 12 \times 5 \mu m\)), hyaline, cylindrical, the apex and base rounded, straight, aseptate, smooth–walled. Sexual morph: Ascomata black, semi-immersed to superficial covered with mycelia, globose to subglobose, solitary or aggregated. Asci 60–80 × 6–10 μm (\(\bar{x} = 72 \times 8 \mu m\)), hyaline, 8–spored, unitunicate, cylindrical, cylindrical–clavate, slightly curved, sessile. Ascospores 15–20 × 3–5 μm (\(\bar{x} = 17 \times 4 \mu m\)), hyaline, fusiform, straight or slightly curved, obtuse to somewhat acute or acute both end, smooth–walled.

Culture characteristics – Colonies on PDA reaching 23 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, velvety appearance, white aerial mycelia and colony from above: brown; reverse: cream.


Notes – Phylogenetic analyses of a concatenated ITS, GAPDH, CHS, ACT and tub2 sequence data showed that our strains (S220, S20 and GMT26) group together with C. fructicola (Fig. 19). The new isolate (S220) shares similar morphology with the type of C. fructicola (MFLU 09–0228) in having hyaline, cylindrical conidia with obtuse to slightly rounded ends and hyaline, slightly curved to curved ascospores with obtuse to slightly rounded ends (Prihastuti et al. 2009). Conidia of new isolate (S220) (12–14 × 4–5 μm) and the type C. fructicola (9.7–14 × 3–4.3 μm)
have overlapping size range (Prihastuti et al. 2009). Ascospores of new isolate (S220) (15–18 × 3–5 µm) are slightly larger than the type C. fructicola (9–14 × 3–4 µm) (Prihastuti et al. 2009). The type of C. fructicola was introduced on berries of Coffea arabica in Chiang Mai Province, Thailand (Prihastuti et al. 2009). Colletotrichum fructicola was isolated as endophytes and saprobes from Coffea arabica (Prihastuti et al. 2009). Colletotrichum fructicola has a wide host range and geographical distribution including Arachis sp. (Fabaceae), Citrus bergamia (Rutaceae) in China; Cymbopogon citrates (Poaceae) in Thailand; Dioscorea alata in Nigeria, Ficus sp. (Dioscoreaceae) in Germany, Fragariaxananassa (Rosaceae) in Canada and the USA, Limonium sp. (Plumbaginaceae) in Israel, Malus domestica (Rosaceae) in Australia, Brazil and Uruguay; Mangifera indica (Anacardiaceae) in India and Brazil; Persea americana (Lauraceae) in Australia, Pyrus pyrifolia (Rosaceae) in Japan, Theobroma cacao (Malvaceae) and Tetragastris panamensis (Burseraceae) in Panama, Pennisetum purpureum (Poaceae) in Thailand; and Camellia sinensis (Wang et al. 2016), Rubus glaucus (Rosaceae) and Vitis vinifera (Vitaceae) in China (Jayawardena et al. 2016). In this study, we report an isolate associate with healthy leaves of Magnolia garrettii in Chiang Mai Province, Thailand. Furthermore, we identified two isolates of C. fructicola and reported its occurrence associate with healthy leaves of Magnolia candolli in Yunnan, China, for the first time.


Index Fungorum number: IF158410; Facesoffungi number: FoF09424

See description in Weir et al. (2012)

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, velvety appearance, white aerial mycelia and colony from above: white; reverse: cream. The culture did not produce any spores even on prolonged incubation.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, S14, living culture, MFLUCC 19–0105 = KUMCC 17–0239.


Notes – Multigene phylogenetic analyses show that the new strain S14 is sister to the ex-type of Colletotrichum gloeosporioides (CBS 112999) with 100% MP, 100% ML and 1.00 BYPP support (Fig. 19). Colletotrichum gloeosporioides has reported from different geographical regions and various host plants, i.e. on Citrus sinensis (ex-epitype culture of C. gloeosporioides, CBS 112999) in Italy, on Carya liliifera in Australia, on Citrus sp., on Ficus sp. in New Zealand, on Mangifera indica in South Africa, on Pueraria lobata, on Citrus sp. on Vitis vinifera in the USA (Weir et al. 2012). Colletotrichum gloeosporioides was recorded from Magnolia liliifera in Thailand as endophytic lifestyle (Promputtha et al. 2005). In addition, C. gloeosporioides was recorded on Magnolia denudata in China, M. grandiflora in India, North Carolina, Magnolia officinalis var. biloba in China and Magnolia sp. in Poland (Farr & Rossman 2021). Therefore, we identify S14 as C. gloeosporioides based on phylogeny and our isolate is introduced here as a new endophytic host record from M. candolli collected in China.

**Colletotrichum xishuangbannaens** N.I. de Silva, Lumyong & K.D. Hyde, sp. nov. Fig. 23

Index Fungorum number: IF556380; Facesoffungi number: FoF09495

Etymology – Referring to the location where the fungus was discovered, Xishuangbanna, Yunnan Province, China.

Holotype – MFLU 20–0593

morph: coelomycetous, conidial masses with mycelia in culture on PDA. Conidiophores not observed. Conidiogenous cells 15–18 × 1.5–2 µm (\(\bar{x} = 16 \times 1.7 \mu m\)), hyaline, cylindrical, smooth-walled. Conidia 9–12 × 3–4 µm (\(\bar{x} = 11 \times 3.5 \mu m\)), hyaline, cylindrical with the apex and base rounded. Appressoria not observed.

Culture characteristics – Colonies on PDA reaching 27 mm diameter after 7 days at 25°C, colonies circular, margin undulate, slightly raised, velvety appearance, cream aerial mycelia and colony from above: light brown; reverse: brown.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, S17 (MFLU 20–0593, holotype, dried culture); ex-type living culture, MFLUCC 19–0107 = KUMCC 17–0242.

Additional materials – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva S118, living culture, MFLUCC 19–0116.

GenBank numbers – (S17); ITS: MW346469, GAPDH: MW537586, CHS: MW660832, ACT: MW652294, (S118); ITS: MW346470, GAPDH: MW537587, CHS: MW660833, ACT: MW652295.

Notes – Colletotrichum xishuangbannaense clustered closely related to C. aeschynomenes with 40% MP, 25% ML, 0.83 BYPP support (Fig. 19). Morphologically, C. xishuangbannaense differs from C. aeschynomenes in having small conidia. Colletotrichum aeschynomenes has 14–20 × 4–5 µm conidia (Weir et al. 2012) and C. xishuangbannaense has 9–12 × 3–4 µm conidia. Colletotrichum magnoliae was isolated from leaves of Magnolia grandiflora in Portugal (Saccardo 1931). Conidia of C. magnoliae (15–18 × 5–6 µm) (Saccardo 1931) are larger than C. xishuangbannaense.

In a BLASTn search of GenBank, the ITS sequence had 100% similarity to C. aenigma (GQH124), the GAPDH sequence had 98% similarity to C. fructicola (TS–B173), the CHS had 98% similarity to C. siamense (ccmi3) and the ACT sequence had 99% similarity to C. siamense (LC0537). Comparison of nucleotides differences between C. xishuangbannaense and the ex-type C. aeschynomenes (ICMP 17673) reveals 3/480 = 0.62% in ITS, 5/220 = 2.27% in GAPDH, 5/250

Figure 23 – Colletotrichum xishuangbannaense (MFLU 20–0593, holotype). a Colonies on PDA. b Conidial masses. c Mycelia and conidial masses. d Conidiogenus cells with conidia. e, f Conidia. Scale bars: c = 50 µm, d–f = 10 µm.
= 2% in CHS and 4/230 = 1.73% ACT. A PHI test revealed no significant recombination event among C. xishuangbannaense and its closely related taxa (Fig. 24). Colletotrichum aeschynomenes was isolated as a pathogen of stem lesion from Aeschynomone virginica in the USA (Weir et al. 2012). In this study, we introduce C. xishuangbannaense as a novel taxon based on phylogeny and morphology.

Figure 24 – The splits graph from the pairwise homoplasy index (PHI) test generated from the concatenated gene set of ITS, GAPDH, CHS, ACT and tub2 sequence data of closely related species of Colletotrichum using both LogDet transformation and splits decomposition. PHI test result (Φw) < 0.05 indicates significant recombination within the dataset. The strain determined in this study is in red.


Index Fungorum number: IF581687; Facesoffungi number: FoF09442

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA, slightly raised, cream, light brown mycelia. Mycelia superficial and immersed composed of septate, branched, 2–3 μm wide, hyaline and light brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: dark brown conidial masses on PDA. Conidiophores not observed. Conidiogenous cells inconspicuous. Conidia 20–22 × 5–7 μm (μ = 21 × 6 μm), hyaline, cylindrical, the apex and base obtuse, straight and or slightly curved, aseptate, smooth-walled with small guttules.

Culture characteristics – Coelomycetous with acervuli in culture. Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin entire, slightly raised, cream and light brown mycelia, colony from above: cream, reverse: light brown.

Material examined – THAILAND, Chiang Mai Province, healthy leaves of Magnolia garrettii (Magnoliaceae), 14 August 2017, N. I. de Silva GMT20, living culture, MFLUCC 18–0939, GMT5, living culture, MFLUCC 18–0934, CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, MT9, living culture, KUMCC 17–0215.


Notes – Colletotrichum karsti was introduced by Yang et al. (2011) associate with anthracnose lesions on leaves of Vanda sp. in China. The new collection differs from the holotype (GZAAS 090006), in having larger (20–22 × 5–7 μm) conidia. Colletotrichum karsti has 12.5–19.5 × 6–8.5 μm conidia (Yang et al. 2011). Colletotrichum karsti has a wide host species range from
Orchidaceae as a pathogen causing dark brown to black, ellipsoid lesions on leaves of *Arundina graminifolia*, *Calanthe argenteo-striata*, *Eria Coronaria* and also isolated as an endophyte of roots of *Pleione bulbocodioides* (Yang et al. 2011). The new collection is reported as a new host record of *C. karsti* associated with healthy leaves of *Magnolia candolli* (Magnoliaceae) for the first time.

**Figure 25** – Phylogram generated from maximum parsimony analysis based on combined ITS, GAPDH, CHS, ACT and *tub2* sequence data. Related sequences of *C. boninense* complex were obtained from Jayawardena et al. (2020). Thirty four strains are included in the combined gene analyses comprising 1810 characters after alignment (540 characters for ITS, 250 characters for GAPDH, 270 characters for CHS, 260 characters for ACT and 490 characters for *tub2*). *Colletotrichum gloeosporioides* (CBS 112999) and *C. proteae* (CBS 132882) are used as outgroup taxa. The best RAxML tree with a final likelihood value of -8649.542487 is presented. The matrix had 715 distinct alignment patterns, with 11.03% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.227512, C = 0.297420, G = 0.250297, T = 0.224771; substitution rates AC = 1.332056, AG = 3.193250, AT = 1.057960, CG = 0.931964, CT = 4.681567, GT = 1.000000; gamma distribution shape parameter α = 1.481581. Bootstrap values for maximum parsimony and maximum likelihood equal to or greater than 75% and Bayesian posterior probabilities ≥ 0.95 (PP) are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold. 
Figure 26 – Colletotrichum karstii (MFLUCC 18–0939, new host record). a Conidial masses on PDA. b Mycelia and conidia. c–e Conidia. Scale bars: b = 10 μm, c–e = 5 μm.

Subclass Xylariomycetidae O.E. Erikss. & Winka
Amphisphaeriales D. Hawksw. & O.E. Erikss.

Sporocadaceae Corda

Sporocadaceae was introduced by Corda (1842) with the type genus Sporocadus. Members of Sporocadaceae are appendaged coelomycetes and they exhibit as endophytes, plant pathogens or saprobes in a wide range of host plants (Liu et al. 2019). Sporocadaceae mostly consists of asexual morph genera with acervular coelomycetes with appendages (Jaklitsch et al. 2016, Liu et al. 2019). Liu et al. (2019) showed that Sporocadaceae as a well-defined family in the Xylariales and identified 30 genera based on phylogenetic analyses and morphological comparison. Hyde et al. (2020b) accepted 32 genera in the family.

Neopestalotiopsis Maharachch., K.D. Hyde & Crous

Neopestalotiopsis was introduced by Maharachchikumbura et al. (2014b). They are plant pathogenic, saprobic and endophytic species commonly present in tropical and subtropical ecosystems (Maharachchikumbura et al. 2014b, Jayawardena et al. 2019). Morphologically Neopestalotiopsis species can differ from Pestalotiopsis and Pseudopestalotiopsis in having somewhat versicolorous median cells (Maharachchikumbura et al. 2014b). Pestalotiopsis and Pseudopestalotiopsis species generally possess concolourous median cells (Maharachchikumbura et al. 2014b). Neopestalotiopsis infects many crops, i.e. blueberry, grapes, strawberry, sweet potatoes, causing leaf spots, trunk diseases, post-harvest fruit rots and root rots worldwide (Jayawardena et al. 2019). There are 43 species epithets are recorded in Index Fungorum (2020). Neopestalotiopsis clavispora and N. saprophytica were recorded from Magnolia species (Magnoliaceae) (Maharachchikumbura et al. 2014b, Farr & Rossman 2021). We report three new records in this study from Magnolia species (Magnoliaceae).

Neopestalotiopsis chiangmaiensis Tibpromma & K.D. Hyde, Fungal Diversity: [139] (2018b)

Index Fungorum number: IF554515; Facesoffungi number: FoF04525

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA flat, spreading, with aerial mycelia. Mycelia superficial and immersed composed of septate, branched, 2–3 μm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial in culture on PDA, globose, solitary or aggregated, embedded or semi-immersed, black. Conidia 18–23 × 3–5 μm (x = 20 × 4 μm), ellipsoid, fusoid, straight to slightly curved, 4–septate; basal cell 3–5 μm long, hyaline, obconic, smooth- and thin-walled; three median cells 12–15 × 3–5 μm (x = 13 × 4 μm) brown, doliform, (the second cell from base 4–6 μm long, third cell 4–5 μm long, fourth cell 4–6 μm long); apical cell 3–4 μm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, smooth- and thin-walled, with 2 tubular apical appendages, 12–15 μm long, arising from the apical crest, filiform, basal appendage 3–5 μm long, single, tubular, unbranched, straight and centric.
Figure 27 – Phylogram generated from maximum likelihood analysis based on combined ITS, tub2 and tef1 sequence data. Fifty nine strains are included in the combined gene analyses comprising 1290 characters after alignment (500 characters for ITS, 420 characters for tub2, and 370 characters for tef1). Pestalotiopsis trachicarpicola (IFRDCCC 2440) is used as outgroup taxon. The best RAxML tree with a final likelihood value of -6323.092983 is presented. The matrix had 508
distinct alignment patterns, with 14.46% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.229682, C = 0.268940, G = 0.212769, T = 0.288609; substitution rates AC = 1.171186, AG = 3.097006, AT = 1.486836, CG = 0.944370, CT = 4.032044, GT = 1.000000; gamma distribution shape parameter α = 0.818017. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold.

Culture characteristics – Colonies on PDA reaching 33 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, slightly raised, white mycelia, colony from above: white; reverse: light yellow.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, MT6 (MFLU 19–0511, dried culture), living culture, MFLUCC 19–0048 = KUMCC 17–0212.

GenBank numbers – ITS: MW248391, tef1: MW259070

Notes – Neopestalotiopsis chiangmaiensis was introduced based on a collection on Pandanus sp. in Thailand by Tibpromma et al. (2018b). In this study, our new isolate clustered with the ex-type of N. chiangmaiensis (MFLUCC 18–0113) with 87% ML, 75% MP and 0.99 BYPP support (Fig. 27). ITS sequence data of the ex-type N. chiangmaiensis (MFLUCC 18–0113) and tub2 sequence data of our isolate (MT6) are not available. The tef1 gene region of both the ex-type N. chiangmaiensis (MFLUCC 18–0113) and the new isolate (MT6) are identical. Therefore, we treat our new isolate as N. chiangmaiensis and this is the first record of N. chiangmaiensis associated with Magnolia candolli in China.

Figure 28 – Neopestalotiopsis chiangmaiensis (MFLUCC 19–0048, new host record). a Colony on PDA producing conidia masses. b, c Conidia. Scale bars: b, c = 10 μm.

Neopestalotiopsis egyptiaca A.M. Ismail, G. Perrone & Crous, Persoonia 35: 271 (2015) Fig. 29

Index Fungorum number: IF813837; Facesoffungi number: FoF09443

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA flat, fluffy appearance with white aerial mycelia. Mycelia superficial and immersed composed of septate, branched, 2–4 μm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial in culture on PDA, globose, solitary or aggregated, embedded or semi-immersed, exuding black conidial masses. Conidia 20–23 × 5–7 μm (x = 20 × 6 μm), ellipsoid, fusoid, straight to slightly curved, 4-septate; basal cell 3–4 μm long, hyaline, obconic, smooth- and thin-walled; three median cells 12–14 × 3–5 μm (x = 13 × 4 μm), brown, doliiform, (the second cell from base 4–5 μm long, third cell 4–5 μm long, fourth cell 4–5 μm long); apical cell 4–5 μm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, smooth-
and thin-walled, with 3 tubular apical appendages, 18–20 µm long, arising from the apical crest, filiform, basal appendage 3–4 µm long, single, tubular, unbranched, straight and centric.

Culture characteristics – Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, fluffy appearance with white mycelia, colony from above: white; reverse: cream.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva MT1, living culture, KUMCC 17–0207.


Notes – The phylogenetic analyses of combined ITS, *tub2* and *tef1* sequence data showed that the new isolate (MT1) clustered with *N. egyptiaca* (Fig. 27). *Neopestalotiopsis egyptiaca* was introduced on a collection from diseased leaves of *Magnifera indica* in Egypt (Crous et al. 2015). This species has not been reported so far from Magnoliaceae (Farr & Rossman 2021). Conidia of new isolate have similar morphology with the ex-type of *N. egyptiaca* (CBS 140162) in having fusiform, 4-septate conidia with brown median cells and 2–3 apical appendages (Crous et al. 2015). Conidia of the new isolate (MT1) (20–23 × 5–7 µm) is slightly smaller than the ex-type *N. egyptiaca* (CBS 140162) (22.5–28 × 6–7.5 µm) (Crous et al. 2015). The ITS gene region of both the ex-type *N. egyptiaca* (CBS 140162) and the new isolate (MT1) is similar. We introduce our new isolate, *N. egyptiaca* as a new host record from *Magnolia candolli* and geographical record from China.

**Figure 29** – *Neopestalotiopsis egyptiaca* (KUMCC 17–0207, new host record). a Colony on PDA producing conidia masses. b, c Conidia. Scale bars: b, c = 10 µm.

*Neopestalotiopsis keteleeria* Y. Song, K.D. Hyde & Yong Wang, Chiang Mai Journal of Science 41(4), 888 (2014a)

Index Fungorum number: IF825388; Facesoffungi number: FoF09444

Endophytic in fresh leaves of *Magnolia candolli*. Colonies on PDA, fluffy appearance, white aerial mycelia. *Mycelium* superficial and immersed composed of septate, branched, 2–3 µm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: *Conidiomata* pycnidial on pine needles, globose, solitary or aggregated, embedded or semi-immersed, exuding black conidial masses. *Conidium* 18–23 × 5–7 µm (x = 20 × 6 µm), ellipsoid, fusoid, fusoid to slightly curved, 4-septate; basal cell 3–5 µm long, hyaline, obconic, smooth- and thin-walled; three median cells 12–15 × 5–7 µm (x = 13 × 6 µm), brown, doliiform, (the second cell from base 4–5 µm long, third cell 4–5 µm long, fourth cell 4–5 µm long); apical cell 4–5 µm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, smooth- and thin-walled, with 3 tubular apical appendages, 25–30 µm long, arising from the apical crest, filiform, basal appendage 3–4 µm long, single, tubular, unbranched, straight and centric.
Culture characteristics – Colonies on PDA reaching 38 mm diameter after 7 days at 25°C, colonies circular, margin entire, fluffy appearance, white mycelia, colony from above: white; reverse: cream.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, MT24 (MFLU 19–0522, dried culture); living culture, MFLUCC 19–0056 = KUMCC 17–0226.


Notes – The new strain is phylogenetically closely related to the ex-type of *N. keteleeria* (MFLUCC 13–0915) (Fig. 27). *Neopestalotiopsis keteleeria* was introduced by Song et al. (2014a) on disease leaves of *Keteleeria pubescens* in China. The ITS gene region of both the ex-type *N. keteleeria* (MFLUCC 13–0915) and the new isolate (MT24) is similar. A comparison of *tub2* gene region indicates ten base pair differences between the ex-type *N. keteleeria* (MFLUCC 13–0915) and the new isolate (MT24) across 400 nucleotides. *tef1* sequence data of the ex-type of *N. keteleeria* (MFLUCC 13–0915) are not available. Conidia of *N. keteleeria* (18.5–24 × 7–9.5 μm) (Song et al. 2014a) and the new isolate (MT24) (18–23 × 6–8) share similar conidial dimensions. Therefore, we identified our current taxon as *N. keteleeria*. This is the first report of endophytic lifestyle of *N. keteleeria* associate with healthy leaves of *Magnolia candolli*.

**Figure 30** – *Neopestalotiopsis keteleeria* (MFLUCC 19–0056, new host record). **a** Conidial masses producing on pine needles. **b, c** Conidia. Scale bars: **b, c** = 10 μm.

**Pestalotiopsis Steyaert**

Species of *Pestalotiopsis* are an appendage-bearing conidial anamorphic form (coelomycetes) commonly found in tropical and temperate ecosystems (Maharachchikumbura et al. 2011). The genus was introduced by Steyaert (1949) with the type *P. maculans*. *Pestalotiopsis* species are known to cause diseases of foliage, stems and cause a considerable reduction on commercial production (Maharachchikumbura et al. 2011, 2013, 2014b, Liu et al. 2017). *Pestalotiopsis* are also isolated as endophytes or occur as saprobes (Maharachchikumbura et al. 2011, 2013, 2014b, Liu et al. 2017). They are able to switch life modes as endophytes, pathogens and saprobes (Maharachchikumbura et al. 2012, Jayawardena et al. 2019). Traditionally, *Pestalotiopsis* species were named according to their host associations and the conidial morphologies such as colour intensities of the median conidial cell (Maharachchikumbura et al. 2014a, Liu et al. 2017). Maharachchikumbura et al. (2012) recommended using combined ITS, *tub2* and *tef1* genes to delimitation species boundaries in *Pestalotiopsis* species. There are 374 species epithets are recorded in Index Fungorum (2021). Wijayawardene et al. (2020) accepted 33 species in *Pestalotiopsis*.

**Pestalotiopsis endophytica** N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.

Index Fungorum number: IF556381; Facesoffungi number: FoF09496

Etymology – The epithet “endophytica” refers to the endophytic lifestyle of this fungus.

Holotype – MFLU 20–0607
Figure 31 – Phylogram generated from maximum likelihood analysis based on combined ITS, tub2 and tefl sequence data. Eighty five strains are included in the combined gene analyses comprising 1460 characters after alignment (530 characters for ITS, 430 characters for tub2, 500 characters for tefl).
Neopestalotiopsis saprophytica (G41) is used as outgroup taxon. The best RAxML tree with a final likelihood value of -11991.194599 is presented. The matrix had 739 distinct alignment patterns, with 13.65% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.234706, C = 0.291700, G = 0.212976, T = 0.260619; substitution rates AC = 0.978127, AG = 2.967800, AT = 1.111125, CG = 0.863582, CT = 3.577924, GT = 1.000000; gamma distribution shape parameter α = 0.727318. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold.

Endophytic in fresh leaves of Magnolia candollii. Colonies on PDA flat slightly raised, spreading, with abundant aerial mycelia. Mycelia superficial and immersed composed of septate, branched, 2–4 µm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial in culture on PDA, globose, solitary or aggregated, embedded or semi-immersed, black, exuding globose, brown to black conidiomata masses. Conidiophores reduced to conidiogenous cells. Conidiogenous cells 7–10 × 3–5 µm (\(\bar{X} = 8 \times 4 \mu m\)), discrete, hyaline, cylindrical to subcylindrical and smooth-walled. Conidia 25–30 × 5–7 µm (\(\bar{X} = 28 \times 6 \mu m\)), ellipsoid, fusoid, straight to slightly curved, 4-septate; basal cell 5–7 µm long, hyaline, obconic, smooth- and thin-walled; three median cells 15–18 × 5–7 µm (\(\bar{X} = 16 \times 6 \mu m\)), brown, doliform, (the second cell from base 4–6 µm long, third cell 6–7 µm long, fourth cell 5–7 µm long); apical cell 5–7 µm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, smooth- and thin–walled, with 2–3 tubular apical appendages (mostly 3), 20–23 µm long, arising from the apical crest, filiform, basal appendage 4–6 µm long, single, tubular, unbranched, straight and centric.

Culture characteristics – Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, slightly raised, velvety appearance, white mycelia, colony from above: white; reverse: cream.

Material examined – THAILAND, Chiang Mai Province, healthy leaves of Magnolia garrettii (Magnoliaceae), 14 August 2017, N. I. de Silva, GMT3 (MFLU 20–0607, holotype, dried culture); ex-type living culture, MFLUCC 18–0932.


GenBank numbers – (GMT3); ITS: MW263946, tef1: MW417119, (GMT31); ITS: MW263947, tef1: MW729384, (GMT222); ITS: MW263948, (GMT333); ITS: MW263949.

Notes – Four new isolates clustered together in a distinct clade with 99% ML, 91% MP and 1.00 BYPP support (Fig. 31) and separated from the sister clade comprising P. australis, P. brassicaceae, P. chamaerops, P. hollandica, P. intermedia, P. italicana, P. Jiangxiensis, P. linearis, P. monochaeta, P. scoparia, P. unicolor and P. verruculosa. Pestalotiopsis endophytica differs from P. brassicaceae, P. hollandica, P. italicana and P. verruculosa in having 3 apical appendages while P. brassicaceae, P. hollandica, P. italicana and P. verruculosa have more than 3 apical appendages (Table 2). Pestalotiopsis monochaeta differs from P. endophytica in having single apical appendage (Maharachchikumbura et al. 2014b). Pestalotiopsis endophytica is morphologically similar to P. australis in having an overlapping range of conidial and appendage dimensions (Table 2). However, P. endophytica has slightly smaller (25–30 × 5–7 µm) conidia when compare to P. australis (26–36) × 7–8.5 µm (Maharachchikumbura et al. 2014b). Further, phylogenetic analyses show P. endophytica is distinctly related to P. australis (Fig. 31). Therefore, we introduce P. endophytica as a novel species bases on morphology and phylogeny.


Index Fungorum number: IF809741; Facesoffungi number: FoF06981
Endophytic in fresh leaves of Magnolia garretti. Colonies on PDA, fluffy appearance, white aerial mycelia. Mycelia superficial and immersed composed of septate, branched, 2–3 μm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial on PDA, globose, solitary or aggregated, embedded or semi-immersed, exuding black conidial masses. Conidia 20–23 × 4–6 μm (μ = 21 × 5 μm), ellipsoid, fusoid, straight to slightly curved, 4-septate; basal cell 3–4 μm long, hyaline, obconic, smooth- and thin-walled; three median cells 14–17 × 4–6 μm (μ = 15 × 6 μm), light brown, doliiform, (the second cell from base 5–6 μm long, third cell 4–5 μm long, fourth cell 5–6 μm long); apical cell 3–5 μm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, smooth- and thin-walled, with 2–3 tubular apical appendages, 18–22 μm long, arising from the apical crest, filiform, basal appendage 2–3 μm long, single, tubular, unbranched, straight and centric.

Culture characteristics – Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin undulate, slightly raised, white mycelia, with black, gregarious conidiomata, colony from above: white; reverse: cream.


GenBank numbers – ITS: MW263950, tef1: MW273931

Notes – The new isolate (GMT13) clustered with the ex-type Pestalotiopsis kenyana (CBS 442.67) in the combined ITS, tub2 and tef1 phylogenetic analysis with 99% ML, 98% MP and 1.00 BYPP support (Fig. 31). ITS and tef1 gene regions of GMT13 is similar to the ex-type P. kenyana (CBS 442.67). The type of P. kenyana was recorded from twigs of Coffea sp. in Kenya (the type CBS H–15657) (Maharachchikumbura et al. 2014b) and on Camellia sinensis in China (Liu et al. 2017). Therefore, we report a new host record considering molecular data and host associations.
<table>
<thead>
<tr>
<th>Species</th>
<th>Conidia size (μm)</th>
<th>Three median cells (μm)</th>
<th>Number of apical appendages</th>
<th>Length of apical appendage/s (μm)</th>
<th>Length of basal appendage (μm)</th>
<th>Host and life mode</th>
<th>Locality</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Pestalotiopsis australis</em></td>
<td>(26–)27–34(–36) × 7–8.5; (x = 30.8 × 7.7)</td>
<td>(16–)17–21(–21.5); (μ = 19.1)</td>
<td>2–3</td>
<td>(11–)12–20(–22); (μ = 15.5)</td>
<td>3–7</td>
<td>Brabejum stellatifolium, Grevillea sp. and Protea neriifolia × susanna</td>
<td>Australia and South Africa</td>
<td>Maharachchikumbura et al. (2014b)</td>
</tr>
<tr>
<td><em>P. brassicae</em></td>
<td>29–30–37(–40) × (8–)8.5–11(–11.5); (μ = 34 × 9.7)</td>
<td>(20–)20.5–24.5(–25); (μ = 22.6)</td>
<td>3–5, (mostly 4)</td>
<td>(27–)28.5–48(–50); μ = 37</td>
<td>10–25</td>
<td>On seeds of Brassica napus</td>
<td>New Zealand</td>
<td>Maharachchikumbura et al. (2014b)</td>
</tr>
<tr>
<td><em>P. chamaeropis</em></td>
<td>(21–)22.5–27(–28) × (6–)7–9(–9.5); (μ = 25.2 × 8)</td>
<td>(15–)16–17.5(–18.5); (μ = 16.7)</td>
<td>2–3, (mostly 3)</td>
<td>13–)14.5–23(–24); μ = 18</td>
<td>4–8.5</td>
<td>On leaves of Chamaerops humilis</td>
<td>Italy</td>
<td>Maharachchikumbura et al. (2014b)</td>
</tr>
<tr>
<td><em>P. endophytica</em></td>
<td>25–30 × 5–7; (μ = 28 × 6)</td>
<td>15–18 × 5–7; (μ = 16 × 6)</td>
<td>2–3</td>
<td>20–23</td>
<td>4–6</td>
<td>Endophytic on healthy leaves of Magnolia candoll</td>
<td>Chiang Mai Province, Thailand</td>
<td>This study</td>
</tr>
<tr>
<td><em>P. hollandica</em></td>
<td>(25–)25.5–33(–34) × 8.5–10(–10.5); (μ = 28 × 9.4)</td>
<td>(16.5–)17–23(–24); (μ = 28 × 9.4)</td>
<td>1–4</td>
<td>20–40; μ = 27</td>
<td>3–9</td>
<td>On Sciadopitys verticillata</td>
<td>Netherlands</td>
<td>Maharachchikumbura et al. (2014b)</td>
</tr>
<tr>
<td><em>P. intermedia</em></td>
<td>24–28 × 5.5–6.5; (μ = 25.7 × 6)</td>
<td>15–19; (μ = 17)</td>
<td>2–3, (rarely 4)</td>
<td>10–28; (μ = 18 × 5)</td>
<td>6–10</td>
<td>Saprobic/endo phytic on unidentified trees,</td>
<td>Hubei, Yunnan Province, China</td>
<td>Maharachchikumbura et al. (2012)</td>
</tr>
<tr>
<td><em>P. italiana</em></td>
<td>26–35 × 8–11; (μ = 30 × 9.6)</td>
<td>18–28; (μ = 23)</td>
<td>2–5, (mostly 3–4)</td>
<td>20–40; (μ = 32)</td>
<td>6–10</td>
<td>Dead twigs of Cupressus glabra</td>
<td>Italy</td>
<td>Liu et al. (2015c)</td>
</tr>
<tr>
<td>Species</td>
<td>Conidia size (μm)</td>
<td>Three median cells (μm)</td>
<td>Number of apical appendages</td>
<td>Length of apical appendage/s (μm)</td>
<td>Length of basal appendage (μm)</td>
<td>Host and life mode</td>
<td>Locality</td>
<td>Reference</td>
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<tr>
<td><em>P. jiangxiensis</em></td>
<td>22–29 × 6–9; ((\bar{x}) = 25.7 × 7.3)</td>
<td>12.5–19; ((\bar{x}) = 15.9)</td>
<td>2–4, (mostly 3)</td>
<td>16.5–32; ((\bar{x}) = 22.4)</td>
<td>6.5–19.5</td>
<td>Camellia sp.</td>
<td>Jiangxi Province, China</td>
<td>Liu et al. (2017)</td>
</tr>
<tr>
<td><em>P. linearis</em></td>
<td>24–33 × 4.7–6; ((\bar{x}) = 29.5 × 5)</td>
<td>17–21; ((\bar{x}) = 19)</td>
<td>2–3, (rarely 1)</td>
<td>10–20</td>
<td>4–7</td>
<td>Endophytic on living leaves of <em>Trachelospermum</em> sp. and <em>Tsuga</em> sp.</td>
<td>Yunnan Province, China</td>
<td>Maharachchikumbura et al. (2012)</td>
</tr>
<tr>
<td><em>P. monochaeta</em></td>
<td>(25–)27–40(–42) × 7–11(–11.5); ((\bar{x}) = 32.8 × 9.6)</td>
<td>(17–)18–25(–26); ((\bar{x}) = 21)</td>
<td>1</td>
<td>(40–)43–67(–75); ((\bar{x}) = 51)</td>
<td>6–14</td>
<td>On <em>Taxus baccata</em> and endophytic in branches of <em>Quercus robur</em> sp.</td>
<td>Netherlands</td>
<td>Maharachchikumbura et al. (2014b)</td>
</tr>
<tr>
<td><em>P. scoparia</em></td>
<td>(22–)23.5–29(–31) × 6–8.5; ((\bar{x}) = 26.3 × 7.4)</td>
<td>15.5–19.5; ((\bar{x}) = 17)</td>
<td>3–5</td>
<td>(20–)23–35(–42); ((\bar{x}) = 29.6)</td>
<td>9–25</td>
<td><em>Chamaecyparis</em> sp.</td>
<td>Unknown</td>
<td>Maharachchikumbura et al. (2014b)</td>
</tr>
<tr>
<td><em>P. unicolor</em></td>
<td>20–24.5 × 4–6; ((\bar{x}) = 22.5 × 1)</td>
<td>13–16; ((\bar{x}) = 14 × 7)</td>
<td>2–3</td>
<td>11–20; ((\bar{x}) = 17.5)</td>
<td>4–10</td>
<td>Endophytic on <em>Rhododendron</em> sp.</td>
<td>Hunan Province, China</td>
<td>Maharachchikumbura et al. (2012)</td>
</tr>
<tr>
<td><em>P. verruculosa</em></td>
<td>28–35 × 9–11; ((\bar{x}) = 30.6 × 10.3)</td>
<td>18–26; ((\bar{x}) = 21.6)</td>
<td>2–6, (mostly 3–4)</td>
<td>25–40; ((\bar{x}) = 34)</td>
<td>8–12</td>
<td>Endophytic on living leaf of <em>Rhododendron</em> sp.</td>
<td>Yunnan Province, China</td>
<td>Maharachchikumbura et al. (2012)</td>
</tr>
</tbody>
</table>
Figure 33 – Pestalotiopsis kenyana (MFLUCC 20–0143, new host record). a Conidial masses on PDA. b, c Conidia. Scale bars: b, c = 10 μm.

*Pestalotiopsis neolitseae* Ariyawansa & K.D. Hyde Mycosphere 9 (5): 1005 (2018) Fig. 34

Index Fungorum number: IF827598; Facesoffungi number: FoF04938

*Endophytic* in fresh leaves of *Magnolia garrettii*. Colonies on PDA slightly raised, white mycelia. *Mycelia* superficial and immersed composed of septate, branched, 2–3 μm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: *Conidiomata* pycnidial on pine needle on PDA, globose, solitary or aggregated, embedded or semi-immersed, exuding black conidial masses. Conidia 17–20 × 4–6 μm (\(\bar{x} = 18 \times 5 \mu m\)), ellipsoid, fusoid, straight to slightly curved, 4-septate; basal cell 2–3 μm long, hyaline, obconic, smooth- and thin-walled; three median cells 11–13 × 3–5 μm, (\(\bar{x} = 12 \times 4 \mu m\)) light brown, doliiform, (the second cell from base 4–5 μm long, third cell 3–4 μm long, fourth cell 4–5 μm long); apical cell 2–3 μm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, smooth- and thin-walled, with 2–3 tubular apical appendages, 10–15 μm long, arising from the apical crest, filiform, basal appendage 2–3 μm long, single, tubular, straight and centric.

Culture characteristics – Colonies on PDA reaching 30 mm diameter after 7 days at 25°C, colonies circular, margin undulate, slightly raised, white mycelia, with black, gregarious conidiomata, colony from above: white; reverse: cream.

Material examined – THAILAND, Chiang Mai Province, healthy leaves of *Magnolia garrettii* (Magnoliaceae), 26 April 2017, N. I. de Silva, GMT1, living culture, MFLUCC 18–0930, GMT2, living culture, MFLUCC 18–0931.

GenBank numbers – (GMT1); ITS: MW263951, *tef1*: MW273932, (GMT2); ITS: MW263952, *tef1*: MW273933

Figure 34 – Pestalotiopsis neolitseae (MFLUCC 18–0930, new host and geographical record). a Conidial masses on pine needles. b, c Conidia. Scale bars: b, c = 10 μm.
Notes – As morphological characters examined largely overlap with the type of *P. neolitseae* (NTUH 17–011), we report our collections (GMT 1 and GMT 2) as a new host record from healthy leaves of *Magnolia candolli*. Both share a similar morphology in having concolourous median cells and 1–3 tubular apical appendages (Ariyawansa & Hyde 2018). The multi-gene (*ITS, tub2* and *tef1*) phylogeny reported herein, also showed that our collections clusters with the ex-type of *P. neolitseae* (NTUCC 17–011) (Fig. 31). *Pestalotiopsis neolitseae* have been recorded from leaf spots of *Neolitsea villosa* as pathogen in Taiwan (Ariyawansa & Hyde 2018) and this is the first record of *P. neolitseae* associated with healthy leaves of *M. candolli* as an endophyte.

**Pseudopestalotiopsis** Maharachch., K.D. Hyde & Crous

The genus was introduced by Maharachchikumbura et al. (2014b) with the type *Pseudopestalotiopsis theae*. *Pseudopestalotiopsis* species are appendage-bearing phenotypically diverse coelomycetes in Sporocadaceae and are commonly found in tropical and subtropical ecosystems (Jaklitsch et al. 2016, Maharachchikumbura et al. 2016a, Jayawardena et al. 2019). *Pseudopestalotiopsis* is characterized by brown to dark brown or olivaceous median cells (concolourous median cells) and knobbed or not knobbed apical appendages (Maharachchikumbura et al. 2014b, 2016a). *Pseudopestalotiopsis* species are considered as plant pathogens; *P. theae* causes grey blight of tea (Maharachchikumbura et al. 2011, 2016a) and *P. ixorae* and *P. taiwanensis* cause a leaf spot (Tsai et al. 2018). *Pseudopestalotiopsis* species are exhibit saprobic and endophytic life modes. *Pseudopestalotiopsis theae* isolated as an endophyte from different hosts (*Camellia nitidissima, C. sinensis, Holarrhena antidisenterica, Podocarpus macrophyllus, Terminalia arjuna*) or as a saprobe (seeds of *Diospyros crassiflora*) (Maharachchikumbura et al. 2011, 2016a, Jayawardena et al. 2019). There are 22 species epithets are recorded in Index Fungorum (2021). In this study, we report two new host records of endophytic *P. ampullacea* and *P. simitheae* from *Magnolia candolli* in China.


Index Fungorum number: IF818922; Facesoffungi number: FoF09445

Endophytic in fresh leaves of *Magnolia candolli*. Colonies on PDA flat, white mycelia. *Mycelia* superficial and immersed composed of septate, branched, 2–3 µm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: *Conidiomata* pycnidial on PDA, globose, solitary, embedded or semi-immersed, exuding black conidial masses. *Conidia* 25–28 × 5–7 µm (X = 26 × 6 µm), ellipsoid, fusoid, straight to slightly curved, 4-septate; basal cell 3–4 µm long, hyaline, obconic, smooth- and thin-walled; three median cells 17–19 × 5–7 µm, (X = 18 × 6 µm) light brown, doliform, (the second cell from base 6–7 µm long, third cell 5–6 µm long, fourth cell 6–7 µm long); apical cell 3–4 µm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, thin-walled, with 2–3 tubular apical appendages, 24–27 µm long, arising from the apical crest, filiform, basal appendage 2–3 µm long, single, tubular, straight and centric.

Culture characteristics: Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, fluffy appearance, white mycelia, colony from above: white; reverse: cream.


Notes – Phylogenetic analysis of combined ITS, tub2 and tef1 sequence data confirmed that our two isolates (S9 and S23) are *P. ampullacea* with 99% ML, 98% MP and 1.00 BYPP support (Fig. 35). The ITS gene region of the ex-type *P. ampullacea* (LC6618) and two new strains are similar. There are two base pair differences in each tub2 and tef1 gene regions of the ex-type *P. ampullacea* (LC6618) and the new strain S9 and S23. *Pseudopestalotiopsis ampullacea* (HMAS
247056, holotype) was isolated from on *Camellia sinensis* in China (Liu et al. 2017). *Pseudopestalotiopsis ampullacea* was also recorded from plant family Lauraceae (Liu et al. 2017). We hence, identify our collection as a new host record of *P. ampullacea*, isolated from leaves of *Magnolia candolli*.

**Figure 35** – Phylogram generated from maximum likelihood analysis based on combined ITS, *tub2* and *tef1* sequence data alignment. Twenty eight strains are included in the combined gene analyses comprising 1880 characters after alignment (500 characters for ITS, 440 characters for *tub2* and 940 characters for *tef1*). *Neopestalotiopsis clavispora* (MFLUCC 12-0281) is used as outgroup taxon. The best RAxML tree with a final likelihood value of -5232.29747 is presented. The matrix had 376 distinct alignment patterns, with 19.27% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.252534, C = 0.258358, G = 0.234842, T = 0.254266; substitution rates AC = 1.785859, AG = 4.601061, AT = 2.011516, CG = 1.468089, CT = 5.532301, GT = 1.000000; gamma distribution shape parameter α = 0.942725. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior
probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex-type strains are in bold.

Figure 36 – *Pseudopestalotiopsis ampullacea* (MFLUCC 19–0100, new host record). a Conidial masses on PDA. b, c Conidia. Scale bars: b, c = 10 μm.

*Pseudopestalotiopsis simitheae* (Yu Song, Tangthir., K.D. Hyde & Y. Wang) Maharachch. & K.D. Hyde, Mycological Progress 15(no. 22): 5 (2016) Fig. 37

Index Fungorum number: IF551720; Facesoffungi number: FoF01631

*Endophytic* in fresh leaves of *Magnolia candolli*. Colonies on PDA flat, white mycelia. *Mycelia* superficial and immersed composed of septate, branched, 2–3 μm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial on PDA, globose, solitary or aggregated, embedded or semi-immersed, exuding black conidial masses. *Conidia* 20–25 × 5–7 μm (x = 23 × 6 μm), ellipsoid, fusoid, straight to slightly curved, 4-septate; basal cell 3–4 μm long, hyaline, obconic, smooth- and thin-walled; three median cells 15–19 × 5–7 μm (x = 16 × 6 μm), light brown, doliiform, (the second cell from base 5–6 μm long, third cell 5–6 μm long, fourth cell 5–7 μm long); apical cell 3–4 μm long, cylindrical to subcylindrical, conic to bell-shaped, hyaline, thin-walled, with 3 tubular apical appendages, 22–27 μm long, arising from the apical crest, filiform, basal appendage 2–3 μm long, single, tubular, straight and centric.

Culture characteristics – Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, fluffy appearance, white mycelia, colony from above: white; reverse: cream.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, S224 (MFLU 20–0603, dried culture); living culture, KUMCC 17–0255.


Notes – Phylogenetic analyses of concatenated ITS, *tub2* and *tef1* sequence dataset indicated that our isolate has a close affinity with *Pseudopestalotiopsis simitheae* with 67% ML, 89% MP, 1.00 BYPP support (Fig. 35). Comparisons of ITS and *tub2* gene regions revealed one base pair difference in each gene region between the ex-type *P. simitheae* (MFLUCC 12–0121) and new isolate S224. There are 10 base pair differences (substitutions) in *tef1* gene regions between the ex-type *P. simitheae* (MFLUCC 12–0121) and the new strain S224 without gaps. Furthermore, the new isolate (S224) shares a size range of conidia (26–30 × 5–7 μm) conidial characters with the holotype *P. simitheae* (MFLU 13–0305) (22–30 × 5–6.5 μm). *Pseudopestalotiopsis simitheae* was collected from dead and living leaves of *Pandanus odoratissimus* in Thailand (Song et al. 2014b). Based on similar morphology and molecular data, here, we report our isolate as a new host and geographical record of *P. simitheae* associate with healthy leaves of *Magnolia candolli* in China.
Figure 37 – *Pseudopestalotiopsis simitheae* (KUMCC 17–0255, new host and geographical record). a Conidial masses on PDA. b, c Conidia. Scale bars: b, c = 10 μm.

**Xylariales** Nannf.  
**Apiosporaceae** K.D. Hyde et al.

Hyde et al. (1998) established Apiosporaceae with the type genus *Apiospora*. Species of this family are saprobes, pathogens, endophytes of plants and occasionally infecting humans (Wang et al. 2017b, Raza et al. 2019, Hyde et al. 2020b, c). Maharachchikumbura et al. (2016b) accepted six genera in Apiosporaceae. Hyde et al. (2020b) accommodated five genera, namely, *Appendicospora*, *Arthrinium*, *Dictyoarthrinium*, *Endocalyx* and *Nigrospora*. Asexual morph of this family are considered as coelomycetous or hyphomycetous comprising sporodochial conidiomata, smooth to finely verruculose conidiogenous cells, unicellular or guttulate to granular conidia (Maharachchikumbura et al. 2016b, Raza et al. 2019, Hyde et al. 2020c). The sexual morphs are characterized by perithecial ascomata immersed in pseudostromata, unitunicate or short-pedicellate asci, smooth-walled ascospores (Maharachchikumbura et al. 2016b, Raza et al. 2019).

**Nigrospora** Zimm.

*Nigrospora* species have been isolated as endophytes from leaves and stems of various plants, or as saprobes from detritus, dead larvae or leaf litter (Mason 1927, Uzor et al. 2015, Wang et al. 2017b). *Nigrospora* is also known as plant pathogen causing leaf spots, leaf blight and squirter disease in many important economic crops, fruits and ornamentals (Wang et al. 2017b) while these fungi cause onychomycosis, hay fever, respiratory and allergic diseases in humans (Raza et al. 2019). *Nigrospora* has a cosmopolitan distribution and a wide host range (Wang et al. 2017b). Asexual morph is characterized by branched micronematous or semimacronematous conidiophores, monoblastic conidiogenous cells and black, shiny, aseptate conidia and sexual morph comprises perithecial ascomata, short–stalked asci with biseriate ascospores (Webster 1952, Wang et al. 2017b, Raza et al. 2019). There are 34 *Nigrospora* epithets in Index Fungorum (2021).

Index Fungorum number: IF820731; Facesoffungi number: FoF09396


Culture characteristics – Colonies on PDA reaching 36 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, fluffy appearance, white mycelia, colony from above: white; reverse: cream.
Figure 38 – Phylogram generated from maximum likelihood analysis based on combined ITS, tub2 and tef1 sequence data. Related sequences of *Nigrospora* were obtained from Raza et al. (2019). Sixty six strains are included in the combined gene analyses comprising 1360 characters after alignment (600 characters for ITS, 430 characters for *tub2* and 330 characters for *tef1*). *Arthrinium malaysianum* (CBS 102053) is used as outgroup taxon. The best RAxML tree with a final likelihood value of -9137.301167 is presented. The matrix had 657 distinct alignment patterns, with
16.46% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.212749, C = 0.307101, G = 0.241426, T = 0.238724; substitution rates AC = 0.995728, AG = 2.485554, AT = 0.913276, CG = 0.873873, CT = 4.084712, GT = 1.000000; gamma distribution shape parameter α = 0.714119. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequences are indicated in red. Type and ex–type strains are in bold.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, S25 (MFLU 20–0599, dried culture); living culture, MFLUCC 19–0114 = KUMCC 17–0248.


Notes – The combined ITS, tub2 and tef1 phylogeny herein, showed that the new isolate (S25) clustered closely related to the ex-type Nigrospora camelliae-sinensis (CGMCC3.18125) and five strains of N. camelliae-sinensis with 74% ML, 75% MP and 0.96 BYPP support (Fig. 38). Conidia of the type N. camelliae-sinensis are spherical or slightly ellipsoidal (Wang et al. 2017b) while conidia of new isolate are spherical. Spherical conidia of new isolate (S25) (8–10 × 8–9μm) are smaller than the type N. camelliae-sinensis (13–18 μm diam.) (Wang et al. 2017b).

There are 7 base pair differences (7/220 = 3.18%) between the ex-type N. camelliae-sinensis and new isolate (S25) in tef1 sequence data. A base pair comparison of ITS and tub2 sequence data between the ex-type N. camelliae-sinensis and S25 revealed 100% similarity.

Nigrospora camelliae-sinensis was introduced by Wang et al. (2017b), which was collected from Guangxi Province, China, on Camellia sinensis. In addition to that N. camelliae-sinensis was recorded from leaves and roots of Saccharum officinarum in China (Raza et al. 2019). The new isolate (S25) was isolated from healthy leaves of Magnolia candolli in Yunnan Province, China. We consider the new isolate as N. camelliae-sinensis and the first report of N. camelliae-sinensis from leaves of Magnolia candolli.

Figure 39 – Nigrospora camelliae-sinensis (MFLUCC 19–0114, new host record). a Colonies on PDA. b Conidiogenous cell with conidia. c Mycelia and conidia. Scale bars: b, c = 5 μm.

Nigrospora chinensis Mei Wang & L. Cai, Persoonia 39: 129 (2017) Fig. 40

Culture characteristics – Colonies on PDA reaching 30 mm diameter after 7 days at 25°C, colonies circular, margin entire, slightly raised, wooly appearance, white aerial mycelia, colony from above: white; reverse: cream.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli (Magnoliaceae), 26 April 2017, N. I. de Silva, S26 (MFLU 20–0600, dried culture); living culture, MFLUCC 19–0115 = KUMCC 17–0249, S126, living culture, KUMCC 17–0253, MT 4, living culture, KUMCC 17–0210 = MFLUCC 19–0046.

GenBank numbers – (S26); ITS: MW285153, tub2: MW620879, tef1: MW296113, (S126); ITS: MW285154, tub2: MW620880, (MT4); ITS: MW285152, tub2: MW620881.

Notes – Three new strains clustered in a well-supported clade with the ex-type Nigrospora chinensis (CGMCC3.18127). Nigrospora chinensis has recorded from various host plants, i.e. Aucuba japonica, Camellia sinensis, Castanopsis sp., Lindera aggregate, Machilus duthiei, Musa paradisiaca, Osmanthus sp. and Smilax ocreata in China (Wang et al. 2017b). Therefore, we report our new collection as endophytic lifestyle of N. chinensis, which was isolated from healthy leaves of Magnolia candolli in Yunnan, China for the first time.

Figure 40 – Nigrospora chinensis (MFLUCC 19–0115, new host record). a Colony on PDA producing conidia masses. b Conidiogenous cell with immature conidium. c Mycelia and conidia. Scale bars: b, c = 10 μm.

Nigrospora magnoliae N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.

Index Fungorum number: IF556382; Facesoffungi number: FoF09497

Etymology – Name reflects the host genus Magnolia, from which the new species was isolated.

Holotype – MFLU 20–0597

Endophytic in fresh leaves of Magnolia candolli. Colonies on PDA brown, flat, spreading, with abundant aerial mycelia. Mycelia superficial and immersed composed of septate, branched, 2–4 μm wide, hyaline and brown with smooth and thick-walled hyphae. Sexual morph: Undetermined. Asexual morph: hyphomycetous, conidial masses with mycelia in culture on PDA. Conidiophores 2–4 μm diam., hyaline, micronematous or semi-macronematous, flexuous or straight, smooth. Conidiogenous cells 5–7× 5–6 μm (x = 6 × 5.4 μm), hyaline, monoblastic, discrete, solitary, determinate, doliiform to amphotriiform. Conidia 10–14 × 10–13 μm (x = 12 × 11 μm), dark brown, solitary, globose or subglobose and smooth-walled.

Culture characteristics – Colonies on PDA reaching 30 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, initially white and becoming dark brown, some white aerial mycelia, colony from above: brown; reverse: dark brown.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli L. (Magnoliaceae), 26 April 2017, N. I. de Silva, S22 (MFLU 20–0597, holotype, dried culture); ex-type living culture, MFLUCC 19–0112 = KUMCC 17–0246.

Additional material – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of Magnolia candolli L. (Magnoliaceae), 26 April 2017, N. I. de Silva, S5, living culture, KUMCC 17–0231.
GenBank numbers – (S22); ITS: MW285092, tub2: MW438334, (S5); ITS: MW699933, tub2: MW727265.

Notes – Phylogenetic results indicated that Nigrospora magnoliae (S22) clustered with other Nigrospora species and in particular, and formed a distinct clade sister to N. camelliae-sinensis, N. singularis and N. pyriformis with 61% ML, 50% MP and 0.75 BYPP support (Fig. 38). Nigrospora magnoliae has dark brown, globose or subglobose (10–14 × 10–13 μm) conidia that are slightly smaller than N. camelliae-sinensis (13–18 μm diam. spherical and 12–18 × 9–14.5 μm ellipsoidal conidia) N. pyriformis (12.5–16.5 μm globose to subglobose and 17.5–27.5 × 10–18.5 μm pyriform conidia) (Wang et al. 2017b) and N. singularis (9.5–13 × 11–15 μm ellipsoid to subglobose conidia) (Raza et al. 2019). Nigrospora sp. 2 (LC6702) clustered with N. magnoliae in our combined multi-gene phylogenetic analysis. Wang et al. (2017b) recorded Nigrospora sp. 2 (LC6702) in their phylogenetic analyses without morphological description, which was isolated from Camellia sinensis in China. Therefore, based on phylogenetic analyses and morphology, N. magnoliae identified as a novel endophytic species in this study.

Figure 41 – Nigrospora magnoliae (MFLU 20–0597, holotype). a Colonies on PDA. b Colony on PDA producing conidia masses. c Mycelia and conidia d, e Conidiogenous cells with conidia. f Mycelium and conidia. Scale bars: c–f = 10 μm.

Nigrospora musae McLennan & Hoëtte, Australian Institute of Science and Industry Reserch Bulletin 75: 15 (1933) Fig. 42

Index Fungorum number: IF271866; Facesoffungi number: FoF09447


Culture characteristics – Colonies on PDA reaching 35 mm diameter after 7 days at 25°C, colonies circular, margin entire, slightly raised, woolly appearance, white and brown mycelia, colony from above: brown and white; reverse: brown.
Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, S4 (MFLU 20–0585, dried culture); living culture, MFLUCC 19–0098 = KUMCC 17–0230.


Notes – Phylogram generated from a combined ITS, *tub2* and *tef1* sequence dataset indicated that our strain is *Nigrospora musae* and it formed a well-supported clade with 95% ML, 100% MP, 1.00 BYPP support (Fig. 38). McLennan & Hoëtte (1993) originally described *N. musae* from fruits of *Musa sapientum* in Australia. The ex-type strain (CBS 319.34) was sterile and therefore, Wang et al. (2017b) emended taxonomic description based on a freshly collected strain from *Camellia sinensis* in China. This species has not been reported from the plant family Magnoliaceae (Farr & Rossman 2021). Herein, we provide the new host record for *N. musae* from healthy leaves of *Magnolia candolli*.

**Figure 42** – *Nigrospora musae* (MFLUCC 19–0098, new host record). a Colonies on PDA producing conidia. b, c Mycelia and young developing conidia. Scale bars: b, c = 10 μm.

*Nigrospora sphaerica* (Sacc.) E.W. Mason, Transactions of the British Mycological Society 12(2–3): 158 (1927)  

Index Fungorum number: IF254776; Facesoffungi number: FoF06599


Culture characteristics – Colonies on PDA reaching 35 mm diameter after 7 days at 25°C, colonies circular, margin entire, slightly raised, woolly appearance, white mycelia, colony from above: white; reverse: light brown.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, S12 (MFLU 20–0589, dried culture); living culture, MFLUCC 19–0103 = KUMCC 17–0237.


Notes – Based on combined multi-gene phylogenetic analysis, the new strain S12 nested with other strains of *Nigrospora sphaerica* with 100% ML, 98% MP, 0.99 BYPP support (Fig. 38). *Nigrospora sphaerica* has a wide host range in China (i.e. on *Camellia sinensis*, *Cleveya japonica*, *Deutzia* sp., *Harpullia longipetala*, *Musa paradisiaca*, *Nelumbo* sp., *Rhododendron arboretum*, *Rosa* sp., *Saccharum officinarum* and submerged wood) (Wang et al. 2017b, Raza et al. 2019). *Nigrospora sphaerica* is a well-known pathogen associated with leaf blight on *Camellia sinensis* in...
China (Liu et al. 2015b), leaf spots and twig and shoot blight on blueberries (*Vaccinium corymbosum*) (Wright et al. 2008). *Nigrospora sphaerica* is also considered as an opportunistic pathogen causing onychomycosis in humans (de Hoog et al. 2000, Fan et al. 2009) and a corneal ulcer (Kindo et al. 2014). The endophytic lifestyle of *N. sphaerica* was recorded from different plant species i.e. *Vinc a rosea* (Apocynaceae) (Metwaly et al. 2014), roots of *Smallanthus sonchifolius* (Asteraceae) (Lopes & Pupo 2011), leaves of *Eupatorium adenophorum* (Asteraceae) (Chen et al. 2016). *Nigrospora sphaerica* was isolated from healthy leaves of *Magnolia candolli* (Magnoliaceae) is regarded as a new host record.

**Figure 43** – *Nigrospora sphaerica* (MFLUCC 19–0103, new host record). a Colony on PDA producing conidia masses. b, c Mycelia and developing conidia. Scale bars: b, c = 10 μm.

**Phylum Basidiomycota** Whittaker ex R.T. Moore  
**Class Agaricomycetes** Doweld  
**Agaricales** Underw  
**Psathyrellaceae** Vilgalys, Moncalvo & Redhead  
Deliquescent type mushroom fruiting bodies are found in this family that they have a special autodigestive phase of ontogeny (hence, the name “deliquescent”) (Nagy et al. 2011, 2013). The type genus of the family is *Psathyrella* and the members of this genus are mainly decomposers of leaf-litter or wood, more rarely living on dung, or parasitizing other fungi (Nagy et al. 2013). Traditionally, the family contained two large genera, *Coprinus* and *Psathyrella* (Nagy et al. 2013). *Ozonium* and *Hormographiella* are the conidial anamorphic taxa in the family that recently reclassified as *Coprinellus* (Nagy et al. 2013). Nagy et al. (2013) revised the taxonomy of the family using ITS sequences data and distinguished 14 major clades within Psathyrellaceae.

**Coprinellus** P. Karst.

*Coprinellus* is one of the most species-rich, coprinoid mushroom genera in Psathyrellaceae that include approximately 80 species (Gomes & Wartchow 2014, Hussain et al. 2018) while Index Fungorum (2021) lists 94 species epithets. Species-level identification of *Coprinellus* is based mainly on the presence or absence and the structure of veil and cystidia on the pileus, of cystidia on the lamellae and on basidiospore morphology and phylogenetic analyses of ITS sequence data (Hussain et al. 2018). *Coprinellus* species are common saprotroph of wood chips, leaf-litter and herbivore dung (Hussain et al. 2018).

**Coprinellus magnoliae** N.I. de Silva, Lumyong & K.D. Hyde, sp. nov.  
**Index Fungorum number:** IF556385; Facesoffungi number: FoF09498  
**Etymology** – Name reflects the host genus *Magnolia*, from which the new species was isolated.  
**Holotype** – MFLU 20–0605  
Endophytic in fresh leaves of *Magnolia candolli*. Colonies on PDA white aerial mycelia. *Mycelia* superficial and immersed composed of septate, branched, 1–2 μm wide, hyaline and light brown, with smooth and thick-walled hyphae. *Spores* 2–3.5 × 1–2 μm (\(\bar{x} = 3 \times 1.4 \mu m\)), hyaline, sub-globose to fusiform with smooth-walled.
Figure 44 – Phylogram generated from maximum likelihood analysis based on ITS sequence data. Related sequences representing *Coprinellus* species were obtained from Hussain et al. (2018). Eighty eight strains are included in the phylogenetic analyses comprising 700 characters after ITS alignment *Psathyrella candolleana* (CBS 300.47) is used as the outgroup taxon. The best RAxML tree with a final likelihood value of -5818.670832 is presented. The matrix had 462 distinct alignment patterns, with 17.55% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.244768, C = 0.236272, G = 0.230683, T = 0.288278; substitution rates AC = 2.005058, AG = 2.692844, AT = 2.306751, CG = 1.041461, CT = 5.240997, GT = 1.000000; gamma distribution shape parameter α = 0.548187. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal or
greater than 0.95 are placed above the branches. The newly generated sequence is indicated in red. Type and ex-type strains are in bold.

Figure 44 – Continued.
Culture characteristics – Colonies on PDA reaching 30 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, white, some white aerial mycelia, colony from above: white; reverse: cream.

Material examined – THAILAND, Chiang Mai Province, Mueang Chiang Mai District, healthy leaves of Magnolia garrettii (Magnoliaceae), 14 August 2017, N. I. de Silva, GMT25 (MFLU 20–0605, holotype, dried culture); ex-type living culture, MFLUCC 18–0942.

GenBank numbers – ITS: MW244022.

Notes – Phylogenetic analyses of ITS sequence data revealed that Coprinellus magnoliae forms a sister lineage with C. disseminates-similis and distinctly separate from C. disseminates-similis with 100% ML, 100% MP, 1.00 BYPP support (Fig. 44). A comparison of ITS sequence shows that C. magnolia differs from C. disseminates-similis (Cr 3w) in 28 base pair positions (28/600 = 4.67%). The important characteristics of C. disseminatus–similis are pileus parabolic to campanulate, greyish-brown, with umbonate centre; surface pruinose to pulverulent, with sparse micaceous-glistening veil, bright white, deeply plicate from centre to margin (Hussain et al. 2018). Basidiopores of C. disseminates-similis are on average 8.5 × 5.2 × 4.9 μm, ellipsoid to cylindrical or obovoid in face view; ellipsoid to amygdaliform in side view (Hussain et al. 2018). However, C. magnoliae formed hyaline, sub-globose to fusiform, (2–3.5 × 1–2 μm), sized spores on PDA media. C. disseminates-similis was found on leaf litter under Populus alba and Morus alba, in lowland northern Pakistan (Hussain et al. 2018). Coprinellus magnoliae was isolated from healthy leaves of Magnolia garrettii in Chiang Mai Province, Thailand. We introduce C. magnoliae as a new species based on phylogeny and morphology.

Figure 45 – Coprinellus magnoliae (MFLU 20–0605, holotype). a Mycelia on PDA. b Mycelia on a rice straw. c Mycelia masses. d Developing spores from mycelia. e, f Spores. Scale bars: c = 10 μm, d, f = 5 μm, f = 2 μm.

Polyporales Gäum.
Phanerochaetaceae Jülich

Taxa of this family are mainly saprobic and some species may also occur as parasites and were recorded worldwide from different dead plants (Larsson 2007, Miettinen et al. 2016). Larsson (2007) suggested accommodating corticioid fungi within Phanerochaete in Phanerochaetaceae. Binder et al. (2013) included the polypore genus Bjerkandera in Phanerochaetaceae. The recent taxonomic revision of Miettinen et al. (2016) accepted 14 genera of corticioid and poroid species in Phanerochaetaceae.
Phanerina Miettinen

The polypore genus *Phanerina* belongs to Phanerochaete clade in Phanerochaetaceae according to phylogenetic analyses (Miettinen et al. 2016). Members of genus *Phanerina* grow on dead dicot trees (Miettinen et al. 2016). *Phanerina* is a monotypic genus closely related to *Riopa* by both morphology and phylogeny (Miettinen et al. 2016).

Figure 46 – Phylogram generated from maximum likelihood analysis based on combined ITS and LSU sequence data. Related sequences representing Phanerochaete clade in Phanerochaetaceae
were obtained from Miettinen et al. (2016). Thirty eight strains are included in the combined gene analyses comprising 1750 characters after alignment (750 characters for ITS and 1000 characters for LSU). *Donkia pulcherrima* (Hausknecht Kovac 6.VII.1998) is used as the outgroup taxon. The best RAxML tree with a final likelihood value of -6571.270371 is presented. The matrix had 471 distinct alignment patterns, with 56.37% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.250421, C = 0.215000, G = 0.265827, T = 0.268752; substitution rates AC = 1.280665, AG = 3.466017, AT = 1.741286, CG = 0.529488, CT = 5.989708, GT = 1.000000; gamma distribution shape parameter α = 0.567128. Bootstrap values for maximum likelihood and maximum parsimony equal to or greater than 75% and Bayesian posterior probabilities equal to or greater than 0.95 are placed above the branches. The newly generated sequence is indicated in red. Type and ex-type strains are in bold.

**Phanerina mellea** (Berk. & Broome) Miettinen, MycoKeys 17: 22 (2016)  

*Index Fungorum number: IF811537; Facesoffungi number: FoF09448*

*Endophytic in fresh leaves of Magnolia candolli.* Colonies on PDA white aerial mycelia and brown spore masses. *Mycelia* superficial and immersed composed of septate, branched, 1–2 μm wide, hyaline and light brown, with smooth and thick-walled hyphae. *Spores* 2–3 μm diameter, initially hyaline, becoming brown at maturity, globose, smooth-walled.

Culture characteristics – Colonies on PDA reaching 24 mm diameter after 7 days at 25°C, colonies circular, margin entire, flat, fluffy appearance, white mycelia, colony from above: white; reverse: cream.

Material examined – CHINA, Yunnan Province, Xishuangbanna, healthy leaves of *Magnolia candolli* (Magnoliaceae), 26 April 2017, N. I. de Silva, S19 (MFLU 20–0594, dried culture); living culture, MFLUCC 19–0109 = KUMCC 17–0244.


Notes – *Polyporus melleus* was synonymized as *Phanerina mellea* by Miettinen et al. (2016) based on phylogenetic analyses and morphological studies. The type was described from dead wood in Sri Lanka (Miettinen et al. 2016). *Phanerina mellea* was also recorded from East Africa (Tanzania, Kenya), Japan (Okinawa), and Indonesia (New Guinea) (Miettinen et al. 2016). *Phanerina mellea* grows on dead dicot trees, both standing and fallen, often in sun-exposed habitats (Miettinen et al. 2016). According to the combined ITS and LSU analyses, our endophytic isolate clustered with other isolates of *P. mellea* with 99% ML, 88 MP, 1.00 BYPP support (Fig. 46). Therefore, this study is the first report of *P. mellea* as an endophyte occurring on *Magnolia candolli* in Yunnan Province, China.

**Figure 47** – *Phanerina mellea* (MFLUCC 19–0109, new host record). a Colonies on PDA with mycelia. b Mycelia masses and spores. c = Spores. Scale bars: b, c = 5 μm.

**Discussion**

This study investigated fungal endophytes associated with healthy leaves of *Magnolia candolli* in Yunnan Province, China and *M. garrettii* in Chiang Mai Province, Thailand. The present collection of fungal endophytes comprises both Ascomycota and Basidiomycota. A total of
56 fungal endophyte isolates were obtained. Among them, two isolates were Basidiomycota and 54 isolates were Ascomycota. Considering the endophytic nature of isolates, higher numbers of samples were Ascomycetes, comprising eight isolates of Dothideomycetes and 46 isolates of Sordariomycetes. The current collection consists of eight new species and 23 new host and or geographical records. In addition, de Silva et al. (2019b) introduced Lasiodiplodia endophytica from Magnolia candollii in Yunnan Province, China and two host records of endophytic Lasiodiplodia thailandica and L. pseudeurobromae from Magnolia candollii.

Promputtha et al. (2005) studied endophytes of leaves of Magnolia liliifera, a commonly found tree in Doi Suthep-Pui National Park, Thailand. They isolated different endophytic fungal species such as Bionectria sp., Colletotrichum sp., Diaporthe sp., Hypoxylon sp., Massarina sp. and Xylaria sp. Promputtha et al. (2007) examined the relationships of fungal saprotrophs and endophytes isolated from Magnolia liliifera in Chiang Mai, Thailand. They identified endophytic Colletotrichum gloeosporioides, Colletotrichum sp., Corynespora cassicola, Diaporthe sp., Fusarium oxysporum, Fusarium sp., Guignardia mangiferae and Leptosphaeria sp. In the current study, we identified Colletotrichum gloeosporioides and Corynespora cassicola, similar to the study of Promputtha et al. (2007).

Colletotrichum and Diaporthe were found asymptomatic in plant leaves in the current study of Magnolia garrettii in Thailand as well as the previous studies of Promputtha et al. (2005, 2007) on Magnolia liliifera in Thailand. Nonetheless, our collection comprised genera including Cladosporium, Coprinellus, Epicoccum, Leptosphaerulina, Letendreae, Neopestalotiopsis, Nigrospora, Pestalotiopsis, Pseudopestalotiopsis and Phanerina (isolated from Magnolia garrettii in Thailand) that differ from the previous studies of Promputtha et al. (2005, 2007). In the current collection, we identified common occurrence of endophytic Colletotrichum and Diaporthe on Magnolia garrettii in Chiang Mai, Thailand and Magnolia candollii in Yunnan Province, China. Apart from that different endophytic fungal genera were recorded between Thailand and China from Magnolia garrettii and M. candollii. Endophytic species were found in Magnolia garrettii in Chiang Mai, Thailand, such as Coprinellus and Pestalotiopsis and some species were found in Magnolia candollii in Yunnan Province, China, such as Cladosporium, Corynespora, Epicoccum, Leptosphaerulina, Letendreae, Neopestalotiopsis, Nigrospora, Phanerina and Pseudopestalotiopsis. These data suggest that two Magnolia hosts in Thailand and China are inhabited by different fungal species while few fungal species common in both hosts. Moreover, fungal species of Magnolia liliifera from previous studies (Promputtha et al. 2005, 2007) and M. garrettii in the current study in Chiang Mai, Thailand are different and only a few fungal species are found common to two host plants.

An endophytic community typically comprises common (generalist) and rare species (host specificity and geographic structure are difficult to assess) (Arnold 2007). According to the current study and previous studies of Magnolia host plants (Promputtha et al. 2005, 2007), common endophytic genera are Colletotrichum, Corynespora and Diaporthe whereas rare species containing genera are Coprinellus, Epicoccum and Letendreae when considering the current investigation and previous studies by Promputtha et al. (2005, 2007). It is hypothesized that each tree species in each locality has a signature community of endophytic symbionts (Hoffman & Arnold 2008). The abundance, diversity, and species composition of endophytic assemblage can be influenced by host plants species, the geographic region, microhabitat and microclimatic conditions (Arnold 2007). Leaves are one of the biologically active tissues in plants and they have the most dynamic interfaces between plants and their environment (Arnold 2007). Different studies agree that endophytic colonization of leaves might vary from one host individual to another (Rashmi et al. 2019). For example, Rodrigues (1994) reported only 30% of leaf colonization in Amazonian palms (Arnold et al. 2003) while it was found up to 73.9% of leaf colonization in Manilkara bidentata. The surface area of the host leaf and the age of the leaf are also important factors that shape the endophytic assemblage (Arnold et al. 2000, Rashmi et al. 2019). Fungal endophytic colonization involves penetration and persistence within host plant tissues and might be influenced by the host plant (Arnold 2007). This is explained by the hypothesis of ‘balanced antagonism’ between
endophyte and their host (Schulz et al. 2015, de Silva et al. 2019a). In general, fungi (fungal virulence factors) can control activation of the host defenses and activate resistance against toxic metabolites of the host. Some fungal endophytes secrete secondary metabolites at low concentrations in the presence of elicitors or substrates of the host plants (Suryanarayanan et al. 2016). If the plant defense withstands fungal virulence, the fungus is unable to colonize plant tissues (Suryanarayanan et al. 2016). If the fungal virulence can act against plant defense, endophytes switch to pathogenic lifestyle and lead to plant diseases (Lo Presti et al. 2015, Suryanarayanan et al. 2016). This implies that the interaction of host plants and endophytic fungi can restrict fungal colonization for specific fungal groups that successfully inhabit host plant tissues without causing visible manifestations of infection.

Previous studies have indicated that incidence, diversity and composition of endophytic fungi from taxonomically related hosts vary related to the biogeographic areas (Arnold et al. 2003, Arnold 2007, U’Ren et al. 2012, Fang et al. 2019). In a general view, it is stated that diversity of endophytes is rich in seasonally wet tropics and decreases with increasing latitude, and relatively few endophyte taxa might be shared among closely related hosts in different biogeographic areas (Arnold & Lutzoni 2007, U’Ren et al. 2012). Fang et al. (2019) studied endophytic fungi from different plant tissues (roots, stems, mature healthy leaves) of Ageratina adenophora in three groups of latitudes (ranging from latitudes 22.63 to 23.22 and elevation 1171–2128 m). Their study revealed that diversity and abundance of endophytic fungi change across different geographic sites. Further, they indicated that latitude is an important factor determining the composition of the endophytic community in plant leaves. They also revealed that the latitude positively correlates with the diversity of the leaf endophytic fungi but negatively correlates with the abundance of the leaf fungi. Different patterns of endophytic assembly are related to the surrounding environment of the biogeographic areas (Fang et al. 2019). Leaves might face fluctuations against climatic changes, including soil and canopy air temperature variations (Fang et al. 2019). Kazenel et al. (2019) indicated that endophytic leaf fungal assemblage varies with altitude. Their study was further detailed that climatic variables such as pH, or soil nutrients might alter some fungal responses, which ultimately change fungal symbiont abundance and composition along with the altitudinal changes. The above explanations justify the variations in fungal communities between Magnolia liliifera from previous studies (Promputtha et al. 2005, 2007) and M. garrettii in the current study in Chiang Mai, Thailand. It also explains the difference in endophytic fungal species in different Magnolia host plants in Thailand and China (in the current study).

Previous studies reported that endophytic fungi of different plant hosts include Ascomycota, Basidiomycota, Chytridiomycota, Mucoromycota and Zygomycota (Carvalho et al. 2012, Rashmi et al. 2019). The majority of endophytic fungi belong to the division Ascomycota while major orders that fall within Ascomycota are Botryosphaeriales, Eurotiales, Hypocreales, Pleosporales, Sordariales and Xylariales (Rashmi et al. 2019). The most frequently recorded taxa of Basidiomycota belong to orders Agaricales, Cantharellales and Polyporales (Rashmi et al. 2019). When considering Ascomycota, most taxa reported being Sordariomycetes followed by Dothideomycetes (Arnold 2007, 2008) and Leotiomycetes mostly in temperate and tropical plants (Arnold 2008, Rashmi et al. 2019). However, Dothideomycetes were reported as dominant members in boreal and Arctic plants followed by Sordariomycetes and Leotiomycetes (Arnold 2007, Rungjindamai et al. 2008). It has not yet been provided robust evidence or explanations for the abundance in Ascomycota species within endophytic community. But there are few possible reasons why the endophytic community is rich in Ascomycota than in Basidiomycota. This might be due to the better adaptations of Ascomycota to colonize the internal tissues of plants than Basidiomycota (Rashmi et al. 2019). Asexual morphs of Ascomycota such as hyphomycetous and coelomycetous would help the endophytic fungi for the successful asymptomatic colonization as well as subsequent saprophytic and pathogenic infections when environmental conditions become unfavorable to the host (Rashmi et al. 2019). Ascomycota species can also produce microscopic fruit bodies or asci directly arising from the hyphae when they switch lifestyle from endophytes either saprophytes or pathogens (Rashmi et al. 2019). The above reasons might increase the
colonization of Ascomycota taxa than Basidiomycota taxa within a particular endophytic community. However, the detection of the abundance of endophytic taxa depends on isolation methods and identification techniques of endophytes (Rashmi et al. 2019). This will be the next important aspect to discuss further that affects the diversity of endophytes.

Studying endophytes is challenging because of their cryptic presence within the healthy host tissues (Rashmi et al. 2019). It is challenging to observe fungal reproductive structures within the host tissues (Rashmi et al. 2019). Fungal hyphae of endophytes can rarely be observed inside and lack any identifying characteristics (Arnold 2008). Therefore, most of the studies depend on in vitro isolations of the emerging endophytes from the sterilized tissues of the hosts on suitable growth media (Rashmi et al. 2019). However, some fungal species are unable to grow on synthetic media (Tao et al. 2008), or some may get overlooked by fast-growing fungi due to their slow growth (Zhu et al. 2008), or require specific nutrients (Jeewon & Hyde 2007, Rashmi et al. 2019). Macro- and micro-morphological characteristics such as culture characteristics, sexual and asexual morph characteristics are considered as main criteria to distinguish different endophytic taxa (Zheng et al. 2016). However, most endophytic isolates cannot sporulate in cultures and are considered as mycelia sterilia (Jeewon et al. 2017). The most reliable alternative option for this problem can be the use of DNA sequence data and molecular phylogenetic analysis (Jeewon et al. 2017). Further, concatenated sequence data were used to determine the phylogenetic relationships of fungal taxa and introduce novel taxa (Zheng et al. 2016). And also, taxonomists mostly use culture-independent molecular approaches, such as PCR-based ITS gene clone libraries, 18S rDNA, and denaturing gradient gel electrophoresis (DGGE), to reveal the complex fungal endophyte communities (Zheng et al. 2016). However, these techniques have limitations and may prevent the identification of heterogeneous species (Zheng et al. 2016). Therefore, a combination of morphological studies and molecular analyses are suggested to identify fungal taxa (Zheng et al. 2016, Jeewon et al. 2017).

Generally, it is considered that most of the endophytes that reside in internal plant tissues are able to switch their lifestyle to be either saprophytic or latent pathogens (Promputtha et al. 2005, Delaye et al. 2013, de Silva et al. 2016, 2019a, Rashmi et al. 2019). Promputtha et al. (2007) provided phylogenetic evidence of exhibiting Corynespora cassiicola as endophytes and saprobes in Magnolia liliifera in Thailand. Further, they revealed endophytic isolates of Colletotrichum, Fusarium, Guignardia, and Phomopsis have high sequence similarity with saprobic isolates of Colletotrichum, Diaporthe, Fusarium and Phylosticta from dead leaves of M. liliifera. In addition to that, two saprobic isolates of Lasiodiplodia pseudotheobromae from dead twigs and an endophytic isolate from fresh leaves of Magnolia candolii were recorded for the first time from China (de Silva et al. 2019b). Moreover, a saprobic isolate of Neopestalotiopsis saprophytica from dead leaves and two endophytic isolates from fresh leaves were identified as endophytic and saprobic lifestyle switching for the first time from Magnolia candolii in China (unpublished data). This confirms Lasiodiplodia pseudotheobromae and Neopestalotiopsis saprophytica species exhibit endophytic and saprobic lifestyles in the same host plant. This situation might be possible because endophytes switch their nutritional mode to saprobic when environmental conditions become unfavorable to the host or during host senescence (de Errasti et al. 2014, de Silva et al. 2019b). In an ecological context switching lifestyle of endophytes to saprobes might be important as saprobes can decay the plant part when the host dies (de Errasti et al. 2014, de Silva et al. 2019b).

Endophytes have attracted increasing attention for novel biological resources for exploitation in the pharmaceutical, industry and agriculture (Zheng et al. 2016). These novel compounds isolated from endophytic fungi show antimicrobial, anti-insect, antioxidant, anticancer, antineoplastic, cytotoxicity and other important biological activities (Zheng et al. 2016). Screening of novel bioactive compounds from endophytic fungi has become a research hotspot. It might be possible to explore novel fungal endophytes from a variety of host plants in different environments in the future. This requires advances in detection methods in both morphological and phylogenetic techniques to identify and introduce taxonomic novelties of unexplored endophytes.
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