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Abstract

This paper is the first in the AJOM series in which we report 100 new collections of fungi which include new species, host and country records. In all, nine new species, 90 new records and one new combination are introduced. The purpose of this series is to provide an outlet for publishing collections with sequence data, so that these observations will not be wasted and mycologists can use the information to update fungal classification and better identification of species. Previously, numerous species were described from the first collection and no further data on the species were published as it was considered low impact. This series will, therefore, increase the knowledge on the host occurrence, biogeography and sequence variability in each taxon dealt with. The distribution and hosts for each listed species are added if backed up by sequence data.

Key words – 9 new taxa – 90 new records – Ascomycota – Dothideomycetes – Leotiomyces – Molecular phylogeny – Sordariomycetes – Taxonomy

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**Introduction**
Fungal diversity is the key to many biological processes on earth. Fungi act as major decomposers and help in the recycling of nutrients in the environment (e.g. Hyde et al. 1998, Tang et al. 2005, Jeewon et al. 2003). Many others live in symbiotic associations with plants and exhibit endophytic lifestyles (Promputtha et al. 2005, 2007, Doilom et al. 2017a, Tibpromma et al. 2018a). A number of fungi also cause damage to crops and are economically important plant pathogens (Jayawardena et al. 2019a, b), but their diversity, nomenclature and roles still warrant further investigation (Hyde & Soytong 2008, Jeewon et al. 2017). Current fungal diversity estimates range from 2.2 to 3.8 million (Hawksworth & Lucking 2017) and it has been suggested that a large proportion of new species still awaits discovery and they possibly lie in tropical regions such as Thailand (Hyde et al. 2018). Our understanding of the ecological and evolutionary context of many fungi relies on their discovery from their natural environment.
Why is there a need for publications of new records based on molecular data?

Taxonomic evaluation is an ongoing process, but a common misconception among many mycologists is that new fungal records are obsolete. The goal in fungal taxonomy lies in reconstituting biological knowledge and providing all information necessary to unravel species relationships and properly describing species. In this context, publishing of new records (even from new habitats or from new hosts) plays a significant role in species documentation. Provision of such data in easily accessed journals is beneficial as this acts as a pool of taxonomic information that can be used and shared by many mycologists and further provides a platform allowing taxonomists to properly identify species and infer more accurate fungal diversity estimates. Publishing and redescriptions of extant taxa should also be encouraged as these 1) provide supplementary data and sometimes insights into better descriptions of morphological data; 2) result in additional information on phylogenetic relationships especially when data are made available from protein and rDNA sequence data; 3) assist in reevaluating, updating and validating current taxonomic classification schemes which are always in transition; 4) provide additional taxonomic documentation to help manage the increasing number of new species and new records; 5) and allow opportunities for mycologists to update nomenclature or from new hosts) plays a significant role in species documentation. Provision of additional data, we anticipate that this paper will provide a more comprehensive taxonomic classification schemes which are always in transition; 4) provide additional taxonomic documentation to help manage the increasing number of new species and new records; 5) and allow opportunities for mycologists to update nomenclature or establish new species, host shift speciation and how they adapt to different environments. Previous checklists of plant pathogens (Jayawardena et al. 2019a, b), freshwater fungi (Luo et al. 2018), terrestrial fungi (Wanasinghe et al. 2018b, Hyde et al. 2019, Phookamsak et al. 2019, Pem et al. 2019) and marine fungi (Jones et al. 2019, Dayarathne et al. 2020) were based mostly on morphological identifications supplemented with DNA based sequence data. The application of phylogenetics based on DNA sequence data in the last decade has largely improved species identification and understanding of taxonomic relationships (Jayasiri et al. 2019, Zeng et al. 2019, Zhang et al. 2019a). These approaches have often shown species to be cryptic, as in genera such as Colletotrichum (Jayawardena et al. 2016, Veloso et al. 2018) and Diaporthe (Senanayake et al. 2017), but in other cases species classification remains unresolved (e.g. Jeewon et al. 2018, Shang et al. 2020). Fungal host association has often been used to either delineate species or establish new species, but this has been reported to be erroneous (Jeewon et al. 2004). However, it is important to document species and gather data that can provide further insights into taxonomic complexity and fungal plant interactions useful for optimization of conservation practices. In this paper, we provide new data on the distribution and host records of taxa mostly in the Ascomycota with emphasis on phylogenetic inferences derived from DNA sequences of commonly used genes in molecular systematics. The paper complements Fungal Diversity notes (Hyde et al. 2019, Phookamsak et al. 2019) and Mycosphere notes (Hyde et al. 2018), where new species have been introduced and new records have been described with detailed descriptions, illustrations and updated information. With additional data, we anticipate that this paper will provide a more comprehensive taxonomic understanding of species.

Materials and methods

Materials and methods follow the previous Fungal Diversity Notes (Hyde et al. 2019, Phookamsak et al. 2019). Fresh and dried specimens in this study were collected from China, India, Italy, Oman, Russia, Taiwan, Thailand and Ukraine (further details for each taxon studied are given in the taxonomy section). Phylogenetic analyses were performed based on Bayesian inference (BI), maximum likelihood (ML) and maximum parsimony (MP) with details as outlined by Tang et al. (2007), Zhang et al. (2008) and Phukhamsakda et al. (2019). Establishment of new species and species differences are based on the guidelines outlined by Jeewon & Hyde (2016). Faces of Fungi numbers and Index Fungorum numbers were registered as described in Jayasiri et al. (2015) and
Index Fungorum (2020), respectively. The numbers of taxa in this study are organized as in the “Outline of Ascomycetes” (Wijayawardene et al. 2018).

**Taxonomy**

**Phylum Ascomycota**

**Class Dothideomycetes**

**Subclass Dothideomycetidae** P.M. Kirk et al.

For recent treatments of Dothideomycetes, we follow Liu et al. (2017b) and Wijayawardene et al. (2018).

**Capnodiales** Woron.

**Phaeothecidiellaceae** K.D. Hyde & Hongsanan

Phaeothecidiellaceae was introduced by Hongsanan et al. (2017a) to accommodate *Chaetothyrina*, *Houjia* and *Phaeothecoidiella* in Capnodiales.

![Phylogram generated from maximum likelihood analysis based on combined LSU, ITS and SSU sequence data. Twenty-five strains are included in the combined gene analyses comprising 2501 characters after alignment (850 characters for LSU, 634 characters for ITS, 1017 characters for LSU). *Capnodium citri* (CBS 131.34) and *C. coffeae* (CBS 147.52) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -12303.332942 is presented. The matrix had 775 distinct alignment patterns, with 24.61% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.244506, C = 0.238481, G = 0.282732, T = 0.234281; substitution rates AC = 1.302334, AG = 1.595992, AT = 1.294978, CG = 1.166465, CT = 5.090925, GT = 1.000000; gamma distribution shape parameter $\alpha = 0.428977$. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.9 are placed above the branches. The newly generated sequence is indicated in blue.

**Chaetothyrina** Theiss.

*Chaetothyrina* was introduced as a member of Microthyriaceae. Singtripop et al. (2016) illustrated the type specimens of *Chaetothyrina*, along with a reference specimen *C. musarum*
Hongsanan et al. (2017a) transferred it to Phaeothecoidiellaceae based on phylogenetic analyses.

**Chaetothyrina guttulata** Hongsanan & K.D. Hyde, Mycosphere 8: 142 (2016)  
Facesoffungi number: FoF01884

*Epiphytes on surface of leaves and petioles of Mangifera indica.* Sexual morph: *Thyriothecia* up to 170 µm in diameter, superficial, solitary, subdense, dark brown, circular, flattened, with an irregular central ostiole. *Upper wall* comprising cells of *textura angularis*, seated on a thin, hyaline basal stroma. *Pseudoparaphyses* filiform, septate, hyaline. *Asci* 26–45 × 11–17 µm (̅x = 35 × 14 µm, n = 30), bitunicate, 8-spored, fusoid, fusiform or clavate, with a short pedicel. *Ascospores* 11–15 × 3–5 µm (̅x = 13 × 4 µm, n = 30), 2–3-seriate, hyaline, ellipsoidal to oblong, 1-septate, constricted at the septum, with 2 guttules in each cell.  

Asexual morph: Undetermined.

**Fig. 2** – *Chaetothyrina guttulata* (MFLU 14–0006, new sequence data). a, b Host plant. c, d Ascomata. e–g Ascomata cross sections. h Ostiole. i Peridium. j Basal wall of ascoma. k Pseudoparaphyses. l–n Asci from immature to mature stage. o–q Ascospores from immature to mature stage, r–s Germinating ascospores. Scale bars: d = 100 µm, e–g = 50 µm, h = 20 µm, i–n = 10 µm, o–q = 5 µm.
Known distribution (based on molecular data) – Thailand (Hongsanan et al. 2017b, this study).
Known hosts (based on molecular data) – Mangifera indica (Hongsanan et al. 2017b, this study).
Material examined – Thailand, Chiang Rai, Tasud, Ban Doo, Pong Phra Bat, on living leaves of Mangifera indica (Anacardiaceae), 10 January 2014, X.Y. Zeng (MFLU 14-0006, new sequence data), living culture (MFLUCC 14-0539).
GenBank numbers – ITS-LSU: MN462949; SSU: MN364417; TUB: MN482717; GAPDH: MN482716.

Notes – This new collection was from the same host and locality as the type specimen, and has identical morphological features. However, there are quite a number of base-pair differences between the two collections (Fig. 1). This species is different from the generic type Chaetothyrina musarum as it lacks setae. Although the author described the characteristic of setae, the latter could not be observed in its photoplate (Hongsanan et al. 2017b). Examination of more samples is needed to understand this species.

Hysteriales Lindau
Hysteriaceae Chevall.
This family currently comprises 14 genera (Wijayawardene et al. 2018, Jayasiri et al. 2019). The latest treatment of Hysteriaceae is by Jayasiri et al. (2019) and a new record for Gloniopsis calami is reported here.

Gloniopsis De Not.
The genus comprises 68 epithets in Index Fungorum (2020), but only five species have sequence data to represent the generic placement. In addition, several species in this genus are polyphyletic (Boehm et al. 2009a, Jayasiri et al. 2019). In this study, we provide an updated phylogenetic tree based on available sequence data.

Gloniopsis calami Konta & K.D. Hyde, Fungal Diversity 80: 34 (2016) Figs 4, 5
Facesoffungi number: FoF02366
Saprobic on dead wood. Sexual morph: Ascomata 174–229 μm high x 136.5–209 μm diameter (x̄ = 196.5 x 162 μm), hysterothecial, erumpent to entirely superficial, solitary, or gregarious, oval or ellipsoid to greatly elongate, straight to curved, scattered, dark, carbonaceous, brittle, with a sunken longitudinal slit-like opening. Peridium 27–40.5 μm wide, composing thick layers of dark brown cells of carbonaceous texture, inner layers composed of several layers of hyaline to brown cells of textura angularis. Hamathecium comprising 1.5–2 μm thin, cylindrical, hyaline, branched, septate, anastomosing pseudoparaphyses. Asci 53–78 x 13–19 μm (x̄ = 64 x 16 μm, n = 10), 8-spored, bitunicate, fissitunicate, saccate, broadly clavate to cylindrical, slightly curved, short pedicellate, with knob-like pedicel, apically rounded, with a well-developed ocular chamber. Ascospores 15–20 x 7–9 μm (x̄ = 18 x 8 μm, n = 20), 2-seriate, overlapping, reddish-brown to brown, dictyosporous, fusiform, oblong or ellipsoidal to cylindrical, 4–6-trans-septate and with 2–4 vertical septa, constricted at the septa, smooth-walled. Asexual morph: Undetermined.

Culture characteristics – Ascospores germinating on WA within 12 h and germ tubes produced from spore cells. Colonies on MEA 18 mm after 7 days at 28 ºC. Mycelium superficial, low convex, slightly effuse hairy, edge entire, dark brown. Aerial mycelium, radiating outwards, superficial, septate, hyaline to brown, smooth-walled.
Known distribution (based on molecular data) – Thailand (Phang-Nga Province) – Hyde et al. (2016), Chiang Rai Province (this study).
Known substrates (based on molecular data) – On dead Calamus sp. (Arecaceae) Hyde et al. (2016), on dead wood of unidentified host (this study).
Material examined – Thailand, Chiang Rai, Huai Kang Pla, on dead wood of unidentified host, 25 October 2010, S. Boonmee, HPK02 (MFLU 10–0973, new geographical record); living culture (MFLUCC 10–0927 = BCC 52145).

GenBank numbers – ITS: MN608546, LSU: MN577415, SSU: MN577426.

**Fig. 3** – Phylogram generated from maximum likelihood analysis based on combined LSU and SSU sequence data. Nineteen taxa are included in the combined gene analyses comprising 1715 characters after alignment (821 characters for LSU, 894 characters for SSU). *Mytilinidion mytilinellum* (CBS 303.34) and *M. rhenanum* (EB 0341) are used as the outgroup taxa. The best RaxML tree with a final likelihood value of -3823.854846 is presented. Estimated base frequencies were as follows: $A = 0.251300$, $C = 0.211072$, $G = 0.294664$, $T = 0.242964$; substitution rates $AC = 0.566402$, $AG = 1.996467$, $AT = 0.358447$, $CG = 0.000000$, $CT = 6.395101$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0$. Bootstrap values for maximum likelihood equal to or greater than 75 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.

**Notes** – *Gloniopsis calami* was introduced by Hyde et al. (2016) based on morphology and phylogenetic analysis. This species was described from a palm from southern Thailand. Our strain shares similar characteristics such as ascomata, asci and ascospores with *Gloniopsis calami* (MFLUCC 15–0739). Further, phylogenetic analysis based on LSU and SSU data indicated that our strain clusters with an asexual species, *Gloniopsis leucaenae* (C289) with poor support (Fig. 3).
Comparison of ITS regions of *Gloniopsis leucaenae* (C289) and *G. calami* MFLUCC 10–0927 reveals 6 base pair (0.92%) differences. However, base pair comparison of ITS, LSU and SSU of *Gloniopsis calami* MFLUCC 15–0739 and *G. calami* MFLUCC 10–0927 are 100% similar (data not shown here). Therefore, we identify our strain as *Gloniopsis calami* and this is the first report of *G. calami* from dead wood in northern Thailand.

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**Fig. 4** – *Gloniopsis calami* (MFLU 10–0973, new geographical record). a, b Material and Appearance of ascomata on woody substrate. c Section of ascoma. d Peridium. e Pseudoparaphyses. f–i Asci. j–n Ascospores. Scale bars: c = 50 µm, d, f–i = 20 µm, e, j–n = 10 µm.

*Gloniopsis calami* Konta. & K.D. Hyde, Fungal Divers 81: 34 (2016)  
Facesoffungi number: FoF02366

*Saprobic* on decaying wood submerged in freshwater habitats. Sexual morph: *Hysterothecia* 130–200 µm high, 150–180 µm diameter, erumpent to superficial, solitary to gregarious, scattered, dark, straight to flexuous. *Peridium* 25–50 µm thick, carbonaceous, thick-walled, relatively smooth on the outer surface. *Hamathecium* comprising dense, branched, hyaline, septate pseudoparaphyses. *Asci* 107–127 × 17–21 µm (x̄ = 117 × 19 µm, n = 10), 8-spored, bitunicate, fissitunicate, cylindrical to cylindric-clavate, short pedicellate, apically rounded, with a well-developed ocular chamber. *Ascospores* 23–27 × 10–12 µm (x̄ = 25 × 11 µm, n = 20), overlapping 1–2-seriate, pale brown to dark brown, dictyosporous, fusiform, slightly curved to straight, 3–5-trans-septate and with 1–4 vertical septa, constricted at the septa, smooth-walled. Asexual morph: Undetermined.

Culture characteristics – Ascospores germinating on PDA within 14 h and germ tubes produced from spore cells. Colonies on MEA reached 20 mm after 10 days at 25 °C. *Mycelium* superficial, low convex, slightly effuse hairy, edge entire, dark brown.
**Fig. 5** – *Gloniopsis calami* (MFLUCC 10–0927, living culture). a Germinating spore. b, c Culture colonies on MEA from surface and reverse. d–f Aerial hyphae in culture. Scale bars: a, d–f = 10 µm.

Known distribution (based on molecular data) – Thailand, Phang-Nga Province (Hyde et al. 2016), Chiang Rai Province (this study).
Known hosts (based on molecular data) – On dead *Calamus* sp. (Arecaceae) (Hyde et al. 2016), on submerged dead wood of unidentified host (this study).
Material examined – Thailand, Chiang Mai Province, saprobic on decaying wood submerged in a freshwater pond, January 2014, Z.L. Luo, ZL–27 (MFLU 15–0074, new to freshwater habitats), living culture MFLUCC 14–0049.
GenBank numbers – ITS: MN 860550, LSU: MN 860555.
Notes – Our strain (MFLUCC 14–0049) shares similar characteristics such as ascomata, asci and ascospores with *Gloniopsis calami* (MFLUCC 15–0739). Further, phylogenetic analysis based on LSU and SSU data indicated that our strain clusters with *Gloniopsis calami* with high statistical support (98% ML) (Fig. 3). This is the first report of *G. calami* from submerged dead wood in freshwater habitats.

**Hysterium** Pers.

*Hysterium* is one of the largest genera of Hysteriales with a cosmopolitan distribution and comprises 435 epithets (Index Fungorum 2020). The genus is characterized by hysterothecial ascomata, carbonaceous, with a longitudinal slit, bitunicate, fissitunicate, cylindrical clavate asci and fusiform, septate, pigmented ascospores (Boehm et al. 2009a). Members of this genus are saprobic on dead wood in terrestrial habitats. An updated phylogenetic tree is provided here.

**Hysterium angustatum** Alb. & Schwein., Consp. fung. (Leipzig): 55 (1805)
Facesoffungi number: FoF 04579

*Saprobic* on dead wood. Sexual morph: *Ascomata* 207.5–294.5 µm high × 242.5–346 µm diameter (x̄ = 238.5 × 278 µm), hysterothecial, erumpent to entirely superficial, solitary, or gregarious, oval or ellipsoid to greatly elongate, straight to curved, scattered, dark, carbonaceous, brittle, with a sunken longitudinal slit-like opening. *Peridium* 42–57 µm wide, composing thick layers of dark brown cells of carbonaceous texture, inner layers composed of several layers of light
brown to brown cells of textura angularis. Hamathecium comprising 1–2 μm thin, cylindrical, hyaline, branched, septate, anastomosing pseudoparaphyses. Asci 67.5–92 × 13–20.5 μm (x̄ = 80 × 17 μm, n = 10), 8-spored, bitunicate, fissitunicate, saccate, broadly clavate to cylindrical, slightly curved, short pedicellate, with knob-like pedicel, apically rounded, with a well-developed ocular chamber. Ascospores 20–31 × 7–10 μm (x̄ = 23.5 × 8.5 μm, n = 20), overlapping 2-seriate, brown to greenish brown, fusiform, oblong or ellipsoidal to cylindrical, 3-septate, constricted at the median septum, smooth-walled. Asexual morph: Undetermined.

Culture characteristics – Ascospores germinating on WA within 12 h and germ tubes produced from both cells. Colonies on MEA 0.2 mm after 7 days at 28 °C. Cylindrical, low convex, slightly effuse hairy, edge entire, grayish-brown to brown. Aerial mycelium, radiating outwards, superficial, septate, hyaline to brown, smooth-walled.

Known distribution (based on molecular data) – Kenya, New Zealand, South Africa, United States (Boehm et al. 2009a), Italy (Jayasiri et al. 2018) and Thailand (this study).

Known hosts (based on molecular data) – On bark of living Pinus rigida (Boehm et al. 2009a), on dead aerial branch of Rubus sp. (Jayasiri et al. 2018), on dead wood of unidentified host (this study).

Material examined – Thailand, Chiang Mai, Chom Thong, on dead wood, 16 November 2010, R. Phookamsak, ITN06 (MFLU 11–0004, new geographical record), living culture (MFLUCC 11–0004 = BCC 52154).

GenBank numbers – ITS: MN608547, LSU: MN577416, SSU: MN577427.

Notes – The strain MFLUCC 11–0004 that was isolated and described in this study, is phylogenetically related to other Hysterium angustatum isolates with high statistical support (94% ML/1.00 BYPP, Fig. 7). In addition, our strain shares similar morphologies in shape, size and pigments of ascomata, asci and ascospores with Hysterium angustatum (CBS 236.34, CBS 123334, CMW 20409, GKM 243a, GKM 5211, MFLUCC 16–0623, SMH 5216). Therefore, based on morphology and phylogenetic affinity, our strain MFLUCC 11–0004 is identified as Hysterium angustatum and it is reported here as a new geographical record from Thailand.

Rhytidhysteron Speg.

Rhytidhysteron was classified in Patellariaceae (Barr 1987, Lumbsch & Huhndorf 2010) but recently accommodated in Hysteriaceae based on multi-gene phylogenetic analyses (Boehm et al. 2009a, b, de Almeida et al. 2014, Wijayawardene et al. 2014). Rhytidhysteron species are endophytes, saprobes and pathogens on plants and humans (Yacharoen et al. 2015, Chander et al. 2017). Several new species have been recently introduced based on morphology and phylogeny (Thambugala et al. 2016b, Doilom et al. 2017b, Kumar et al. 2019).

Rhytidhysteron thailandicum Thambug. & K.D. Hyde, Cryptog. Mycol. 37(1): 110 (2016)

Facesoffungi number: FoF01841

Saprobic on dead twigs of Afzelia xylocarpa. Sexual morph Ascomata 1300–1700 μm long × 380–790 μm high × 1160–1815 μm wide (x̄ = 1500 × 585 × 1505 μm, n = 10), hysterothecial with a longitudinal slit, black, solitary to aggregated, semi-immersed to superficial, with irregular opening when wet, folded at the margin when dry, with striation, subiculum, coriaceous, yellow at the center, forming an elongate slit. Exciple 50–100 μm (x̄ = 75 μm, n = 15), composed of textura angularis, comprising two cell layers, outer layer comprising black, thick-walled cells, inner layer comprising dark reddish to hyaline, thin-walled cells. Hamathecium comprising 1.–2.3 μm wide, dense, septate, filiform, cellular pseudoparaphyses, forming epithecium above the asci and enclosed in a gelatinous matrix. Asci 140–157 × 10–12 μm (x̄ = 150 × 11 μm, n = 15), 8-spored, bitunicate, subcylindrical to cylindrical, with a short pedicel, apically rounded with an ocular chamber. Ascospores 21–25.5 × 7.5–9.6 μm (x̄ = 23.5 × 8.7 μm, n = 20), uni-seriate, partially overlapping, hyaline to pale brown, becoming pale brown to dark brown, initially subglobose, 1-septate, slightly constricted at the central septum, becoming 3-septate, ellipsoidal to fusiform, slightly rounded or...
pointed at both ends, guttulate, without a mucilaginous sheath. Asexual morph Undetermined (see Thambugala et al. 2016b for the type).

**Fig. 6** – *Gloniopsis calami* (MFLU 15–0074, new to freshwater habitats). a Specimen. b–d Appearance of hysterothecia on host. e, f Section of hysterothecium. g Structure of peridium. h–j Asci. k–n Ascospores. o Germinating ascospore. p Colony on MEA. Scale bars: b = 1000 μm, c = 200 μm, d = 150 μm, e, f = 60 μm, g–j = 30 μm, k–n = 10 μm, o = 20 μm, p = 30 mm.
Fig. 7 – Phylogram generated from maximum likelihood analysis based on combined LSU and SSU sequence data. Twenty-one taxa are included in the combined gene analyses comprising 2065 characters after alignment (1043 characters for LSU and 1022 characters for SSU). *Mytilinidion mytilinellum* (CBS 303.34) and *M. rhenanum* (EB 0341) are used as the outgroup taxa. The best RaxML tree with a final likelihood value of -5388.433176 is presented. The matrix had 332 distinct alignment patterns, with 32.25% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.254589, C = 0.217165, G = 0.291200, T = 0.237046; substitution rates AC = 0.327416, AG = 3.974093, AT = 1.042057, CG = 0.582695, CT = 12.493723, GT = 1.000000; gamma distribution shape parameter α = 0.075104. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.
Fig. 8 – *Hysterium angustatum* (MFLU 11–0004, new geographical record). a Material. b Appearance of ascomata on woody substrate. c, d Sections of ascomata (Fig. c transverse section and Fig. d longitudinal section). e Peridium. f Pseudoparaphyses. g–j Asci. k–n Ascospores. Scale bars: b = 500 µm, c, d = 100 µm, e = 50 µm, f, k–n = 10 µm, g–j = 20 µm.

Culture characteristics – Ascospores germinating on PDA within 24 h and germ tubes produced from one or both ends or second or third cells. Colonies on PDA filamentous to irregular in shape, undulate edge, flat, initially white, becoming grey.
Fig. 9 – *Hysterium angustatum* (MFLUCC 11–0004, living culture). a Germinating spore. b, c Culture colonies on MEA from surface and reverse at 30 days. d–f Aerial hyphae in culture. Scale bars: a, d–f = 10 µm.

Known distribution (based on molecular data) – Thailand (Thambugala et al. 2016b, this study).

Known hosts (based on molecular data) – *Afzelia xylocarpa* (this study).

Material examined – Thailand, Chiang Rai Province, Mae Suai District, on dead twigs of *Afzelia xylocarpa* (Leguminosae), 20 November 2012, M. Doilom (MFLU 19–2701, new host record), living culture (MFLUCC 13–0051).

GenBank numbers – ITS: MN509433, LSU: MN509434, tef1: MN509435.

Notes – The holotype of *Rhytidhysteron thailandicum* has been reported from a dead twig in Thailand (Thambugala et al. 2016b). Our collection has similar morphologies to the holotype (Thambugala et al. 2016b) in its clavate to cylindrical asci, ellipsoidal to fusiform ascospores with 3 septa, but they are different with regards to the hysterothecia on host substrate along with their sizes. In combined analysis of sequence data of LSU, ITS and tef1 of our strain (MFLUCC 13–0051) clusters with *R. thailandicum* (MFLUCC 14–0503, ex-type) and (MFLUCC 12–0530) with high bootstrap and Bayesian probabilities (100% ML/ 1.00 PP) (Fig. 10). Thus, we identify our collection as *R. thailandicum*. This is the first record of *R. thailandicum* on *Afzelia xylocarpa*.

*Rhytidhysteron thailandicum* Thambug. & K.D. Hyde, Cryptog. Mycol. 37(1): 110 (2016)

Fig. 12

Facesoffungi number: FoF 01841

*Saprobic* on decaying wood. Sexual morph: *Hysterothecia* 1000–1800 µm long × 300–500 high × 500–1200 µm diameter (x̄ = 1400 × 420 × 850 µm, n = 10), arising singly or in small groups, sessile, slightly erumpent from the substrate. *Receptacle* cupulate, black, flat or slightly concave, yellowish brown when fresh, with slightly dentate margin. *Excipulum* 40–70 µm wide, ectal excipulum narrow layered, deep, thick-walled, with black cells of textura globulosa to textura angularis; medullary excipulum composed of narrow, long, thin-walled, hyaline to brown cells of textura angularis. *Hamathecium* comprising 2–5 µm wide, numerous, propoloid, apically swollen, branched paraphyses, exceeding asci in length, apices form a layer on the hymenium to develop the epithecium. *Asci* 120–150 × 12–16 µm (x̄ = 130.1 × 15.3 µm, n = 30), 8-
spored, long cylindrical, short pedicellate, rounded at the apex. Ascospores 20–32 × 7.5–10.5 µm (\(\bar{x} = 23.3 \times 8.6 \mu m, n = 40\)), uniseriate, dark brown, ellipsoid with conical ends, regularly 3-septate, smooth-walled, guttulate. Asexual morph: coelomycetous. See Thambugala et al. (2016b, Pages 15, 16) for more details.

Fig. 10 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS and tef1 sequence data. Sixteen strains are included in the combined gene analyses comprising 2358 characters after alignment (868 characters for LSU, 637 characters for ITS, 853 characters for tef1). Gloniopsis calami (MFLUCC 15-0739) is used as the outgroup taxon. The best RaxML tree with a final likelihood value of -6327.236134 is presented. The matrix had 393 distinct alignment patterns, with 15.91 % undetermined characters or gaps. Estimated base frequencies were as follows: \(A = 0.229826, C = 0.263523, G = 0.280734, T = 0.225917\); substitution rates \(AC = 1.425723, AG = 2.098406, AT = 0.841566, CG = 0.571540, CT = 7.348273, GT = 1.000000\); gamma distribution shape parameter \(\alpha = 0.502627\). Bootstrap values for maximum likelihood equal to or greater than 60 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.

Known distribution (based on molecular data) – Thailand, Chiang Rai, Phitsanulok (Yacharoen et al. 2015, Thambugala et al. 2016b), China, Yunnan Province (this study).

Known hosts (based on molecular data) – deciduous plants
Material examined – China, Yunnan Province, Qujing, 24.68703° N, 104.24653° E, 1618m, on dead twigs of an undetermined tree, 6 May, 2019, Dhanushka N. Wanasinghe (MFLU 19-2373).
GenBank numbers – ITS: MN989428, LSU: MN989429, SSU: MN989430, tef1: MN989431.
Notes – Rhytidhysteron thailandicum was introduced by Thambugala et al. (2016b), which was collected from Thailand (on a dead twig). During our investigation on diversity of microfungi in China, a specimen was recovered from a dead twig in Qujing, Yunnan Province. Morphological characters such as ascomata, asci and ascospores fit well within the sexual morph of Rhytidhysteron thailandicum. We did not obtain an isolate and therefore we isolated DNA directly from the fruiting bodies. Comparison of ITS and tef1 sequence data reveals there is no significant difference (< 3 bp differences) between our new collection and the type strain (MFLUCC 14-0503). Therefore, we introduce our new collection as a new geographical record herein.
Fig. 11 – *Rhytidhysteron thailandicum* (MFLU 19–2701, new host record). a Host substrate. b Appearance of hysterothecia on host. c, d Vertical section through hysterothecia. e Exciple. f Pseudoparaphyses. g Pseudoparaphyses and immature and mature asci. h Asci and ascospores enclosed in a gelatinous matrix. i–l Ascospores. o Germinated ascospore. Scale bars: b = 1000 µm, c, d = 300 µm, e, h = 20 µm, g, i–o = 10 µm, f = 5 µm.

**Pleosporales** Luttrell ex M.E. Barr

**Anteagloniaceae** K.D. Hyde, J.K. Liu & A. Mapook

Anteagloniaceae comprises two genera i.e., *Anteaglonium* and *Flammeascoma* (Hyde et al. 2013, Jayasiri et al. 2016, Wijayawardene et al. 2018). Its members are characterized by hysterothecial to conical or lenticular ascomata, uni- to bi-loculate, with a longitudinal slit- to pore-like opening, bitunicate, fissitunicate, cylindrical-clavate asci and 1-septate, hyaline ascospores. The latest treatment of Anteagloniaceae is by Jayasiri et al. (2019) and is followed here.

**Anteaglonium** Mugambi & Huhndorf

*Anteaglonium* is characterized by hysterothecial ascomata, carbonaceous navicular with a longitudinal slit, bitunicate, fissitunicate, cylindrical-clavate asci and 1-septate, hyaline ascospores (Mugambi & Huhndorf 2009). Members of this genus are saprobes and are commonly found on dead wood in terrestrial habitats and have a cosmopolitan distribution. The genus currently includes eight species (Index Fungorum 2020). An updated phylogenetic tree is provided here.
Fig. 12 – *Rhytidhyster*on *thailandicum* MFLU 19-2373, new geographical record (a, b Appearance of hysterothecia on host. c, d Vertical section through hysterothecium. e Cells of peridium. f–h Pseudoparaphyses. i–l Asci. m–r Ascospores. Scale bars: c, d = 200 µm, e, i–l = 20 µm, f–h, m–r = 10 µm.)
Fig. 13 – Phylogram generated from maximum likelihood analysis based on combined LSU and SSU sequence data. Twenty-one taxa are included in the combined gene analyses comprising 1769 characters after alignment (827 characters for LSU, 942 characters for SSU). *Flammeascoma bambusae* (MFLUCC 10-0551) is used as the outgroup taxon. The best RaxML tree with a final likelihood value of -3014.773692 is presented. The matrix had 139 distinct alignment patterns, with 36.86% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.247411$, $C = 0.226484$, $G = 0.303542$, $T = 0.222563$; substitution rates $AC = 0.807595$, $AG = 3.459156$, $AT = 1.159395$, $CG = 0.412993$, $CT = 12.612093$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.020000$. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.
**Fig. 14** – *Anteaglonium parvulum* (MFLU 10–0974, new geographical record). a Appearance of ascomata on woody substrate. b, c Longitudinal and vertical sections of ascomata with slightly hyphae penetrating into the host tissue. d Peridium. e Pseudoparaphyses. f, g Asci (Fig. g stained in cotton blue reagent). h Ascospores. Scale bars: a = 500 µm, b, c = 100 µm, d, f, g = 20 µm, e = 5 µm, h = 10 µm.

*Anteaglonium parvulum* (Schwein.) Mugambi & Huhndorf, Syst. Biodiv. 7(4): 460 (2009)

Facesoffungi number: FoF01931

*Saprobic* on dead wood. Sexual morph: *Ascomata* 128–192 µm high × 134–304 µm diameter (\( \bar{x} = 223 \times 166.5 \) µm), hysterothecial, superficial, semi-immersed or sunken at the base, oval to elongate, or subglobose, black, carbonaceous, straight or curved, with slightly hyphae penetrating into the host tissue. *Ostiole* central, with longitudinal slit. *Peridium* 34–58 µm thick, relatively thick, strongly carbonaceous texture, composed of thick-walled dark angular or relatively compressed pseudoparenchymatous cells, inner layers composed of hyaline to brown cells of *textura angularis*. *Hamathecium* comprising 1–1.5 µm thin, cylindrical to filiform, hyaline, branched, septate, anastomosing pseudoparaphyses. *Asci* 34–53 × 4–5.5 µm (\( \bar{x} = 44 \times 4.5 \) µm, n =
20), 8-spored, bitunicate, fissitunicate, elongate cylindric-clavate, straight or slightly curved, sessile or with short pedicel, apically rounded, with a minute ocular chamber. Ascospores 5–8 × 2.5–3 μm (x̄ = 6 × 3 μm, n = 20), 1-seriate, hyaline, ellipsoidal, 1-septate, constricted at the septa, upper cell wider and tapering towards the narrow ends, guttulate, smooth-walled. Asexual morph: Coelomycetous. Conidiomata 42–85 μm high, 42–77 μm diameter, superficial, grouped to scattered, subiculum, uniloculate or multi-loculate, subglobose to globose, dark brown. Conidioma wall 42–77 μm wide, pseudoparenchymatous, composed of dark brown cells of textura angularis to subglobose. Conidiophores 11–21 μm long, 2–2.5 μm wide, cylindrical, hyaline. Conidiogenous cells phialidic, hyaline, simple, smooth, with a conspicuous collarette at the apex. Conidia 3–4 × 1.5–2 μm (x̄ = 4 × 2 μm, n = 20), 1-celled, oblong to ellipsoidal or oval, rounded ends, slightly curved, aseptate, hyaline, smooth-walled.

Culture characteristics – Ascospores germinating on MEA with 12 h and germ tubes produced from both ends. Colonies on MEA reaching 9 mm after 7 days at 28 °C, slightly effuse, radially with undulate edge, greyish to light brown, with reddish pink pigmented in media. Mycelium superficial, branched, septate, hyaline to light brown, smooth-walled, asexual state formed within 60 days.

Known distribution (based on molecular data) – USA (Louisiana and Michigan States) (Mugambi & Huhndorf 2009), India (Hongsanan et al. 2020), Thailand (Chiang Rai Province) (Jayasiri et al. 2016, and this study).

Known substrates (based on molecular data) – On dead wood (this study).

Material examined – Thailand, Chiang Rai, Huai Kang Pla, on dead wood, 25 October 2010, S. Boonmee, HPK03 (MFLU 10–0974, new geographical record); living culture (MFLUCC 10–0928 = BCC 52146).

GenBank numbers – ITS: MN608542, LSU: MN577411, SSU: MN577422.

Notes – The new collection is a typical Anteaglonium species, and it shares similar morphological characters such as hysterothecial ascomata, black, carbonaceous, cylindric-clavate, bitunicate, fissitunicate asci and uniseriate, ellipsoidal, hyaline, 1-septate, small ascospores (less than 10 μm). In addition, this species always developed a coelomycetous asexual morph in culture (Fig. 15). Multi-gene phylogenetic analysis placed these four strains (MFLUCC 10–0928, MFLUCC 11–0374, MFLUCC 11–0380, MFLUCC 11–0511) with other Anteaglonium parvulum isolates (SMH 5210, MFLUCC 14–0815, MFLUCC 14–0817, MFLUCC 14–0821, MFLUCC 14–0823) with moderate-support and type sequence of this species is unavailable (Fig. 13). These four strains share similar sexual morphological features, but they differ in culture and asexual characteristics.

**Anteaglonium parvulum** (Schwein.) Mugambi & Huhndorf, Syst. Biodiv. 7(4): 460 (2009)

Figs 16, 17

Facesoffungi number: FoF01931

Saprobic on dead wood. Sexual morph: Ascomata 155–261 μm high × 180–284 μm diameter (x̄ = 196 × 240 μm), hysterothecial, superficial, semi-immersed or sunken at the base, oval to elongate, or subglobose, uniloculate or multi-loculate, carbonaceous, brittle, straight or curved, black. Ostiole central, with longitudinal slit. Peridium 32–48(–77.5) μm thick, relatively thick, strongly carbonaceous texture, composed of thick-walled dark angular or relatively compressed pseudoparenchymatous cells, inner layers composed of hyaline to brown cells of textura angularis. Hamathecium comprising 1–2.5 μm thin, cylindrical to filiform, hyaline, branched, septate, anastomosing pseudoparaphyses. Asci 28–41 × 4–5.5 μm (x̄ = 35.5 × 4.5 μm, n = 20), 8-spored, bitunicate, fissitunicate, elongate cylindric-clavate, straight or slightly curved, sessile or with short pedicel, apically rounded, with a minute ocular chamber. Ascospores 5–8 × 2–3 μm (x̄ = 6.5 × 2.5 μm, n = 20), 1-seriate, hyaline, ellipsoidal, 1-septate, constricted at the septa, upper cell wider and tapering towards the narrow ends, guttulate, smooth-walled. Asexual morph: Coelomycetous. Conidiomata grouped to scattered, superficial, semi-immersed at the base, subglobose to globose, dark brown. Conidioma wall pseudoparenchymatous, composed of dark brown cells of textura
angularis. Conidiophores 9–21 μm long, 1.5–3 μm wide, cylindrical, hyaline. Conidiogenous cells holoblastic. Conidia 2.5–4 × 2–3 μm (\(\bar{x} = 3 \times 2 \mu m\), n = 20), 1-celled, globose, subglobose to ellipsoidal, aseptate, hyaline, guttulate, smooth-walled.

**Fig. 15** – *Anteaglonium parvulum* (MFLUCC 10–0928, living culture). a Germinating spore. b, c Culture colonies on MEA from surface and reverse. d Aerial hyphae in culture. e Developing conidiomata in culture. f, g Section of conidiomata. h, i Peridium with developing of conidiophores and conidiogenous cells. j Conidia. Scale bars: a, d, j, i = 5 μm, f, g = 20 μm, h = 10 μm.

Culture characteristics – Ascospores germinating on MEA with 12 h and germ tubes produced from both ends. Colonies on MEA reaching 8 mm after 7 days at 28 °C, slightly effuse, radially with undulate edge, greyish, light brown, reddish pink. Mycelium superficial, branched,
septate, hyaline to light brown, with reddish pink pigmented, smooth-walled, asexual state formed within 60 days.

Known distribution (based on molecular data) – USA, Louisiana and Michigan States (Mugambi & Huhndorf 2009), India (Hongsanan et al. 2020), Thailand, Chiang Rai Province (Jayasiri et al. 2016, this study).

Known substrates (based on molecular data) – On dead wood (this study).

Material examined – Thailand, Chiang Rai, Muang, Doi Pui, on dead wood, 10 May 2011, S. Boonmee, DP05 (MFLU 11–0389, new geographical record); living culture (MFLUCC 11–0380 = BCC 52031).

GenBank numbers – ITS: MN608544, LSU: MN577413, SSU: MN577424

**Fig. 16** – *Anteaglonium parvulum* (MFLU 11–0389, new geographical record). a Appearance of ascomata on woody substrate. b, c Vertical and longitudinal sections of multi-loculate stromata. d Peridium. e Pseudoparaphyses. f–h Asci (Fig. h stained in Melzer’s reagent). i–k Ascospores. Scale bars: a = 1000 µm, b, c = 100 µm, d = 40 µm, e–h = 10 µm, i–k = 5 µm.

*Anteaglonium parvulum* (Schwein.) Mugambi & Huhndorf, Syst. Biodiv. 7(4): 460 (2009)

Facesoffungi number: FoF01931

*Saprobic* on dead wood. Sexual morph: *Ascomata* 230–259 µm high × 209–268 µm diameter (\(\bar{x} = 247.5 \times 239 \) µm), hysterothecial, superficial, semi-immersed or sunken at the base, oval to
elongate, or subglobe, carbonaceous, brittle, straight or curved, black. Ostiole central, with longitudinal slit. Peridium 52–69 μm thick, relatively thick, strongly carbonaceous texture, composed of thick-walled dark angular or relatively compressed pseudoparenchymatous cells, inner layers composed of hyaline to brown cells of textura angularis. Hamathecium comprising 1–2 μm thin, cylindrical to filiform, hyaline, branched, septate, anastomosing pseudoparaphyses. Asci 38.5–59 × 3.5–6 μm (μ = 49 × 5 μm, n = 20), 8-spored, bitunicate, fissitunicate, elongate cylindric-clavate, straight or slightly curved, sessile or with short pedicel, apically rounded, with a minute an ocular chamber. Ascospores 5–9 × 2–4 μm (μ = 6.5 × 3 μm, n = 20), 1-seriate, hyaline, ellipsoidal, 1-septate, constricted at the septa, upper cell wider and tapering towards the narrow ends, smooth-walled. Asexual morph: Coelomycetous. Conidiomata grouped to scattered, superficial, subglobe to globose, dark brown, surrounded by aerial mycelia. Conidiophores and conidiogenous cells not observed. Conidia 2.5–5 × 3–5 μm (μ = 3 × 4 μm, n = 20), 1-celled, globose to subglobe, aseptate, hyaline, guttulate, smooth-walled.

Culture characteristics – Ascospores germinating on MEA with 12 h and germ tubes produced from both ends. Colonies on MEA reaching 10 mm after 7 days at 28 °C, effuse, velvety to hairy, radially with entire edge, dark brown and sparse mycelium in the outer ring on the surface with light brown margin. Mycelium superficial and partially immersed, branched, septate, light brown, smooth-walled, asexual state formed within 60 days.

Known distribution (based on molecular data) – USA, Louisiana and Michigan States (Mugambi & Huhndorf 2009), India (Hongsanan et al. 2020), Thailand, Chiang Rai Province (Jayasiri et al. 2016), Thailand, Chiang Mai Province (this study).

Known substrates (based on molecular data) – On dead wood (this study).

Material examined – Thailand, Chiang Mai, Muang, on dead wood, 21 April 2011, S. Boonmee, DST02 (MFLU 11–0133, new geographical record); living culture (MFLUCC 11–0374 = BCC 52032).

GenBank numbers – ITS: MN608543, LSU: MN577412, SSU: MN577423.

Anteaglonium parvulum (Schwein.) Mugambi & Huhndorf, Syst. Biodiv. 7(4): 460 (2009)

Figs 20, 21

Facesoffungi number: FoF01931

Saprobic on dead wood. Sexual morph: Ascomata 155–242 μm high × 216–292 μm diameter (μ = 206 × 240 μm), hysterothecial, superficial, semi-immersed or sunken at the base, oval to elongate, or subglobe, black, carbonaceous, straight or curved. Ostiole central, with longitudinal slit. Peridium 34–83 μm thick, relatively thick, strongly carbonaceous texture, composed of thick-walled dark angular or relatively compressed pseudoparenchymatous cells, inner layers composed of hyaline to brown cells of textura angularis. Hamathecium comprising 1–2 μm thin, cylindrical to filiform, hyaline, branched, septate, anastomosing pseudoparaphyses. Asci 45–55 × 3–6 μm (μ = 50 × 4 μm, n = 20), 8-spored, bitunicate, fissitunicate, elongate cylindric-clavate, straight or slightly curved, sessile or with short pedicel, apically rounded, with a minute ocular chamber. Ascospores 6–8 × 3–4 μm (μ = 7 × 3.5 μm, n = 20), 1-seriate, hyaline, ellipsoidal, 1-septate, constricted at the septa, upper cell wider and tapering towards the narrow ends, smooth-walled. Asexual morph: Coelomycetous. Conidiomata grouped to scattered, superficial, subglobe to globose, dark brown. Conidiophores holoblastic, cylindrical, hyaline. Conidiogenous cells not observed. Conidia 2–4 × 2 μm (μ = 3 × 2 μm, n = 20), 1-celled, globose to subglobe, oblong to ellipsoidal or oval, aseptate, hyaline, smooth-walled.

Culture characteristics – Ascospores germinating on MEA with 12 h and germ tubes produced from both ends. Colonies on MEA reaching 8 mm after 7 days at 28 °C, effuse, radially with entire edge, initially white cream, eventually turning unevenly brown after 4 weeks. Mycelium superficial, branched, septate, brown, smooth-walled, asexual state formed within 60 days.

Known distribution (based on molecular data) – Thailand, USA, Louisiana and Michigan States (Mugambi & Huhndorf 2009), India (Hongsanan et al. 2020), Thailand, Chiang Rai Province (Jayasiri et al. 2016), Thailand, Chiang Mai Province (this study).
Astrosphaeriellaceae Phookamsak & K.D. Hyde

Astrosphaeriellaceae was introduced by Phookamsak et al. (2015) to accommodate Astrosphaeria and Pteridiospora. Another five genera viz. Astrosphaeriellopsis (Wanasinghe et al. 2018a), Javaria (Wijayawardene et al. 2018), Pithomyces (Wanasinghe et al. 2018a), Quercicola and Xenoastrosphaeriella (Jayasiri et al. 2019) were included in this family. The family is characterized by conical, carbonaceous ascostromata, bitunicate asci with trabeculate pseudoparaphyses, fusiform or obclavate ascospores with or without sheath and both coelomycetous and hyphomycetous asexual morphs (Phookamsak et al. 2015, Wanasinghe et al. 2018a).

Astrosphaeria Syd. & P. Syd.

Astrosphaeria introduced by Sydow & Sydow (1913) with A. fusispora as the type, is a common genus and can be found on bamboos, palms and stout grasses (Barr 1990a, Zhou et al. 2003, Tanaka & Harada 2005, Hu 2010, Zhang et al. 2012, Phookamsak et al. 2015, this study). The genus is similar to Acrocordiopsis, Caryospora, Caryosparella, Mamillisphaeria and Trematosphaeria in having carbonaceous ascostromata, and trabeculate pseudoparaphyses (Hawksworth 1981, Hyde & Fröhlich 1998, Hu 2010, Liu et al. 2011, Zhang et al. 2012), however, it can be distinguished based on asci and ascospores shape and hosts (Phookamsak et al. 2015). The genus has coelomycetous asexual morph as seen in the holomorph of Astrosphaeria bambusae (Phookamsak et al. 2015).

Astrosphaeria neostellata D.Q. Dai, Phookamsak & K.D. Hyde, Fungal Diversity 74: 172 (2015)

Facesoffungi number: FoF01226

Saprobic on dead bamboo culms. Sexual morph: Ascomata 300–700 μm diameter, 250–500 μm high, black, scattered, solitary, 2–3-gregarious, erumpent to superficial, breaking the host tissue, conical, uni-loculate, glabrous, carbonaceous, with centrally ostiole. Peridium 40–50 μm wide, of unequal thickness, poorly developed at the base, composed of dark brown to black pseudoparenchymatous cells arranged in textura angularis. Hamathecium comprising dense, 1–1.5 μm wide, branched, filamentous pseudoparaphyses, anastomosing at the top, embedded in gelatinous matrix. Asci 130–200 × 10–14 μm (x̅ = 165 × 12 μm, n = 20), 8-spored, bitunicate, fissitunicate, cylindrical to clavate, with a long furcate pedicel, apically rounded with a distinct ocular chamber. Ascospores 40–50 × 4.5–5.5 μm (x̅ = 45 × 5 μm, n = 20), overlapping 1–2-seriate, hyaline to subhyaline, fusiform, tapering at both ends, rarely aseptate, normally 1-septate, constricted at the septum, smooth-walled, guttulate, surrounded by a thin gelatinous sheath. Asexual morph: Undetermined.

Culture characteristics – Ascospores germinating on PDA within 48 h, at 23–28 ℃. Colonies growing on PDA, reaching 15 mm diameter in 2 weeks at 23–28 ℃, cottony, circular, umbonate, pale brown at the middle, dark brown at the margin from above; dark brown from below. Mycelium superficial to immersed in media, with dark brown, branched, septate, smooth hyphae.

Known distribution (based on molecular data) – Thailand, Chiang Rai (Phookamsak et al. 2015), China, Yunnan (this study).

Known hosts (based on molecular data) – Dead bamboo culms (Phookamsak et al. 2015, this study).

Material examined – China, Yunnan Province, Chuxiong Yi Autonomous Prefecture, Chuxiong City, Biji Village, on dead bamboo culms, 5 February 2019, H.B. Jiang, CX003 (KUN-
HKAS 101775, new geographic record), living culture (KUMCC 19–0218).

GenBank numbers – ITS: MN629351, LSU: MN629352, SSU: MN629353, tef1: MN635787.

![Images of Anteaglonium parvulum](image)

**Fig. 17** – *Anteaglonium parvulum* (MFLUCC 11–0380, living culture) a Germinating spore. b, c Culture colonies on MEA from surface and reverse. d, e Aerial hyphae in culture. e Compressed hyphae and developing conidiomata in culture. f Squash mount of conidioma. g, h Developing of conidiophores and conidiogenous cells. i Conidia. Scale bars: a, d, g, h = 10 µm, e = 20 µm, f = 50 µm, i = 5 µm.

Notes – The new strain forms a sister clade to *Astrosphaeriella neostellata* (MFLUCC 11–0625) with high statistical support (Fig. 22). It also shares similar morphological characteristics with the type specimen (MFLU 15–1195) of *A. neostellata* and has small differences in the size of ascomata. A comparison of the LSU, SSU and tef1 nucleotides of these two strains reveals less than 1.5% nucleotide differences, which demonstrates that our new collection is *Astrosphaeriella neostellata* (Jeewon & Hyde 2016). Based on the known distribution of *Astrosphaeriella neostellata* (Phookamsak et al. 2015), the new collection is reported in China for the first time in this study.
Based on broad morphology and phylogeny investigations, Wanasinghe et al. (2017) introduced Camarosporidiellaceae to accommodate species with conidial morphology resembling Camarosporium sensu stricto and other camarosporium-like genera. The family includes a single genus Camarosporidiella (Wanasinghe et al. 2017, Wijayawardene et al. 2018). The members of Camarosporidiellaceae have coelomycetous asexual morphs, comprising pycnidial conidiomata, with a single, papillate ostiole, enteroblastic, annellidic, integrated to discrete, doliiform, lageniform or cylindrical, hyaline conidiogenous cells and pale to dark brown conidia that are phragmosporous to muriform and mostly ellipsoidal. Sexual morphs are gregarious to solitary, globose to subglobose ascomata having a papillate, central ostiole, a peridium containing cell layers of textura angularis, cylindrical, (2–)4–8-spored asci and uniseriate, ellipsoidal, brown, muriform ascospores. Camarosporidiellaceae members are cosmopolitan in distribution and exist as saprobes, endophytes or pathogens of various host species (Wanasinghe et al. 2017, Hyde et al. 2018). An updated phylogenetic tree for the family is presented in Fig. 24 and we introduce a new host record of Camarosporidiella laburni from Colutea cilicica (Fabaceae).
**Fig. 19** – *Anteaglonium parvulum* (MFLU 11–0374, living culture). a Germinating spore. b, c Culture colonies on MEA from surface and reverse. d–f Growth of asexual state on plant tissues and developing conidiomata in water agar culture. g Aerial hyphae in culture. h, i Squash mount of conidiomata. j Close up of conidiophore and conidiogenous cell with developing of conidium at the apex. k, l Conidia. Scale bars: a, j–l = 10 µm, g = 5 µm, h = 50 µm, i = 100 µm.

*Camarosporidiella* Wanas., Wijayaw. & K.D. Hyde

*Camarosporidiella* was introduced by Wanasinghe et al. (2017) to accommodate *C. caraganicola* as the type species. *Camarosporidiella* is characterized in having camarosporium-like asexual morph and cucurbitaria-like sexual morph. Based on morphology and phylogeny support, Wanasinghe et al. (2017) placed 20 species in this genus and Hyde et al. (2018) introduced another new species, *Camarosporidiella populina* from Russia. *Camarosporidiella* species show a wide distribution as saprobes, endophytes or pathogens of various host species. In this study, we follow the latest treatment and updated account of *Camarosporidiella* in Wanasinghe et al. (2017) and Hyde et al. (2018). *Camarosporidiella laburni* is introduced as a new host record from *Colutea cilicica* (Fabaceae).
**Fig. 20** — *Anteaglonium parvulum* (MFLU 11–1147, new geographical record). a Appearance of ascomata on woody substrate. b, c Vertical and longitudinal sections of ascomata. d Peridium. e Pseudoparaphyses. f–h Asci. i–l Ascospores. Scale bars: a = 500 µm, b, c = 100 µm, d = 50 µm, e, i–l = 5 µm, f–h = 20 µm.

*Camarosporidiella celtidis* (Shear) Thambug., Wanas. & K.D. Hyde, Stud. Mycol. 87: 226 (2017)

Facesoffungi number: FoF 03533

*Saprobic* on dead twigs and thin branches. Asexual morph: *Conidiomata* pycnidial, 310–350 µm high, 330–400 µm diameter (\( \bar{x} = 323.5 \times 358.8 \) µm, n = 10), solitary or gregarious, black, immersed to semi-erumpent, unilocular. *Pycnidial wall* multi-layered, 22–26 µm, thick, comprising 4–5 layers, light-brown cells of *textura angularis*, cells towards the inside lighter. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* enteroblastic, annellidic, doliiform, integrated, solitary, hyaline, smooth-walled, and formed from the inner layer of pycnidium wall. *Conidia* 11–14 × 5–7 µm (\( \bar{x} = 12.8 \times 5.7 \) µm, n = 30), oblong, straight, rounded at both ends, sometimes narrowly rounded ends, 3-transversely septate, without longitudinal septa, smooth-walled, initially hyaline, becoming brown to dark brown at maturity. Sexual morph: See Wanasinghe et al. (2017).

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 2 weeks at 20–25 °C, colonies medium sparse, circular, flat, surface slightly rough with edge entire, margin well defined, cottony to fairly fluffy with sparse aspects, colony from above: light brown to yellowish grey at the margin, light brown to grey at the centre; reverse, yellowish at the margin, dark brown to black at the centre; mycelium light brown to whitish grey with tufting; not producing pigments in PDA.
Known distribution (based on molecular data) – USA (Shear 1902), Russia (Wanasinghe et al. 2017 and this study).

Known hosts (based on molecular data) – *Ailanthus altissima*, *Betula pendula*, *Celtis occidentalis*, *Elymus repens*, *Gleditsia tracanthis*, *Maclura pomifera*, *Morus alba*, *Prunus padus*, *Spiraea* sp. (Shear 1902, Wanasinghe et al. 2017), *Prunus armeniaca* (this study)

Material examined – Russia, Krasnodar region, Novorossiyssk City, trees near Sudzhuk lagoon (44.6836366° N 37.7952325° E), on dead twigs of *Prunus armeniaca* (Rosaceae), 14 June 2016, Timur S. Bulgakov, NK071 (MFLU 17–0804, new host record), living culture, MFLUCC 17–1765.

GenBank numbers – LSU: MN750588, SSU: MN750603, ITS: MN750609.

Notes – *Camarosporidiella celtidis* was introduced as *Cucurbitaria celtidis* by Shear (1902) from *Celtis occidentalis*. Thambugala et al. (2016a) placed this species in *Camarosporium* based on DNA sequence data from a fresh collection and introduced *Cm. uniseriatum*. Based on morphology and phylogeny evidence, Wanasinghe et al. (2017) accommodated *Cucurbitaria celtidis* in *Camarosporidiella* and the asexual morph of the species is described and illustrated herein.

**Fig. 21** – *Anteaglonium parvulum* (MFLUCC 11–0511, living culture). a Germinating spore. b, c Culture colonies on MEA from surface and reverse. d Aerial hyphae in culture. e, f Squash mount of conidiomata. g Close up of conidiophore and conidiogenous cell with developing of conidium at the apex. h, i Conidia. Scale bars: a, h, i = 5 µm, e, f = 200 µm, g = 10 µm.
Fig. 22 – Phylogram generated from maximum likelihood analysis based on a combined LSU, SSU and tef1 sequence dataset. Twenty-one strains are included in the combined gene analyses comprising 2,768 total characters including gaps (LSU: 1–855 bp, SSU: 856–1896 bp, tef1: 1897–2768 bp). The ML tree with the best scores is selected to represent the phylogenetic relationships of Astrosphaeriella species, with the final ML optimization likelihood: -7773.014991. The matrix had 482 distinct alignment patterns, with 19.13% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.244553, C = 0.251159, G = 0.280413, T = 0.223875; substitution rates AC = 1.475476, AG = 4.502665, AT = 0.996575, CG = 1.808357, CT = 17.545595, GT = 1.000000; Tree-Length = 0.380939; gamma distribution shape parameter α = 0.724752; The proportion of invariable sites I = 0.641084. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.009646. Bootstrap support for ML greater than 70% and Bayesian posterior probabilities greater than 0.95 are defined above the nodes as MLBP/BYPP. The tree is rooted to Aigialus parvus (BCC 32558). The type strains are indicated in bold and newly generated sequence is shown in blue.
As morphological characters examined largely overlap with *Camarosporidiella celtidis* (MFLU 17–0466), we, therefore, report our collection (MFLU 17–0804) as a new host record of *C. celtidis* from dying twigs of *Prunus armeniaca* (Rosaceae). The multi-gene phylogeny herein also shows our collection clusters with other *Camarosporidiella celtidis* species with high bootstrap support (88% ML, 1.00 BYPP) (Fig. 24).

**Fig. 23** – *Astrosphaeriella neostellata* (KUN-HKAS 101775, new geographic record). a, b Ascomata on host surface. c Vertical section through ascoma. d Pseudoparaphyses. e–g Asci. h Culture frontage and back. i–k Ascospores. l Peridium. Scale bars: d, e = 30 μm, f, g = 20 μm, j, k = 15 μm, i, l = 10 μm.
Fig. 24 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU, ITS, and tef1 sequence data. Eighty three strains are included in the combined gene analyses comprising 3334 characters after alignment. *Coniothyrium palmarum* (strains CBS 400.71 and CBS 758.73) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -6761.700472 is presented. The matrix had 323 distinct alignment patterns, with 12.37 % undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.242758, C = 0.240900, G = 0.267664, T = 0.248678; substitution rates AC = 1.300383, AG = 5.109489, AT = 2.415853, CG = 0.406655, CT = 8.867918, GT = 1.000000; gamma distribution
shape parameter $\alpha = 0.705373$. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.

**Fig. 25** – *Camarosporidiella celtidis* (MFLU 17–0804, new host record). a Conidiomata on host surface. b Close-up of conidiomata. c, d Vertical section through conidioma. e Conidioma wall. f Conidiogenous cells producing conidia. g–i Conidia. j Colony from above. k Colony from below. Scale bars: c, d = 100 μm, e = 20 μm, C = 20 μm; f–i = 10 μm.

*Camarosporidiella laburni* (Pers.) Wanas., Bulgakov, Camporesi & K.D. Hyde. Stud. Mycol. 87: 233 (2017)

Facesoffungi number: FoF 03540

*Saprobic* on woody branches. Sexual morph: Ascomata 350–500 μm high, 400–550 μm diameter ($\bar{x} = 425.8 \times 452.8 \mu m$, n = 10), black, superficial to semi-immersed, clustered, fully or partly erumpent, globose, multi-loculate, with an ostiole. *Peridium* 44–60 μm wide, thick, comprising 7–9 layers, outermost layer heavily pigmented, thin-walled, comprising blackish to dark brown amorphous layer, middle layer thick-walled, light brown, loosely packed cells of textura angularis, inner layer composed of 3–4 layers, hyaline, flattened, thick-walled cells of textura angularis. *Hamathecium* comprising numerous, 2–3 μm (n = 30) wide, filamentous, branched septate, pseudoparaphyses. *Asci* 155–180 × 11–14 μm ($\bar{x} = 169.4 \times 12.4 \mu m$, n = 40), 8-spored, bitunicate, fissitunicate, cylindrical, short-pedicellate, apex rounded, with a minute ocular chamber. *Ascospores* 25–30 × 8–11 μm ($\bar{x} = 27.5 \times 9.3 \mu m$, n = 30), overlapping uniseriate, initially hyaline, becoming pale brown at maturity, with slightly paler ends, muriform, mostly ellipsoidal, 5–7-transversely septate, with 4–5 longitudinal septa, deeply constricted at the middle septum, slightly constricted at remaining septa asymmetrical, conical and narrowly rounded at the ends, not surrounded by a mucilaginous sheath. Asexual morph: See Wanasinghe et al. (2017).

Culture characteristics – Colonies on PDA reaching 25 mm diameter after 2 weeks at 20–25 °C, colonies medium sparse, circular, flat, surface slightly rough with entire edge, margin well defined, cottony to fairly fluffy with sparse aspects, colony from above and reverse: greenish grey; mycelium grey to whitish grey with tufting; producing pigments (pink) in PDA.

Known distribution (based on molecular data) – Italy, Russia (Wanasinghe et al. 2017 and this study).
Known hosts (based on molecular data) – *Laburnum anagyroides* (Wanasinghe et al. 2017), *Colutea cilicica* (this study).

Material examined – Russia, Republic of Crimea, Bakhchysarai District, Sel-Bukhra Mountain, shrubs on the slopes (44.736148° N, 33.989779° E), on dying twigs of *Colutea cilicica* Boiss. & Balansa (Fabaceae), 4 July 2016, Timur S. Bulgakov, CR088 (MFLU 17–0795, new host record), living culture, MFLUCC17–1763.

GenBank numbers – LSU: MN750589, SSU: MN750604, ITS: MN750610.

**Notes** – As morphological characters examined largely overlap with *Camarosporidiella laburni* (MFLU 16–0094), we report our collection (MFLU 17–0795) as a new host record from dying twigs of *Colutea cilicica* (Fabaceae). Both species share a similar morphology, viz. black, superficial to semi-immersed, clustered ascomata, cylindrical, short-pedicellate asci and muriform, mostly ellipsoidal ascospores (Wanasinghe et al. 2017). The multi-gene (LSU, SSU, ITS and *tef1*)
phylogeny herein, also shows that our collection clusters with other Corynesporascaceae species (Fig. 24). Corynesporascaceae laburni species have been recorded from Italy and Russia (on dead aerial branches of Laburnum anagyroides) and this is the first record of Corynesporascaceae laburni from Colutea ciliaris (Fabaceae).

Corynesporascaceae Sivan.

Corynesporascaceae introduced by Sivanesan (1996) with Corynesporascaceae caryotae as the type species, was collected from a decaying leaf of Caryota urens collected in Sri Lanka and linked the sexual (Corynesporascaceae caryotae) and asexual (Corynesporascaceae) state were linked in culture. Rossman et al. (2015) recommended to use Corynesporascaceae rather than Corynesporascaceae in case of Corynesporascaceae has been widely used and includes approximately 200 species names. Corynesporascaceae is accepted in Corynesporascaceae, but this family is in invalidly published (Index Fungorum 2020). In the previous phylogenetic analysis, species of Corynesporascaceae form a distant clade together with the generic type, C. cassiicola, and is classified in Corynesporascaceae (Voglmayr & Jaklitsch 2017).

Corynesporascaceae Güssow

Corynesporascaceae was described by Güssow (1906), and more than 200 epithets have been recorded (Index Fungorum 2020). However, there are only six species in this genus with DNA sequence data (Crous et al. 2019b). Corynesporascaceae has a widespread distribution (Kirk et al. 2008a) and are saprobes, pathogens, and endophytes on woody and herbaceous plants, other fungi, nematodes, and human skin (Dixon et al. 2009, Kumar et al. 2012, Singh et al. 2012). In this study, an additional three new species are described.

Corynesporascaceae doipuiensis J.F. Li & Phookamsak, sp. nov.

Index Fungorum number: IF557020; Facesoffungi number: FoF07056

Etymology – Name reflects the location from which it was collected, Doi Pui, Chiang Rai, Thailand.

Holotype – MFLU 14–0388

Saprobic on branches of hanging dead branch of unidentified plant. Sexual morph: Undetermined. Asexual morph: Hyphomycetes, colony on natural substrate effuse, dark brown or black. Colonies effuse, thin, cottony, pale to dark grey. Mycelium partly superficial, consisting of branched, septate, smooth, thin-walled, pale hyphae. Conidiophores 212–426 × 10–15 μm (x̄ = 313.45 × 13.89 μm, n = 40), macronematous, setiferous, erect, straight or flexuous, often arising in groups, septate, unbranched, thick-walled, dark brown, sometimes light brown at the tip, percurrently proliferating from cut ends. Conidiogenous cells 27–5 × 10–16 μm (x̄ = 33.71 × 13.34 μm, n = 100) monotretic, integrated, terminal, light to moderately dark brown, often percurrently proliferating. Conidia 136–165 × 5–25.5 μm (x̄ = 143.7 × 21.31 μm, n = 100), solitary, simple, obconical, curved, subhyaline to moderately brown, acrogenous, pseudoseptate, guttulate, thick-walled, smooth, with a dark basal scar, wide at lower part, narrow and elongate at the upper part.

Culture characteristics – Conidia germinating on PDA within 12 hours and germ tubes produced from both ends. Colonies growing on PDA, cottony, white to pale grey, reaching 5 mm in 7 days at 30 °C, mycelium partly superficial, partly immersed, slightly effuse, radially striate, with irregular edge, white; Asexual and sexual spores are not formed within 60 days.

Material examined – Thailand, Chiang Rai Provence, Doi Pui, on dead herbaceous branch, 9 January 2014, J.F Li, H-15 (MFLU 14–0388, holotype), ex-type living culture at MFLUCC 14–0022.

GenBank numbers – ITS: MN648322, LSU: MN648326, SSU: MN648318.

Notes – In this study, our strain (MFLUCC 14–0022) shares a size range of the conidial characters with the type and other representative specimens (Voglmayr & Jaklitsch 2017). However, Corynesporascaceae doipuiensis is unique in having lanky, curved, hyaline to light brown conidia with long and brown conidiophores. Furthermore, phylogenetic analysis shows that our strain (MFLUCC 14–0022) forms a well-supported subclade (0.99 BYPP) (Fig. 27) with Corynesporascaceae...
Corynespora submersa (MFLUCC 16–1101) with support. Our specimen is identified as a novel species Corynespora doipuiensis collected from herbaceous aerial dead branches from Chiang Rai, Thailand.

Fig. 27 – Phylogram generated from the best scoring of the RAxML tree based on combined ITS, LSU, SSU and tef1 sequenced data of taxa in Corynespora and other related families (Periconiaceae). Cyclothyriella rubronotata (TR9) and C. rubronotata (TR) were selected as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RAxML tree with a final likelihood value of -9787.150160 is presented. The value of Tree-Length is 0.422339. The matrix had 590 distinct alignment patterns, with 26.28% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.240881, C = 0.256132, G = 0.272893, T = 0.230094; substitution rates AC = 1.532399, AG = 2.164946, AT = 1.539029, CG = 1.017301, CT = 7.622600, GT = 1.000000; gamma distribution shape parameter α = 0.169372. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with final average standard deviation of split frequencies = 0.008695. Bootstrap support for maximum likelihood (ML, black) equal to or greater than 70%. BYPP (red) equal to or greater than 0.95 are given above or below the nodes. Newly generated sequences are indicated in blue.

Corynespora submersa Z.L. Luo, H.Y. Su & K.D. Hyde, sp. nov.

Index Fungorum number: IF557058; Facesoffungi number: FoF07072

Etymology – Referring to the submerged habitat.

Holotype – HKAS 92703

Saprobic on decaying wood submerged in freshwater habitats. Sexual morph Undetermined. Asexual morph Colonies on natural substrate effuse, dark brown to black. Mycelium partly superficial, partly immersed in the substrate, composed of branched, septate, subhyaline to brown, smooth-walled hyphae. Conidiophores 150–370 μm (x̄ = 260 μm, SD = 110, n = 10) long, 10–12
μm (\(\bar{x} = 11 \mu m, SD = 1, n = 10\)) wide, macronematous, mononematous, erect or ascending, simple, straight or flexuous, pale brown to dark brown, septate, with up to four successive cylindrical proliferations. Conidiogenous cells monotretic, cylindrical, pale brown to brown, often with proliferation through the apical pore and formation of another conidium at the apex of the proliferation. Conidia 100–150 μm (\(\bar{x} = 125 \mu m, SD = 25, n=20\)) long, 16–24 μm (\(\bar{x} = 20 \mu m, SD = 4, n = 20\)) wide, formed singly or in a short chains through a wide pore at the apex of the conidiophore, almost obclavate, sometimes rostrate, straight or slightly curved, smooth, subhyaline to golden brown, 9–13-distoseptate.

**Fig. 28** – *Corynespora doipuiensis* (MFLU 14-0388, holotype). a Colonies on handing dead herbaceous branch. b–c Conidiophores. d Conidiophores bearing conidia. e–d Conidiogenous cells. g–f Conidia. n Germinated conidia. Scale bars: a = 200 μm, b–i, k, l, n = 20 μm, j, m = 10 μm.
Culture characteristics – Colonies on PDA attaining 20 mm diameter within 25 days at 25 °C under natural light, velvety, centrally raised, pale brown or greyish olivaceous, reverse dull green or grey olivaceous.

Material examined – China, Yunnan Province, saprobic on decaying wood submerged in Dulong River, May 2015, Z.L. Luo, S-504 (HKAS 92703, holotype), ex-type living culture MFLUCC 16–1101.

GenBank numbers – ITS: MN860548, LSU: MN860553.

Notes – *Corynespora submersa* resembles *C. titrarpaniensis* in having macronematous, erect, straight or flexuous, smooth conidiophores, straight or slightly curved, smooth, distoseptate conidia. However, *Corynespora submersa* differs from *C. titrarpaniensis* in having subhyaline to golden brown, smaller conidia which are formed singly or in a short chain through a wide pore at the apex of the conidiophore (Kushwaha et al. 2017). Phylogenetic analysis also shows that *Corynespora submersa* is distinct from other species presently known from culture or DNA sequence.

*Corynespora lignicola* Z.L. Luo, H.Y. Su & K.D. Hyde, sp. nov.

Index Fungorum number: IF557059; Facesoffungi number: FoF07073

Etymology – Referring to this taxon dwelling on wood.

Holotype – HKAS 92792

*Saprobic* on decaying wood submerged in freshwater habitats. Sexual morph Undetermined. Asexual morph Colonies on natural substrate effuse, dark brown to black. Mycelium partly superficial, partly immersed in the substrate, composed of branched, septate, subhyaline to brown, smooth-walled hyphae. Conidiophores (350–)470–670 (–700) μm (x̄ = 570 μm, SD = 100, n = 10) long, 9–13 μm (x̄ = 11 μm, SD = 2, n = 10) wide, macronematous, mononematous, erect or ascending, simple, straight or flexuous, septate, smooth. Conidiogenous cells monotretic, cylindrical, pale brown to brown, often with proliferation through the apical pore and formation of another conidium at the apex of the proliferation, branched. Conidia 110–156 μm (x̄ = 133 μm, SD = 23, n = 20) long, 7–9 μm (x̄ = 8 μm, SD = 1, n = 20) wide, formed singly or in a short chain through a wide pore at the apex of the conidiophore, cylindrical, straight or slightly curved, smooth, subhyaline to pale brown, distoseptate.

Culture characteristics – Colonies on MEA attaining 15 mm diameter within 20 days at 25 °C under natural light, velvety, centrally raised, pale brown or greyish olivaceous, reverse grey olivaceous to dark brown.

Material examined – China, Yunnan Province, saprobic on decaying wood submerged in Nujiang River, May 2015, S.M. Tang, S-334 (HKAS 92792, holotype), ex-type living culture MFLUCC 16–1301.

GenBank numbers – ITS: MN 860549, LSU: MN860554.

Notes – *Corynespora lignicola* resembles *C. encephalarti* in having macronematous, erect, straight, smooth-walled conidiophores, monotretic, cylindrical conidiogenous cells and distoseptate conidia which are similar in size. However, *Corynespora lignicola* differs from *C. encephalarti* in having cylindrical, straight or slightly curved, subhyaline to pale brown conidia, while *C. encephalarti* has obclavate, medium olivaceous brown to dark brown conidia (Crous et al. 2019b), and *Corynespora lignicola* is phylogenetically distinct from all species that have DNA sequences.

**Dictyosporiaceae** Boonmee & K.D. Hyde

Dictyosporiaceae was validly introduced by Boonmee et al. (2016) based on type genus, *Dictyosporium* Corda. Typical asexual morphs in Dictyosporiaceae are characterized by cheiroid conidia (Boonmee et al. 2016). Currently, 13 genera are accepted in Dictyosporiaceae (Boonmee et al. 2016, Liu et al. 2017b, Iturrieta-González et al. 2018, Yang et al. 2018).
Fig. 29 – *Corynespora submersa* (HKAS 92703, holotype). a Conidiophore with conidia on natural substrate. b Conidiophore with conidia. c Conidiophore with conidiogenous cells. d Conidiogenous cells with conidia. e–i Conidia. j, k Germinating conidia. l, m Colony on MEA. Scale bars: b, c = 100 μm, d–k = 50 μm.

*Dendryphiella* Bubák & Ranoj.

*Dendryphiella* was established by Ranojevic (1914) with the type species *D. interseminata* (Berk. & Ravenel) Bubák. Fifteen species are accepted in this genus (Crous et al. 2016a, Liu et al. 2017c, Hyde et al. 2018, Iturrieta-Gonzalez et al. 2018).
**Dendryphiella eucalyptorum** Crous & E. Rubio, Persoonia 32: 231 (2014b)

Facesoffungi number: FoF06712

*Saprobic* on decaying wood in terrestrial habitats. Sexual morph: Undetermined. Asexual morph: Hyphomycetous. *Colonies* on natural substrate superficial, effuse, greyish brown. *Mycelium* partly immersed, composed of septate, branched, hyaline hyphae, 1–2 μm wide. *Conidiophores* up to 320 μm long, macronematous, mononematous, solitary, erect, brown, thick-walled, straight or slightly flexuous, finely verruculose, septate, branched, wider at the subsection. *Conidiogenous cells* 22–41 μm long (\(\bar{x} = 31 \mu m, n = 15\)), polytretic, terminal, later becoming subterminal, proliferating asymmetrically, integrated, brown, finely verrucose, enlarged at vertex. *Conidia* 16–26 × 4–6 μm (\(\bar{x} = 21 \times 5 \mu m, n = 30\)), catenate in acropetal chain, fusiform to ellipsoidal, rounded at apex, subtruncate at base, pale brown, aseptate when young, brown or dark brown, 3-septate when mature, slightly constricted at the septa, thick-walled, verruculose.

**Culture characteristics** – Conidia germinating on water agar within 24 h. Germ tubes produced from one or both ends. Mycelia superficial, circular, with entire edge, mycelia dense at center, sparse towards circumference, yellowish white from above, yellow at center, paler towards circumference from below.

Known distribution (based on molecular data) – Spain, Asturias (Crous et al. 2014b) and China (this study).

Known hosts (based on molecular data) – *Eucalyptus globulus* (Myrtaceae) (Crous et al. 2014b) and decaying wood (this study).

Material examined – China, Guizhou Province, Guiyang, Guiyang Botanical Garden, on decaying wood, 8 August 2017, Ningguo Liu, ZWY006 (GZAAS 20−0002, new geographical record), living culture, GZCC 20−0001.

GenBank numbers – ITS: MN999925; LSU: MN999929.

Notes – *Dendryphiella eucalyptorum* was introduced by Crous et al. (2014b) on small branches of *Eucalyptus globulus* (Myrtaceae) in Spain. Our collection differs from the holotype (CBS H-21699) (Fig. 31) in having shorter conidiophores (up to 320 μm vs up to 500 μm), but there are no nucleotide differences in the ITS and LSU regions and therefore, we identify it as *D. eucalyptorum*. This is a new geographical record of *D. eucalyptorum* from China.

**Neodendryphiella** Iturrieta-González

*Neodendryphiella* was introduced by Iturrieta-González et al. (2018) based on the type species *N. tarraconensis* Iturrieta-González, Gené & Dania García. *Neodendryphiella* species have been reported as coprophilous, saprobic and soil fungi. Only three species are accommodated to this genus (Iturrieta-González et al. 2018).

**Neodendryphiella tarraconensis** Iturrieta-González, Gené & Dania García, MycoKeys 37: 30 (2018)

Facesoffungi number: FoF06713

*Saprobic* on decaying wood in terrestrial habitats. Sexual morph: Undetermined. Asexual morph: Hyphomycetous. *Colonies* on natural substrate superficial, effuse, golden brown, velvety. *Mycelium* mostly immersed, composed of septate, branched, median brown hyphae, 1–2 μm wide. *Conidiophores* up to 530 μm long, 3–5 μm wide at tip, 8–10 μm at base, macronematous, mononematous, erect or slightly flexuous, branched or unbranched, cylindrical, dark brown at base, slightly paler towards the apex, smooth to finely verrucose, thick-walled. *Conidiogenous cells* polyblastic, terminal and intercalary, mostly cylindrical. *Ramoconidia* 0–1-septate, pale brown, smooth to verruculose, mostly cylindrical. *Conidia* 9–15.5 × 3–5 μm (\(\bar{x} = 11 \times 4 \mu m, n = 30\)), catenate, ellipsoidal or subcylindrical with more or less rounded ends, 0–2-septate, slightly constricted at the septum; when 2-septate, usually constricted at only one septum, pale brown, verruculose.

Culture characteristics – Conidia germinating on water agar within 24 h. Germ tubes produced from one or both ends. Mycelia superficial, circular, slightly umbonate at center with
entire edge, olivaceous brown from above, dark brown at center, paler towards circumference from below.

Known distribution (based on molecular data) – Spain (Tarragona) (Iturrieta-González et al. 2018) and China (this study).

Known hosts (based on molecular data) – From garden soil (Iturrieta-González et al. 2018) and decaying wood (this study).

Material examined – China, Guizhou Province, Zunyi, Wangcao (28°12.30’N, 107°10.24’E), on decaying wood, 15 September 2018, N.G. Liu, KKS016 (GZAAS 20–0003, new geographical record), living culture, GZCC 20–0002.

GenBank numbers – ITS: MN999922, LSU: MN999927.

Notes – Neodendryphiella tarracunensis was introduced by Iturrieta-González et al. (2018) from soil in Spain. This species was previously identified as Dendryphiella sp. Our collection differs from the holotype (CBS H-23479) in having much longer conidiophores (up to 530 μm vs 19–185 μm). This may be because of the differences when grown in natural substrate (our collection) and in culture (holotype). ITS comparison between our strain and FMR 16234 showed that there are only 3 bp differences in a total of 452 bp. Thus, we identify it as Neodendryphiella tarracunensis and it is a new geographical record for China.
Fig. 31 – Phylogram generated from maximum likelihood analysis based on combined LSU and ITS sequence data. Twenty-eight strains are included in the combined gene analyses comprising 1404 characters after alignment (825 characters for LSU, 579 characters for ITS). *Periconia igniaria* (CBS 379.86) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -7037.019449 is presented. The matrix had 473 distinct alignment patterns, with 12.81% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.238381, C = 0.235435, G = 0.277387, T = 0.248797; substitution rates AC = 1.542553, AG = 2.179153, AT = 2.337602, CG = 0.362500, CT = 6.291882, GT = 1.000000; gamma distribution shape parameter α = 0.834067. Bootstrap values for maximum likelihood equal or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in bold and blue.

**Didymellaceae** Gruyter, Aveskamp & Verkley

The most recent taxonomic treatments of Didymeallaceae are from Valenzuela-López et al. (2018), Wanasinghe et al. (2018a), and Jayasiri et al. (2019). This family was established by de Gruyter et al. (2009) with species which are traditionally classified as *Phoma* and phoma-like.
Fig. 32 – *Dendryphiella eucalyptorum* (GZAAS 20–0002, new geographical record). a, b Colonies on natural substrate. c Conidiophore. d–f Conidiogenous cells and conidia. g–n Conidia. p Germinated conidium. Scale bars: c = 20 μm, d–p = 10 μm.

**Ascochyta** Lib.

*Ascochyta* with *Ascochyta pisi* Lib. as its type species, contains numerous endophylic, pathogenic and saprobic species associated with a wide range of hosts worldwide (Wijayawardene et al. 2017).

**Ascochyta herbicola** Qian Chen & L. Cai, Stud. Mycol. 82: 187 (2015)  
Facesoffungi number: FoF07122

*Saprobic* on dead aerial stem of *Orobanche* sp. Sexual morph: Undetermined. Asexual morph: Coelomycetous. *Conidiomata* visible as dark round dots on the host surface. *Conidiomata* 120–340 μm diameter, pycnidial, solitary, scattered or gregarious, globose to irregular, semi- immersed to immersed, sometimes erumpent, unilocular, black. *Conidioma wall* thick, 2–4 layered, hyaline to dark brown cells of *textura angularis*. *Conidiogenous cells* hyaline, phialidic, globose. *Conidia* 5–7
× 2–3 μm wide, cylindrical to subcylindrical, guttulate, aseptate.

**Fig. 33** – *Neodendryphiella tarraconensis* (GZAAS 20–0003, new geographical record). a–c Colonies on natural substrate. d, e Conidiophores and conidia. f, g Conidiogenous cells and conidia. h–k Conidia. l Germinated conidium. Scale bars: d, e = 50 μm, f, g = 10 μm, h–l = 5 μm.
Fig. 34 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, RPB2 and TUB sequence data. Thirty-three strains are included in the combined gene analyses and *Phoma herbarum* (CBS 502.91 and 377.92) strains are used as the out-group taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of \(-5091.923675\) is presented. The matrix had 214 distinct alignment patterns, with 22.94\% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.234220, C = 0.256120, G = 0.244025, T = 0.265634; substitution rates AC = 0.657233, AG = 4.062723, AT = 1.701821, CG = 0.583489, CT = 9.457386, GT = 1.000000; gamma distribution shape parameter \(\alpha = 0.669698\). Bootstrap values for maximum likelihood equal to or greater than 60 and Bayesian posterior probabilities equal or greater than 0.90 are placed above or below the branches. The newly generated sequences are indicated in blue.
**Fig. 35** – *Ascochyta herbicola* (JZB 380047, new host record). a Appearance of fruiting bodies on host substrate. b Section through the fruiting body. c Peridium cell wall. d Immature and mature conidia attached to conidiogenous cells. e Conidia. Scale bars: a = 200 μm, e = 10 μm.

Culture characters – Colonies on OA 4–5cm diameter after 7 days, regular, colorless to dark concentric zones, aerial mycelium, fluffy, white, reverse colorless.

Known distribution (based on molecular data) – Italy, Forlì-Cesena Province, United States, Wyoming (Chen et al. 2015, this study).

Known hosts (based on molecular data) – *Orobanche* sp., Water, stems of *Syntheris dissecta*

Material examined – Italy, Province of Forlì-Cesena [FC], Forlì – Via Pietro Nenni, on dead aerial stem of *Orobanche* sp., (Orobanchaceae) 9 March 2017, E. Camporesii, IT 3850 (MFLU 380047), living culture, JZB380047.

GenBank number – ITS: MN989422.

Notes – The collection obtained from dead aerial stem of *Orobanche* sp. was identified as *Ascochyta herbicola* with support from morphology and phylogeny. Our isolate clustered with the reference strain of the *Ascochyta herbicola* (CBS 629.97 7) in the combined LSU, ITS, RPB2 and TUB sequence phylogeny (Fig. 34). We could not obtain RPB2 and TUB sequence data from our isolate, and were unable to perform further analysis. This is the first record of *Ascochyta herbicola* from *Orobanche* sp. in Italy.
**Calophoma** Qian Chen & L. Cai

Chen et al. (2015) introduced *Calophoma clematidina* as the type species and there are ten epithets recorded under *Calophoma* (Chen et al. 2015). All species have been reported with sexual and asexual morphs. *Calophoma clematidina* was reported from the stem of *Clematis* sp. in the Netherlands (Woudenberg et al. 2009). *Calophoma* is characterised by solitary, pycnidial conidiomata which have conspicuously papillate ostioles. The pycnidial wall comprises several layers of oblong to isodiametric cells. Conidiogenous cells are phialidic, hyaline, smooth, and ampulliform to doliiform. Conidia are cylindrical to ellipsoidal, smooth and thin, and aseptate or 1-septate. Chlamydospores are tan to dark brown. In this study, we introduce a novel species to this genus with support from combined LSU and ITS sequence data.

**Fig. 36** – Phylogram generated from maximum likelihood analysis based on combined LSU and ITS sequence data. Twenty-two strains are included in the combined gene analyses comprising 1,358 characters after alignment (888 characters for LSU, 470 characters for ITS). *Didymella macropodii* CBS100191 is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -2481.811840 is presented. The matrix had 66 distinct alignment patterns, with 0.63% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.246751, C = 0.220789, G = 0.275533, T = 0.256927; substitution rates AC = 6.461538, AG = 18.966230, AT = 9.278572, CG = 2.883910, CT = 46.053729, GT = 1.000000; gamma distribution shape parameter α = 0.020000. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.

**Calophoma humulicola** Chaiwan, T.C Bulgakov, Jayaward & K.D. Hyde, sp. nov.

- Index Fungorum number: IF557012; Facesoffungi number: 06956
- Etymology – The specific epithet “humulicola” refers to the meaning “borne on *humulus*”
*lupulus*”

Holotype – MFLU 17–2130  
*Saprobic* on dead wood of *humulus lupulus*. Sexual morph: Undetermined. Asexual morph: *Conidiomata* pycnidal, solitary, globose to subglobose, mostly with some hyphal outgrows, produced on the agar surface or immersed, (120–)135–165 × 85–130 μm. *Ostioles* 1(–3), conspicuously papillate. *Pycnidial wall* pseudoparenchymatous, 2–4-layered, 13–21 μm thick, composed of oblong to isodiametric cells. *Conidiogenous cells* phialidic, hyaline, smooth, ampulliform to doliiform, 5.5–7.5 × 4–7 μm. *Conidia* ellipsoidal to cylindrical, smooth- and thin-walled, aseptate or occasionally 1-septate, 4.5–7 × 2–3 μm, with (0–)2–4(–8) polar guttules. Conidial matrix pale pink. *Chlamydospores* usually scanty, uni- or multi-cellular, 1-celled, intercalary, guttulate, thick-walled, green brown, 8–10 μm diameter, multi-cellular irregular dictyo/phragmosporous, somewhat botryoid and in combination with 1-celled chlamydospores, tan to dark brown, 3–50 × 12–25 μm.

Culture characteristics – Conidia germinating on PDA within 24 hours. Germ tubes produced around conidia. Colonies on PDA entire edge, velvety, medium dense, flat or effuse, brownish grey. *Mycelium* 1.5–3.8 μm broad, partly superficial, partly immersed, white, septate.

Material examined – Russia, on dead wood of *Humulus lupulus* (Cannabaceae), 11 May, T.S. Bulgakov, T-1828 (MFLU 17–2130, holotype).  
GenBank numbers – ITS: MN796328, LSU: MN796329.

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*Fig. 37* – *Calophoma humulicola* )MFLU 17–2130, holotype(, a–c Conidia observed on host substrate. d Conidiomata. e pycnidial wall. f Conidia. g Conidiogenous cell. Scale bars: a = 500 μm, b = 200 μm, c = 100 μm, d, e = 50 μm, f = 20 μm, g = 10 μm.
Fig. 38 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, and RPB2 and TUB sequence data. Twenty-three strains are included in the combined gene analyses comprising 2331 characters after alignment. *Phoma herbarum* (CBS 377.92, 502.91 and 615.75) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -10101.832594 is presented. The matrix had 465 distinct alignment patterns, with 6.60% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.237653, C = 0.243234, G = 0.276939, T = 0.242174; substitution rates AC = 1.371854, AG = 5.338217, AT = 1.639327, CG = 0.874686, CT = 12.527823, GT = 1.000000; gamma distribution shape parameter \( \alpha = 0.656489 \). Bootstrap values for maximum likelihood and parsimony equal to or greater than 50 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.
Notes – *Calophoma humulicola* is morphologically similar to *Calophoma humuli* (Hyde et al. 2019), but phylogenetically distinct and constitutes an independent lineage basal to an unidentified species of *Calophoma* (CBS 186.55). The new taxon groups with *C. petasitis* with 99% ML statistical support (Fig. 36), which has been reported from a dead stem of *Petasites* sp. (Asteraceae), near Campigna-Santa Sofia, Province of Forlì-Cesena, Italy, on 9 June 2014 (Tibpromma et al. 2017).

A comparison of LSU and ITS with these two strains reveals 3/960 (0.31%) and 5/522 (0.95%) base pair differences following the guidelines of Jeewon and Hyde (2016). Therefore, we introduce our collection as a new species in *Calophoma*.

**Epicoccum** Link

*Epicoccum* was introduced by Chen et al. (2015). Species belonging to this genus are characterized with epicoccoid, subcylindrical conidia, and irregular pycnidial conidiomata. The most recent taxonomic treatments to this family are Jayasiri et al. (2017), Valenzuela-Lopez et al. (2018), Wanasinghe et al. (2018a) and Chethana et al. (2019).

**Epicoccum latusicollum** Qian Chen, Crous & L. Cai, Stud. Mycol. 87: 144 (2017)  
Facesoffungi number: FoF06580

*Saprotrophic or pathogenic on Phragmites australis* leaves. Sexual morph: Undetermined. Asexual morph: Growing well on PDA. *Conidiomata* pycnidial, mostly solitary, sometimes aggregated, globose to subglobose or pyriform, glabrous, produced on the agar surface, ostiolate, sometimes elongate with a short, slightly papillate neck. *Pycnidial wall* pseudoparenchymatous, composed of oblong to isodiametric cells, with 2–3 cell layers of which outer layers are brown. *Conidiogenous cells* phialidic, hyaline, smooth, ampulliform to doliiform. *Conidia* phialidic, hyaline, smooth, walled, 4.5–6 × 2–3 μm (μ = 5 × 2.5 μm, n = 40). *Chlamydospores* aseptate, light brown, smooth and thin-walled, in chains or single, ovoid.

Culture characteristics – Colonies on PDA reach 80–85 mm diameter after 7 d, margin regular, floccose aerial mycelia covering the whole colony, dense, white to grey, forming several white mycelial pellets; reverse white.

Known distribution (based on molecular data) – China and Japan (Chen et al. 2017, Valenzuela-Lopez et al. 2018), China (this study).

Known hosts (based on molecular data) – *Acer palmatum*, *Camellia sinensis*, *Podocarpus macrophyllus*, *Sorghum bicolor*, *Vitex negundo* (Chen et al. 2017, Valenzuela-Lopez et al. 2018), *Phragmites australis* (this study).

Material examined – China, Shandong, Yellow River Park, on leaf spots of *Phragmites australis* (Poaceae), 7 October 2017, Y.Y. Hao (JZBH380037, new host record) living culture, JZB380037.

GenBank submissions – ITS: MN533797, LSU MN533800.

Notes – *Epicoccum latusicollum* was introduced by Chen et al (2017). In our phylogenetic analysis, this isolate clusters with the type (CGMCC 3.18346) with 100% ML bootstrap support and 1.0 BYPP (Fig. 38). The collection shares similar colony morphology, conidial structure and sizes with the protologue (Chen et al 2017). Pairwise comparison of nucleotide sequences in four gene regions reveals the following: LSU is 100% similar; ITS less than 1% dissimilar; protein-coding regions (BT and RPB2) less than 5% dissimilar. Based on these facts isolate (JZBH380037) is considered to be *Epicoccum latusicollum*. This species has been reported on *Acer palmatum*, *Camellia sinensis*, *Podocarpus macrophyllus*, *Sorghum bicolor* and *Vitex negundo* (Chen et al. 2017). In the present study, we collected this taxon associated with leaf spots on *Phragmites australis* for the first time (Farr & Rossman 2020).

**Nothophoma** Qian Chen & L. Cai

*Nothophoma* was introduced by Chen et al. (2015) to accommodate five species. This genus is comprised of important plant pathogens. Currently, there are ten species accepted in this
Nothophoma pruni

Chethana, J.Y. Yan, X.H. Li & K.D. Hyde, Mycosphere 10(1): 520 (2019)

Facesoffungi number: FoF 04917

Saprobic on diseased leaves of Castanea mollissima. Sexual morph: Undetermined. Asexual morph: Pycnidia on the PDA surface, 0.21–0.45 mm (x̄ = 0.29 mm, n = 10) diameter, solitary, scattered, globose to irregularly shaped, black, ostiolate. Conidiogenous cells phialidic, hyaline, simple, doliform to ampulliform, variable in size. Conidia 4.7–8.6 × 2.5–3.5 μm (x̄ = 6 × 3.2 μm, n = 40), cylindrical to obvoid or oblong, hyaline, aseptate, smooth-walled. Conidial exudates hyaline to buff.

Known distribution (based on molecular data) – China, Beijing (Chethana et al. 2019 and this study).

Known hosts (based on molecular data) – Prunus avium (Chethana et al. 2019), Castanea mollissima (this study).

Culture characteristics – Colonies on PDA reach 80 mm diameter after 7 days at 25 °C, with regular margin, dull white aerial mycelium surface floccose, with reverse pale vinaceous.

Material examined – China, Shandong, Yellow River Park, on leaf spots of Castanea mollissima (Fagaceae), 7 October 2017, Y.Y. Hao (JZBH380038, new host record), living culture, JZB380038.

GenBank numbers – ITS: MN533798, LSU: MN533801, TUB: MN991303, RPB2: MN991306.

Notes – Nothophoma pruni was introduced by Chethana et al. (2019) as a leaf spot causing microfungus associated with Prunus avium. The taxon collected in this study fits well within Didymellaceae with its morphological characteristics. In our combined phylogenetic analysis of the LSU, ITS, TUB and RPB2 gene regions, our isolate clusters with the N. pruni type strain (MFLUCC 18–1601) with 100% ML bootstrap and 1.0 BYPP. Conidial dimensions and shape are similar to the original description. Nucleotide sequences were compared in MFLUCC 18–1601 and JZBH380038. LSU and RPB2 sequences are 100% similar and ITS has less than 1% dissimilarity. Despite some nucleotide differences in the TUB gene region we identify JZBH380038 as Nothophoma pruni. This is the first report of N. pruni from China, associated with Castanea mollissima worldwide (Farr & Rossman 2020).
Quercus sp. and Ziziphus jujuba (Jianyu et al. 2016, Moral et al. 2017, Chethana et al. 2019, Farr & Rossman 2020).

Fig. 39 — Epicoccum latusicollum JZB380037, new host record. a, b Material examined. c Pycnidial wall. d Conidia on PDA. e Developing chlamydospores. f Chlamydospores g Culture on PDA (upper view). h culture on PDA (reverse view). Scale bars: d–f = 20 µm.

Material examined – China, Shandong, Yellow River Park, on leaf spots of Rosa multiflora (Rosaceae), 7 October 2017, Y.Y. Hao (JZBH380039, new host record, living culture, JZB380039.

GenBank numbers – ITS: MN533799, LSU: MN533802, TUB: MN991304, RPB2: MN991307.

Notes – Our Nothophoma quercina isolate investigated in this study clusters with other Nothophoma quercina strains (UTHSC:D116-270 and CBS 633.92; type) in the multiple gene phylogenies with 63% ML bootstrap value. The morphological features of our collection are also identical to those published (Chen et al. 2013). Pairwise comparison of nucleotide sequences in the four gene regions, LSU, ITS and RPB2 of both the type and our taxon also reveal a high percentage of similarity that confirms strain JZBH380039 as Nothophoma quercina. Nothophoma quercina is a pathogenic species in Didymellaceae (Chen et al. 2013). Previously this species was known as Ampelomyces quercinus and Phoma fungicola until it was synonymised with this current name by
Chen et al. (2013). This is the first report of *N. quercina* from *Rosa multiflora* (Farr & Rossman 2020).

**Fig. 40** – *Notophoma pruni* JZB380038, new host record. a Material examined. b–c Pycnidia. d Pycnidial wall. e–g Conidia on PDA. h Culture on PDA (upper view). i Culture on PDA (reverse view). Scale bars: b = 100 μm, c = 100 μm, d–f = 20 μm, g = 10 μm.

**Phomatodes** Qian Chen & L. Cai, Stud. Mycol. 82: 191 (2015)

*Phomatodes* was introduced by Chen et al (2015) to accommodate phoma-like taxa. The type species of this genus is *Phomatodes aubrietiae* which is characterized by pycnidial, globose to subglobose, ostiolate conidiomata, solitary or confluent, with a 3–5-layered, pigmented pseudoparenchymatous pycnidial wall, phialidic, hyaline, smooth, ampulliform to doliiform conidiogenous cells and cylindrical to allantoid, hyaline, thin-walled, smooth, aseptate, guttulate conidia (Chen et al 2015). In this study, we introduce a new host record of *P. nebulosa*.

**Phomatodes nebulosa** (Pers.) Qian Chen & L. Cai, Stud. Mycol. 82: 191 (2015)  
Facesoffungi number: FoF06803

*Saprobic or necrotrophic on Urtica* spp. Sexual morph: Undetermined. Asexual morph *Conidiomata* pycnidial, solitary, globose to subglobose, glabrous, semi-immersed or immersed. *Ostiole* single, slightly papillate. *Pycnidial walls* pseudoparenchymatous, 4–6-layered, thick, composed of isodiametric cells. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* phialidic, hyaline, smooth, ampulliform to doliiform. *Conidia* 5–7 × 1.5–2.5 μm (x̄ = 2 × 5.5 μm, n = 30), ellipsoidal to cylindrical, smooth- and thin-walled, aseptate.

Culture characteristics – Colonies on PDA, 35–40 mm diameter after 7 days at 25 °C, margin regular, floccose, initially yellowish white and becoming smoke-grey, with immersed thin mycelia mat, reverse dark olivaceous with a yellow entire margin.

Known distribution (based on molecular data) – Netherlands, Poland, Uzbekistan (Farr & Rossman 2020), Italy (this study)

Known hosts (based on molecular data) – *Armoracia rusticana, Datisca cannabina, Mercurialis perennis, Thlaspi arvense* (Farr & Rossman 2020), *Urtica* sp. (this study).

Material examined – Italy, Province of Forli-Cesena, Fiumana di Predappio, on dead aerial branch of *Urtica* spp. (Urticaceae), 15 December 2015, E. Camporesi, IT 3808 (JZBH380041, new host record, living culture, JZB380041.

GenBank numbers – ITS: MN648212, LSU: MN640407.
Notes – In the present study, *Phomatodes* species was collected from dead and dying twigs and branches of *Urtica dioica* in Forlì-Cesena, Italy. The taxon clustered together with the *Phomatodes nebulosa* type (CBS 100191) with 100% ML bootstrap and 1.0 BYPP support (Fig 38). Spores and colony characters are similar to the type species (Chen et al. 2015). Therefore, we confirm the taxon collected in the present study as *Phomatodes nebulosa*. Currently, there are only two species associated with this genus, viz. *P. aubrietiae* and *P. nebulosa*. This is the first report of *Phomatodes* associated with *Urtica* sp. (Farr & Rossman 2020).

**Fig. 41** – *Nothophoma quercina* JZB380039, new host record. a Material examined. b Pycnidia. c–d Conidiogenous cell. e–g Conidia on PDA. h Mycelia. i Culture on PDA (upper view). j Culture on PDA (reverse view). Scale bars: b–d = 20 µm, e–h = 10 µm.
Didymosphaeriaceae Munk = Montagnulaceae M.E. Barr, Mycotaxon 77: 194 (2001)

We follow the latest treatments and updated accounts of Didymosphaeriaceae presented in Ariyawansa et al. (2014a), Wanasinghe et al. (2018b) and Tibpromma et al. (2018b). This family comprises 26 accepted genera (Wijayawardene et al. 2018).

Fig. 42 – Phomatodes nebulosa (JZBH380041, new host record). a–c Appearance of conidiomata on host surface. d Pycnidial wall. e Conidia on host. f Conidioma on PDA. g Conidia on PDA. h Upper view of colony on PDA. i reverse view of colony on PDA. Scale bars: a = 500 μm, c–f = 20 μm.

Neokalmusia Ariyaw. & K.D. Hyde

Neokalmusia currently comprises five species and is typified with *N. brevispora*. Previously *N. brevispora* was categorized under *Phaeosphaeria* as *Phaeosphaeria brevispora* by Hedjaroude (1968). Studies by Ariyawansa et al. (2014c) revealed that placement of *Phaeosphaeria brevispora* and *Kalmusia scabrispora* were not acceptable and hence *Neokalmusia* was erected to accommodate those two species.

Neokalmusia didymospora D.Q. Dai & K.D. Hyde, Sydowia 68: 20 (2016)  
Facesoffungi number: FoF00061

*Saprobic* on dead culms of *Microstegium* sp. Sexual morph: *Ascomata* 420–460 μm high, 140–220 μm diameter, immersed and raising host tissue, becoming erumpent, solitary, scattered, globose, brown to dark brown, ostiolate, wide, brown to reddish brown, smooth. *Peridium* 20–35 μm, wider at the apex and thinner at the base, composed of 4–6-layers of dark brown, cells of *textura angularis*, cells towards the inside lighter, at the outside, darker, fused with the host tissues.
*Hamathecium* comprising numerous, 1.5–1.8 µm wide, filamentous, branched, septate, pseudoparaphyses. *Asc* 135–140 × 8–9.5 µm, 8-spored, bitunicate, fissitunicate, elongate-clavate to short cylindrical, long pedicellate, thick-walled at the apex, with a minute ocular chamber. *Ascospores* 12–13.5 × 5–5.5 µm, overlapping 1–2-seriate, hyaline to light brown when immature, becoming brown to blackish brown when mature, ellipsoidal, unequally 1-septate and strongly constricted at the septum, with the upper cell wider and pointed and lower cell longer and rounded, smooth-walled, thick-walled, without a mucilaginous sheath.

![Phylogenetic tree](image)

**Fig. 43** – Phylogenetic tree generated from maximum likelihood (ML) based on LSU and SSU sequences. Bootstrap support (BS) values above 50% and Bayesian posterior probabilities (BYPP) equal or greater than 0.90 are placed above the branches are shown at nodes. The tree is rooted with *Bimuria novae-zelandiae* (CBS 107.79). The best RaxML tree with a final likelihood value of -3717.445755 is presented. The matrix had 80 distinct alignment patterns, with 12.08% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.255042, C = 0.219029, G = 0.282988, T = 0.242942; substitution rates AC = 2.742405, AG = 5.807474, AT = 3.101080, CG = 0.711167, CT = 15.431181, GT = 1.000000; gamma distribution shape parameter α = 97.241967.

Known distribution (based on molecular data) – Thailand, Chiang Rai (Dai et al. 2015), China, Yunnan, Kunming (this study).

Known hosts (based on molecular data) – Bamboo (Dai et al. 2015), *Microstegium* sp. (Poaceae) (this study)

Material examined – China, Kunming, KIB premises, on decaying stems of *Microstegium* sp. (Poaceae), 20 October 2016, A. Karunarathna, AKKIB 48 (MFLU 17–0371, KUN 97364, new host and geographic record); living culture, MFLUCC 17–381; KUMCC 16–0233.

GenBank number – LSU: MN989184.

Notes – Our strain is phylogenetically related to *N. didymospora* (Fig. 43) and both are morphologically similar. *Neokalmusia didymospora* is a new host and geographic record.

**Paracamarosporium** Wijayaw. & K.D. Hyde

*Paracamarosporium* was introduced by Wijayawardene et al. (2014) to accommodate *Camarosporium psoraleae* Crous & M.J. Wingf. Both *Camarosporium* and *Coniothyrium*-like species are morphologically similar and group in this genus, hence taxa in this genus are difficult to be distinguished. Currently, there are seven species epithets under *Paracamarosporium* (Index Fungorum 2020).
Paracamarosporium fagi Crous & R.K. Schumach., Sydowia 67: 109 (2015)

Facesoffungi number: FoF 06223

Saprobic on twigs of Ziziphus jujuba. Sexual morph: Undetermined. Asexual morph: Conidiomata 336–449 × 84–277 µm (± = 381 × 238.2 µm, n = 10) solitary, immersed to erumpent, globose with central ostiole. Peridium comprising 2–3 layers of brown textura angularis. Conidiophores reduced to conidiogenous cells or with a supporting cell lining the inner cavity. Conidiogenous cells 1.8–3.1 × 4.7–6.9 µm (± = 2.22 × 5.8 µm, n = 10), annellidic, hyaline, smooth, ampulliform, apex with prominent periclinal thickening, or with percurrent proliferation. Conidia 8–10 × 3–4 µm (± = 10.1 × 4.3 µm, n = 10) solitary, subcylindrical with obtuse ends, initially hyaline, smooth, aseptate, becoming brown, medially 1-septate and surface becoming warty when mature.

Fig. 46 – Neokalmusia didymospora (MFLU 17–0371, new host and geographic record). a–c Appearance of ascoma on host d Longitudinal section of ascoma e Longitudinal section of ostiole f Peridium g Pseudoparaphyses h–j Asci k–o Ascospores p Germinating ascospore. Scale bars: d = 50 µm, e = 50 µm, f = 10 µm, g = 5 µm, h–j = 20 µm, k–p =10 µm.
Culture characteristics – Colony on MEA slow growing, with sparse to moderate aerial mycelium and irregular edge, hairy, above umber colour and whitish in the middle, reverse rust to umber.

Known distribution (based on molecular data) – Germany, Republic of Latvia (Crous et al. 2015b), Ukraine (this study).

Known hosts (based on molecular data) – Betula pendula (Betulaceae), Elaeagnus rhamnoides (Elaeagnaceae), Fagus sylvatica (Fagaceae) (Crous et al. 2015b), Ziziphus jujuba (this study).

Material examined – Ukraine, Donetsk region, Donetsk City, Donetsk Botanical Garden, arboretum, on Ziziphus jujuba Mill. (Rhamnaceae), 18 May 2017, Timur Bulgakov (MFLU 17–2476, new host and geographic record), living cultures MFLUCC 18–0778.

GenBank numbers – LSU: MN244202, SSU: MN244179, ITS: MN244221.

Notes – Our isolate MFLUCC 18–0778 clustered with the ex-type strain of Paracamarosporium fagi (Fig. 45), formerly described by Crous et al. (2015b) from Fagus sylvatica (Fagaceae) in Germany. However, it differs from the type strain of Paracamarosporium fagi in having larger conidiomata (336–449 × 84–277 μm vs. 200 μm diameter). By considering the phylogenetic results, morphology and the host, we consider our collection as a new record on Ziziphus jujuba from Ukraine.

Paraconiothyrium Verkley

Paraconiothyrium was established by Verkley et al. (2004) to accommodate Parac. estuarinum (type species), Parac. brasiliense, Parac. cyclothyrioides and Parac. fungicola. However, subsequent molecular studies resulted in the transfer of a few species from this genus to other genera. Currently, Paraconiothyrium includes 24 accepted species as listed in Index Fungorum (2020).

Paraconiothyrium cyclothyrioides Verkley, da Silva, Wicklow & Crous, Stud. Mycol. 50(2): 330 (2004)

Facesoffungi number: FoF06572

Saprobic on decaying mangrove wood. Sexual morph: Ascomata 200–455 μm high, 150–300 μm diameter (x̄ = 345 × 210 μm, n = 10), lenticular-subglobose, immersed to semi-immersed, mostly covered by the epidermis and perforating the cortex with a small ostiole, thin-walled, black. Necks 170–230 μm long, 45–80 μm wide (x̄ = 285 × 297 μm, n = 10), cylindrical, dark-colored. Peridium 15–35 μm (x̄ = 25 μm, n = 10), composed of 3–5 layers of brown to dark brown cells of pseudoparenchymatous cells, arranged in a textura angularis type. Hamathecium comprising 1.5–2.5 μm wide, filamentous, broad, cellular branched, pseudoparaphyses, anastomosing between and above the asci, embedded in a mucilaginous matrix. Asci 55–120 × 5–10 μm (x̄ = 78 × 8.4 μm, n = 10), 8-spored, bitunicate, cylindrical-clavate, obtusely rounded at the apex, tapering to a short pedicel at the base, thick-walled. Ascospores 10–17 × 2.5–5 μm (x̄ = 12 × 3.8 μm, n = 10), biseriate, dirty olive-brown, cylindrical to ellipsoidal, obtusely rounded at both ends, 3-septate, constricted at the septa, second cell from the top largest. Asexual morph: Undetermined.

Culture characteristics – Ascospores germinating on 50% sea water agar within 24 hours, with germ tubes developed from both ends of ascospores. Colonies on MEA fast growing, reaching 55–70 mm diameter after 15 days of incubation at 25 °C, surface olivaceous to pale olivaceous grey with a hyaline margin, reverse pale brown, floccose, irregular.

Known distribution (based on molecular data) – Papua New Guinea, Central Province, tropical countries (Verkley et al. 2004, Guégan et al. 2016), India (this study).

Known hosts – Soil sample, humans (Verkley et al. 2004, Gordon et al. 2012, Guégan et al. 2016), Suaeda monoica (this study).

Material examined – India, Tamil Nadu, Tiruvarur, Muthupet mangroves (10.4°N 79.5°E), on decaying wood of Suaeda monoica (Amaranthaceae) 30 March 2016, B. Devadatha (AMH-10014, new host record), living culture, NFCCI-4387.
GenBank numbers – ITS: MN242780, LSU: MN241143, SSU: MN241147.

**Fig. 45** – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU and ITS sequence data. Eighty-two strains are included in the combined gene analyses comprising...
2306 characters after alignment (857 characters for LSU, 980 characters for SSU and 467 characters for ITS). *Pleospora tarda* (CBS 714.67), *Pleospora herbarum* (CBS 191.856) and *Pleospora herbarum* (IT 956) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -11598.386568 is presented. The matrix had 816 distinct alignment patterns, with 23.53% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.247509, C = 0.225025, G = 0.279478, T = 0.247988; substitution rates AC = 1.541963, AG = 2.727864, AT = 1.472638, CG = 1.108201, CT = 6.541487, GT = 1.000000; gamma distribution shape parameter α = 0.568674. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.90 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.

**Fig. 46** – *Paracamarosporium fagi* (MFLU 17–2476, new host and geographic record). a, b Appearance of conidiomata on host surface. c Vertical section through the conidioma. d Peridium e–j Conidiogenesis. k–r Conidia s Germinating conidium t, u Culture characteristics on MEA ( t: above view; u: reverse view) Scale bars: a = 500 μm, b = 1000 μm, c = 50 μm, d = 25 μm, e = 10 μm, f, g = 5 μm, h = 10 μm, j = 5 μm, k = 20 μm, l = 5 μm, m–r= 5 μm, s= 10 μm.
Fig. 47 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU and ITS sequence data. Thirty-seven strains are included in the combined gene analyses comprising 2457 characters after alignment (872 characters for LSU, 1035 characters for SSU, 550 characters for ITS). *Nigrograna mycophila* TDK is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -7749.585520 is presented. The matrix had 937 distinct alignment patterns, with 18.57% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.246023, C = 0.226607, G = 0.278252, T = 0.249118; substitution rates AC = 1.565451, AG = 1.781597, AT = 1.275177, CG = 0.882883, CT = 5.629469.
Notes – BLAST search analyses of LSU and ITS sequence data revealed that our taxon is 100% and 99.79% similar to Paraconiothyrium cyclothyrioides. Further phylogenetic analyses based on combined LSU, SSU and ITS showed that our taxon is nested within a monophyletic clade with other Paraconiothyrium cyclothyrioides strains with significant support from ML 82%, MP 76%, and 0.97 BYPP (Fig. 47). However, our collection of Paraconiothyrium cyclothyrioides (NFCCI-4387) did not sporulate even after 2 months’ incubation and hence could not be compared with the asexual morph of Parac. cyclothyrioides (CBS 972.95T) (Verkley et al. 2004). Base pair differences of Paraconiothyrium cyclothyrioides (NFCCI-4387) and Parac. cyclothyrioides (UTHSC: DI16-367) are 1 out of 563bp (0.1%) of ITS gene region which indicates both are similar species. Recent reports support that Paraconiothyrium cyclothyrioides is an emerging opportunistic human pathogen in immunocompromised patients known to cause cutaneous phaeohyphomycosis in renal transplant cases mainly from tropical regions (Gordon et al. 2012, Colombier et al. 2015, Guégan et al. 2016, Valenzuela-Lopez et al. 2017, Garcia-Hermoso et al. 2019). Our sexual morph of Paraconiothyrium cyclothyrioides resembles Parac. thysanolaenae in having cylindrical asci and ascospores that are phragmosporous, ellipsoidal, 3-septate, lack mucilaginous sheaths and constricted at the septa. However, the former is distinct from the latter in having lenticular-subglobose, uniloculate ascomata while the latter produces pseudostromata with uni- to multilocules (Liu et al. 2015). Paraconiothyrium fuckeli, Parac. magnoliae and Parac. cyclothyrioides (NFCCI-4387) share similar ascomata and ascal characters with Parac. cyclothyrioides (NFCCI-4387) (Ariyawansa et al. 2014c, Liu et al. 2015). However, Paraconiothyrium fuckeli differs from Parac. cyclothyrioides (NFCCI-4387) in having narrowly ovoid to clavate ascospores (Ariyawansa et al. 2014c). Paraconiothyrium magnoliae is clearly distinguished from Parac. cyclothyrioides (NFCCI-4387) in having ascospores that are yellowish-brown, with bipolar appendages and a gelatinous sheath (Ariyawansa et al. 2014c).

Paraconiothyrium cyclothyrioides (AMH-10014) shares closer morphological characters with Leptosphaeria peruviana in having immersed ascomata, cylindrical-clavate asc and cylindrical to ellipsoidal, 3-septate, dirty olive-brown ascospores (Pang et al. 2011). However, our taxon is distinct from Leptosphaeria peruviana in having significantly larger subglobose ascomata and asc and narrower ascospores. Leptosphaeria peruviana is a marine species which was described from Salicornia ambiguа (= S. peruviana) by Spegazzini (1881), but has also been reported from mangrove wood (Pang et al. 2011). Leptosphaeria peruviana was poorly described and subsequently Kohlmeyer & Kohlmeyer (1979) examined the scarce type material and its placement under Leptosphaeria (Spegazzini 1881) remains extremely doubtful (Kohlmeyer & Kohlmeyer 1979). This species is very rare and has not been isolated or sequenced.

This is the first report of this taxon as a saprobe and a first report of it being reported from a decaying plant substrate, as otherwise it is widely known as an opportunistic human pathogen. This is also the first report on a sexual morph of this species. This is the first time that this taxon is reported from marine/mangrove environment and its collection from India extends its geographical range.

**Leptosphaeriaceae**

Leptosphaeriaceae species are saprobes, hemibiotrophs or parasites on stems and leaves of herbaceous or woody plants in terrestrial habitats, as well as possibly in aquatic habitats (Hyde et al. 2013, Ariyawansa et al. 2015, Liu et al. 2015, Tennakoon et al. 2017). Barr (1987) introduced Leptosphaeriaceae, and designated Leptosphaeria Ces. & De Not. as the type of the family (Hyde et al. 2013, Wanasinghe et al. 2016). Leptosphaeriaceae species are characterized by immersed or erumpent, perithecial ascomata with single papillate ostioles, fissitunicate, cylindrical asci and
hyaline to brown, transversely septate ascospores (Hyde et al. 2013, Ariyawansa et al. 2015). The asexual morph of taxa in Leptosphaeriaceae can be either coelomycetes or hyphomycetes (Alves et al. 2013, de Gruyter et al. 2013, Hyde et al. 2013, Wanasinghe et al. 2016, Tennakoon et al. 2017). There are 12 genera within Leptosphaeriaceae (Wijayawardene et al. 2020).

**Plenodomus Nees**

*Plenodomus* Preuss was introduced by Preuss (1851) with *P. rabenhorstii* Preuss, as the type species (de Gruyter et al. 2013, Ariyawansa et al. 2015). The type species of *Plenodomus* was replaced by *P. lingam* (Tode) Höhn. (sexual morph: *Leptosphaeria maculans* (Desm.) Ces. & De Not.) by Boerema & Kesteren (1964) due to the type material of *P. rabenhorstii* being lost during World War II (Torres et al. 2005, Ariyawansa et al. 2015). The connection between *L. maculans* (sexual morph) and *P. lingam* (asexual morph) has been confirmed by single spore isolation (Boerema & Kesteren 1964). Boerema et al. (1997) classified *Phoma* species based on their morphological characters in nine sections including *Plenodomus* and each of the sections were systematically illustrated in Boerema et al. (2004). Recently, de Gruyter et al. (2013) reclassified the *Phoma* section of *Plenodomus* with supporting molecular phylogeny. We describe the sexual morph of *Plenodomus collinsoniae* for the first time.

**Plenodomus collinsoniae** (Dearn. & House) Gruyter, Aveskamp & Verkley, Stud. Mycol. 75: 21 (2012)

Facesoffungi number: FoF07379

*Saprobie* on dead branch of an unidentified host. Sexual morph: *Ascomata* 245–260 μm high, 360–390 μm diameter (μx̅ = 255 × 380 μm, n = 20), solitary, scattered, appearing as small, raised black dots on the host surface, superficial or semi immersed, globose to sub globose, smooth, with a flattened top and base and, brownish black, ostiolate, papillate. *Papilla* central circular. *Peridium* 45–55 μm (μx̅ = 52 μm, n = 10) at sides, 20–30 μm (μx̅ = 28 μm, n = 10) at the base, slightly thin at the base, composed of three layers of scleroplectenchymatous cells, inner layer comprising 2–3 cell layers of flattened, light brown cells, arranged in a *textura angularis*, middle layer comprising several layers of subhyaline cells arranged in a *textura globulosa*, outer layer composed of heavily pigmented, thick-walled, dark brown cells, *textura angularis*. *Hamathecium* comprising 1.5–2.5 μm (μx̅ = 2.2 μm, n = 10) wide, septate, cellular pseudoparaphyses, branching between the asci, embedded in a gelatinous matrix. *Asci* 105–210 × 15–25 μm (μx̅ = 170 × 23 μm, n = 20), 8-spored, bitunicate, fissitunicate, cylindrical, short pedicel, apically rounded, with a distinct ocular chamber. *Ascospores* 40–70 × 6–10 μm (μx̅ = 65 × 9 μm, n = 20), overlapping uni- or bi-seriate, olivaceous to yellowish, cylindrical to sub fusoid, with obtuse ends, 6–7 euseptate, not constricted at each septum, cell above central septum slightly wider, guttulate, thick and smooth-walled, with mucilaginous globoid-shaped apical and basal appendages. Asexual morph: Undetermined.

Known distribution (based on molecular data) – China, Japan (De Gruyter et al. 2013, Marin-Felix et al. 2017b), China (this study).

Known hosts – *Collinsonia canadensis*, *Vitis coignetiae*, Unidentified dead wood (De Gruyter et al. 2013, Marin-Felix et al. 2017b).

Material examined – China, Yunnan Province. Kunming, dead branch of an unidentified host, 15 December 2018, V. Thyagaraja, DXH 015 (MFLU 19–2279, new sexual morph record).

GenBank accessions – LSU MN982862, SSU MN982863, ITS MN982858.

Notes – *Plenodomus collinsoniae* is only known from its phoma-like asexual morph (De Gruyter et al. 2013). In our phylogram, new strain (MFLU 19–2279) grouped together with *P. collinsoniae* (CBS 120227) with high statistical support (99 % ML, 1.00 PP) (Fig. 49). Furthermore, DNA sequences of SSU, LSU and ITS of our strain (MFLU 19–2279) are almost similar to *P. collinsoniae* (CBS 120227). Therefore, it is confirmed that our new strain is the asexual morph of *P. collinsoniae*. *Plenodomus collinsoniae* showed close morphological and phylogenetic similarities to the type of the recently introduced *P. sinensis*. However, they are morphologically different by their asci (105–210 × 15–25 vs. 80–100 × 10–12 μm) and ascospore...
sizes (40–70 × 6–10 vs. 27–40 × 3.8–4.4 μm). Additionally, 9 base pairs of the ITS sequence of *Plenodomus collinsoniae* and *P. sinensis* are different out of 536 bp without gaps (1.7 %).

**Fig. 48** – *Paraconiothyrium cyclothyrioides* (AMH-10014, new host record). a Ascomata semi-immersed on decaying wood. b–d Longitudinal sections of ascomata d. Peridial wall layers e Filamentous pseudoparaphyses. g–n Immature and mature asci. l–n Immature and mature ascospores. o, q–v Ascospores. x Germinating ascospore. Scale bars: b–c = 100 μm, f, b, d, f–h = 50 μm, c, e, i–o = 10 μm.
Fig. 49 – Maximum likelihood analysis with 1000 bootstrap replicates yielded a best tree with the likelihood value of -8974.532790. The combined LSU, SSU and ITS sequence dataset comprised 79 strains with *Phaeosphaeria oryzae* (CBS 110110) and *Phaeosphaeriopsis glaucopunctata* (MFLUCC 13-0265) and *Paraphoma radicina* (CBS 111.79) as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum
likelihood analysis. The matrix had 403 distinct alignment patterns, with 10.72% of undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.232805, C = 0.250707, G = 0.235491, T = 0.280998; substitution rates AC = 1.078239, AG = 3.114105, AT = 1.506841, CG = 0.917572, CT = 3.843879, GT = 1.000000; gamma distribution shape parameter α = 0.753063. Maximum likelihood bootstrap (ML) values > 65% and Bayesian posterior probabilities (PP) > 0.80% are given above the nodes. The scale bar indicates 0.04 changes. The ex-type strains are in bold and the new isolate in blue bold.

Lophiostomataceae Sacc.

This family currently comprises 18 genera (Wijayawardene et al. 2018). The latest treatments of Lophiostomataceae is by Hyde et al. (2019) and a new record Vaginatispora nypae is reported here.

Vaginatispora K.D. Hyde

The genus comprises ten species and its members are commonly found on dead wood in aquatic and terrestrial habitats (Hyde et al. 2019, Jayasiri et al. 2019). This genus is characterized by immersed, globose to subglobose, dark ascomata, cylindric-clavate, bitunicate asci and fusiform, septate, guttulate, hyaline ascospores, with a distinctively and thickened mucilaginous sheath. An updated phylogenetic tree is provided here.

Vaginatispora nypae Jayasiri, E.B.G. Jones & K.D. Hyde, Mycosphere 10(1): 84 (2019)

Fig. 52

Facesoffungi number: FoF05264

Saprobic on dried twigs of unidentified host. Sexual morph: Perithecia 305–332 μm high × 269–359 μm diameter (x̄ = 317 × 332 μm, n= 3), immersed, semi-erumpent, solitary, scattered globose to subglobose, dark brown. Peridium 42–96 μm wide, 5–8 layers of dark brown cells of textura angularis, almost black at outside. Hamathecium comprising 3 μm wide, anastomosing, septate, branched, hyaline, pseudoparaphyses. Asci 62–102 × 9–14 μm (x̄ = 87 ×11.5 μm, n= 20), 8-spored, bitunicate, fissitunicate, cylindric-clavate, sessile or with short pedicel, apically rounded, with minutely an ocular chamber. Ascospores 23–33 × 4–7 μm (x̄ = 28 ×6 μm, n = 20), overlapping 1–2-seriate, hyaline, fusiform with narrow, acute ends, 1-septate, constricted at the septum, with large guttules, with a distinctively and thickened mucilaginous sheath, smooth-walled. Asexual morph: Undetermined.

Culture characteristics – Ascospores germinating on MEA within 24 h at room temperature and germ tubes produced from the ends of the ascospore. Colonies on MEA reaching 2.4 cm diameter after 4 weeks at 25 °C. Initially aerial mycelium white, slightly raised, in old cultures dark green to black, flattened on surface, crenate, green to dark green from below, light green margin.

Known distribution (based on molecular data) – Thailand (Jayasiri et al. 2019, this study).

Known hosts (based on molecular data) – On fallen fruit pericarp of Nypa fruticans (Jayasiri et al. 2019), on submerged decaying twigs (this study).

Material examined – Thailand, Phitsanulok, Noen Maprang District, Chomphu, Wat Pa Nong Thap Ruea, submerged decaying twigs, 30 December 2018, S. Boonmee PSL16–0484 (MFLUCC 19–0484, new geographical record); living culture MFLUCC 19–0484.

GenBank numbers – ITS: MN608549, LSU: MN577418, SSU: MN577428, tefL: MN612112.

Notes – Our new isolate Vaginatispora nypae (MFLUCC 19–0484) is phylogenetically related to Vaginatispora nypae (MFLUCC 18–1543) with good statistical support (100 % ML/1.00 BYPP, Fig. 51). This new collection shares similarities in ascomata, asci and ascospores with Vaginatispora nypae (MFLUCC 18–1543, see Fig. 68 in Jayasiri et al. 2019). However, these two strains are from different substrates and geographical locations. Vaginatispora nypae (MFLUCC 18–1543) is described from fallen fruit pericarp of Nypa fruticans (Arecales) from Krabi Province, while our strain is described from dried twigs from Phitsanulok Province. Therefore,
based on morphology and phylogenetic affinity, we identify our strain MFLUCC 19–0484 as *Vaginatispora nypae* and it is reported here as a new geographical record.

**Fig. 50** – *Plenodomus collinsoniae* (MFLU 19–2279, new sexual morph record). a–b Ascomata on host. c Vertical section of ascoma. d Section of peridium. e–f Asci. g Pseudoparaphyses. h–j Ascospores. Scale bars: b–c = 50 μm, d–j = 20 μm, k– o = 5 μm.
Fig. 51 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU, ITS and *tef1* sequence data. Twenty-one taxa are included in the combined gene analyses comprising 3311 characters after alignment (848 characters for LSU, 984 characters for SSU, 585 characters for ITS and 893 characters for *tef1*). *Lentistoma bipolare* (strains KH 216 and KH 214) are used as the outgroup taxa. The best RaxML tree with a final likelihood value of -8141.591393 is presented. The matrix had 450 distinct alignment patterns, with 13.29% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.247229, C = 0.244566, G = 0.270729, T = 0.237477; substitution rates AC = 1.560615, AG = 2.113756, AT = 0.863778, CG = 1.118506, CT = 6.049102, GT = 1.000000; gamma distribution shape parameter α = 0.020000. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.
Fig. 52 – Vaginatispora nypae (MFLU 19–2440, new geographical record). a Appearance of ascomata on woody substrate. b Section of ascoma. c Closed up apical ostiole. d Peridium. e Pseudoparaphyses. f–h Asci. i–l Ascospores (Fig. k stained in nigrosin reagent and Fig. l stained in India ink reagent). m Germinated spore. n–o Colonies on PDA from surface and reverse at 30 days. Scale bars: b = 50 µm, c = 25 µm, d = 25 µm, e = 50 µm, f = 20 µm, g–h = 50 µm, i–l = 20 µm, m = 50 µm.

Lophiotremataceae K. Hiray. & Kaz. Tanaka

Lophiotremataceae was introduced by Hirayama & Tanaka (2011), with Lophiotrema as the type genus. The taxonomic placement of the family was re-evaluated by Hashimoto et al. (2017). Until now, there are six genera accommodated in the family viz. Atrocalyx, Crassimassarina, Cryptoclypeus, Galeaticarpa, Lophiotrema and Pseudocryptoclypeus (Wijayawardene et al. 2018, Jayasiri et al. 2019).

Atrocalyx A. Hashim. & Kaz. Tanaka

Atrocalyx was introduced by Hashimoto et al. (2017), with A. acutisporus as the type species. Until now, there are seven species in this genus reported from Belgium, China, Japan, Spain and Thailand (Hashimoto et al. 2017, de Silva et al. 2018, Jaklitsch et al. 2018, Jayasiri et al. 2019).
**Atrocalyx bambusae** (Phookamsak, S.C. Karunarathna & K.D. Hyde) N.I. de Silva & K.D. Hyde, Phytotaxa 333: 204 (2018)

Facesoffungi number: FoF06536

*Saprobi*c on dead bamboo culms, forming wedged-shaped, blackened perithecia. Sexual morph: *Ascomata* 345–465 μm diameter, 130–200 μm high, scattered, solitary, gregarious, immersed to semi-immersed in host cortex, dark brown to black, conical to subglobose, uniculate, coriaceous. *Peridium* composed of host and fungal tissues; laterally 25–35 μm thick, composed of several layers of dark brown to hyaline cells of *textura angularis*; basal part, with 50–80 μm thick sides, comprising dark brown to hyaline cells, arranged in *textura epidermoidea*. *Hamathecium* comprising 1–1.3 μm broad, septate, branched, anastomosing, hyaline, filiform, pseudoparaphyses. *Asci* 70–95 × 7.5–10 μm (\(\bar{x} = 82.5 \times 8.7 \mu m, n = 20\)), 8-spored, bitunicate, fissitunicate, cylindrical-clavate, curved, with short furcate pedicel, apical rounded with developed ocular chamber. *Ascospores* 15–23 × 3.5–4.5 μm (\(\bar{x} = 19 \times 4 \mu m, n = 20\)), 2-seriate, hyaline, fusiform, 1- or 3-septate, constricted at the septa, straight to curved, smooth-walled, guttulate. Asexual morph: Undetermined.

Culture characteristics – Colonies on PDA reaching 20–25 mm diameter after 2 weeks at room temperature (20–30 °C), dark brown to black in the centre, brown, radiated at the margin from above and reverse; medium dense, woolly, irregularly flattened, slightly raised at the middle; not producing pigment in agar.

Known distribution (based on molecular data) – Thailand, Chiang Rai (Hyde et al. 2016), China, Yunnan (this study).

Known hosts (based on molecular data) – dead bamboo culms (Hyde et al. 2016; this study).

Material examined – China, Yunnan Province, Southwest Forestry University, Bamboo Garden, on dead bamboo culms (Poaceae), 8 August 2016, H.B. Jiang, XNLD001 (KUN–HKAS 101764, new geographic record), living culture KUMCC 18–0117.

GenBank numbers – ITS: MN511733, SSU: MN511734, tef1: MN525566.

Notes – *Lophiotrema bambusae* was introduced by Hyde et al. (2016) based on morphological and phylogenetic analyses. However, the species was transferred to *Atrocalyx* and became *Atrocalyx bambusae* based on combined SSU, ITS, LSU and tef1 sequences analyses (de Silva et al. 2018). Morphological characteristics of our collection are similar to *Atrocalyx bambusae*. Based on a nucleotide comparison of ITS and tef1 pairwise (Jeewon & Hyde 2016), the new isolate has consistent with the base pairs with the type strain of *Atrocalyx bambusae* (Fig. 53). Thus, we identify the new collection as *Atrocalyx bambusae*. Because the record of *A. bambusae* was so far only reported from Thailand (Hyde et al. 2016), we report this species from China for the first time.

**Lophiotrema** Sacc., *Michelia* 1 (3): 338 (1878)

*Lophiotrema* established by Saccardo is a group that comprised of lophiostomataceous taxa, characterised by hyaline, multi-septate ascospores (Saccardo 1883). However, it was later treated as a synonym of *Lophiostoma* as they pointed out that both ascospore colour and number of transverse septa are not sufficient for generic distinctions (Chesters & Bell 1970). However, Hirayama & Tanaka (2011) separated *Lophiotrema* from *Lophiostoma* based on morphological traits and phylogenetic analyses, and established *Lophiotremataceae* to accommodate *Lophiotrema*.

**Lophiotrema neohysterioides** M.E. Barr, Mycotaxon 45: 208, 1992

Facesoffungi number: FoF 05217

*Saprobi*c on dead wood of herbaceous plant. *Ascomata* 250–300 μm high × 270–320 μm diameter (\(\bar{x} = 298 \times 311 \mu m, n = 10\)), dark brown to black, scattered, gregarious, immersed to erumpent throughout the host tissue, globose to subglobose, uniloculate, glabrous, coriaceous. *Ostiole* central, carbonaceous, with a crest-like opening. *Peridium* 15–20 μm thick, comprising several layers, an inner stratum with hyaline to light brown cells of *textura angularis* and an outer stratum with light brown to dark brown cells of *textura angularis* fusing with the host tissues.
Hamathecium comprising 1.5–2 μm wide, branched, cellular pseudoparaphyses, anastomosing among and between the asci. Asci 70–90 × 7–10 μm (\( \bar{x} = 75 \times 10 \) μm, n = 20), 8-spored, bitunicate, fissitunicate, cylindric-clavate, short pedicellate, apically rounded with a minute ocular chamber. Ascospores 20.5–25 × 3–5 μm (\( \bar{x} = 23 \times 4 \) μm, n = 20), hyaline, fusiform, 3-septate, with the primary septum median, the second cell from the apex slightly enlarged downward and shortest although longer than wide, with guttules, smooth, no sheath or appendages detected.

**Fig. 53** – Phylogram generated from maximum likelihood analysis based on a combined ITS, LSU, SSU and tef1 sequence dataset. Twelve strains are included in the combined gene analyses comprising 3,354 total characters including gaps (ITS: 1–566 bp, LSU: 567–1426 bp, SSU: 1427–2453 bp, tef1: 2454–3354 bp). The best scoring of the ML tree is selected to represent the phylogenetic relationships of Atrocalyx species, with the final ML optimization likelihood: -8153.910961. The matrix had 366 distinct alignment patterns, with 16.26% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.242968, C = 0.259079, G = 0.267179, T = 0.230774; substitution rates AC = 1.994630, AG = 3.374777, AT = 1.799513, CG = 1.818183, CT = 10.748211, GT = 1.000000; Tree-Length = 0.634901; gamma distribution shape parameter \( \alpha = 0.585697 \); The proportion of invariable sites I = 0.547872. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.002895. Bootstrap support for maximum likelihood (MLBP) greater than 70% and Bayesian posterior probabilities (BYPP) greater than 0.95 are defined above the nodes as MLBP/BYPP. The tree is rooted to Pseudocryptocylnus yakushimensis (KT 2186). The type
strains are indicated in bold and newly generated sequence is shown in blue.

**Fig. 54** – *Atrocalyx bambusae* (KUN-HKAS 101764, new geographic record). a–c Ascomata on bamboo host. d Vertical section of ascoma. e, f Peridium. g Pseudoparaphyses. h–k Asci. l Culture characteristics. m–o Ascospores. p Germinating ascospore. Scale bars: d = 100 μm, e = 30 μm, f, h–k, p = 15 μm, g = 10 μm, m–o = 5 μm.
Fig. 55 – Phylogram generated from maximum likelihood analysis based on combined SSU, ITS, LSU and tef1 partial sequence data. Thirty one strains were included in the sequence analysis, which comprised 2545 characters including alignment gaps. Lophiotrema spp. (Lophiostomataceae) were used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring RAxML tree with a final likelihood value of -34728.193523 is presented. The matrix had 286 distinct alignment patterns, with 4.84% of undetermined characters or gaps. Estimated base frequencies were as follows; A = 0.253516, C = 0.226561, G = 0.278443, T = 0.241480; substitution rates AC = 1.845417, AG = 3.129372, AT = 1.390649, CG = 1.720102, CT = 9.650566, GT = 1.000000. ML bootstrap support (first set) equal or greater than 70 % and Bayesian posterior probabilities equal or greater than 0.95 are given near to each branch. The new strain is in blue.

Known distribution (based on molecular data) – Japan (Hashimoto et al. 2017), China (Zhang et al. 2018).

Known hosts (based on molecular data) – *Phyllostachys bambusoides, Robinia pseudoacacia* (Hashimoto et al. 2017, Zhang et al. 2018), *Pinus* sp. (this study).

Material examined – China, Yunnan, Kunming Institute of Botany, on decaying wood of *Pinus* sp. (Pinaceae), 25 June 2018, A. Ekanayaka, C 467 (MFLU 18–2171, new host record).
Fig. 56 – *Lophiotrema neohysterioides* (MFLU 18–2171, new host record). a Ascomata on host substrate. b, c Vertical section through ascomata. d–g Asci h–i Ascospores. Scale bars: a = 500 µm, b–c = 100 µm, d–g = 20 µm, h–l = 10 µm.

GenBank numbers – ITS: MN862638, LSU: MN862641, SSU: MN862640, *tef1*: MN862639.

Notes – The new collection shares similar morphological characters and DNA sequence data with *Lophiotrema neohysterioides*. Our collection is morphologically similar to the latter in having clavate asci and hyaline, fusiform, three equidistant septate ascospores as described in Tanaka & Harada (2003). A comparison of the SSU, ITS, LSU and *tef1* nucleotides of these strains revealed less than ≤ 1.5% nucleotide differences which indicates that our isolate is a new record of *Lophiotrema neohysterioides* (Jeewon & Hyde 2016). This is the second record of *Lophiotrema neohysterioides* from China, with a previous record which was collected from wood of an unidentified plant (Zhang et al. 2018).
**Massariaceae** Nitschke

Massariaceae was introduced by Nitschke (1869) to accommodate the type genus *Massaria* and is a well-resolved family in Pleosporales. The species of Massariaceae have been reported as hemibiotrophs, saprotrophs or weak pathogens. Consequently, another monotypic genus *Neomassaria* was introduced by Hyde et al. (2016). The family is characterized by relatively large pseudothecia, central or eccentric ostioles, relatively wide, firm, opaque compressed cells of *textura angularis*, darkly pigmented peridium, branching and anastomosing pseudoparaphyses, 4–8-spored, bitunicate, fissitunicate asc and relatively large, oblong, cylindrical, ellipsoidal or fusoid brown ascospores. Huanraluek et al. (2018) introduced a new genus *Massarioiramusculicola* based on phylogenetic analysis of large subunit (LSU) rRNA sequence data and morphological studies. The asexual morph is coelomycetous where known. Species of Massariaceae are distributed on a wide range of hosts namely, Aceraceae, Fabaceae, Moraceae, Rosaceae, Rutaceae, Ulmaceae and Viburnaceae worldwide especially in North America and Europe.

**Massaria** De Not.

_Massaria_ was introduced by De Notaris with _Sphaeria inquinans_ (= _Massaria inquinans_ as type species. The genus is characterized by large subglobose, pyriform to strongly depressed pseudothecia, central or eccentric ostioles, a wide, firm, opaque pseudothecial wall composed of numerous rows of thick-walled cells of *textura angularis*, numerous persistent, indistinctly septate anastomosing pseudoparaphyses, bitunicate, fissitunicate, cylindrical or fusoid asci and large oblong, cylindrical, ellipsoidal or fusoid, hyaline, light to dark brown ascospores, rounded or tapered to subacute ends. Voglmayr & Jaklitsch (2011) studied 17 species (with seven new species) and also reported their occurrence predominantly on _Acer_ and Rosaceae hosts. Currently, 31 species are listed under _Massaria_. The asexual morph is coelomycetous.

*Massaria anomia* (Fr.) Petr., Anns Mycol. 21:114 (1923)  
Facesoffungi number: FoF 06536

_Saprobic_ on dead branch of _Robinia pseudoacacia_. Sexual morph: _Ascomata_ (pseudothecia) relatively large 500–1500 µm wide, 800–1400 µm wide, scattered or clustered, pyriform to strongly depressed, immersed in bark and typically in pseudostromatic tissues intermixed with substrate cells, with blackened zones, clypeate, and ostiolate. _Ostioles_ 200–325 µm long, 250–400 µm wide, central, long, solitary projecting through the bark, stout papillate, surmounted by peaks of stromatic tissues that form coarsely sulcate tips above the bark surface. _Peridium_ 29–43 µm relatively wide, firm, opaque, composed of numerous rows of thin-walled, darkly pigmented, compressed cells of *textura angularis*. _Hamatheecium_ comprising 1–2 µm of numerous persistent, indistinctly sepatate, branching and anastomosing pseudoparaphyses embedded in a gelatinous matrix. _Asci_ 115–194 × 18–24 µm (\( \bar{x} = 150.7 \times 21.4 \) µm, n = 20), 4–spored, bitunicate, fissitunicate, basal and peripheral, oblong, cylindrical or fusoid, pedicellate, apically with wide oculur chamber (2–5 µm) and refractive ring. _Ascospores_ 46–49 × 13–14 µm (\( \bar{x} = 47.9 \times 13.9 \) µm, n = 20), 1–2-seriate, dark brown, oblong, cylindrical, ellipsoidal or fusoid, rounded or tapered to subacute ends, hyaline or light to dark brown, slightly inequilateral, symmetric, biconid and symmetrically 1-euspatate in early states, becoming 3-distoseptate at maturity, slightly constricted at the septa, secondary septa usually closer to primary septum than to ends of ascospore wall, smooth, without mucilaginous sheath, dark-brown lumina rhomboid or lenticular in the central cells, conid in the end cells.  
Asexual morph: Undetermined.

Culture characters – Colonies growing on MEA, reaching 7 mm diameter after 7 d at 16 °C, circular to slightly irregular, flat to slightly raised, dense, surface white, middle dark-grey, reverse saffron to dark-brown, white at the margin, smooth surface with entire to slightly undulate edge.

Known distribution (based on molecular data) – Austria, France, Northern Hemisphere (North America, Europe) (Voglmayr & Jaklitsch 2011), Italy (this study).

Known hosts (based on molecular data) – on woody host of Fabaceae (Amorpha, Gleditsia, Laburnum) (Voglmayr & Jaklitsch 2011), Robinia pseudacacia (Fabaceae) (this study).
Fig. 57 – Phylogram generated from maximum likelihood analysis based on combined LSU and SSU sequence data. Sixty-nine strains are included in the combined gene analyses comprising 1939 characters after alignment (906 characters for LSU and 1035 characters for SSU). *Delitschia didyma* (UME 31411) and *Delitschia winteri* (CBS 225.62) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -5560.059600 is presented. The matrix had 298 distinct alignment patterns, with 32.99% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.262352, C = 0.201082, G = 0.289067, T =
Material examined – Italy, Province of Forlì-Cesena [FC], near Predappio, on dead terrestrial branch of *Robinia pseudoacacia* (Fabaceae), 11 February 2017, E. Camporesi (MFLU 17–0490, new geographical record); living culture MFLUCC 18–1128.

GenBank numbers – LSU: MN244203, SSU: MN244180.

Notes – Our taxon is morphologically similar and phylogenetically related to *Massaria anomia*, but collected in different geographical locations. Both are associated with the same host, *Robinia pseudoacacia* (Fabaceae). *Massaria anomia* was recorded in Austria, France and is widely distributed in the Northern Hemisphere (Voglmayr & Jaklitsch 2011) while our collection is from Italy. The phylogenetic placement of our strain (MFLUCC 18–1128) is shown in Fig. 57.

**Massarinaceae** Munk

Massarinaceae was introduced by Munk (1956) with *Massarina* as the type genus. The morphological characters of this family include immersed, flattened or spherical ascomata, cellular pseudoparaphyses, clavate to cylindro-clavate asci, and hyaline, fusiform to narrowly fusiform, 1 to 3-septate ascospores with or without a mucilaginous sheath. Barr (1987) treated Massarinaceae as a synonym of Lophiostomataceae. These two families are now recognized as different lineages based on morphological and molecular data (Liew et al. 2002, Zhang et al. 2009). Species of Massarinaceae are distributed in terrestrial habitats and are saprobic on wood or twigs (Hyde et al. 2013). *Massarina* was established to separate taxa with hyaline ascospores based on *Massarina, Keissleriella, Metasphaeria, Pseudotrichia* and *Trichometasphaeria* (Munk 1956, Hyde et al. 2013). Following its introduction, many studies have been conducted on the above mentioned genera, with the exception of *Massarina* and have been transferred to other families (Suetrong et al. 2009, Zhang et al. 2012, Hyde et al. 2013, Wijayawardene et al. 2014, Boonmee et al. 2017). Currently, nine genera are accommodated in Massarinaceae namely, *Bambusistroma, Helminthosporium, Longiostiolum, Massarina, Pseudodidymosphaeria, Pseudosplanchnonema, Semifissispora, Stagonospora* and *Suttonomyces*.

**Pseudosplanchnonema** Chethana & K.D. Hyde

The monotypic genus *Pseudosplanchnonema* was introduced by Chethana et al. (2015) to accommodate *Pseudosplanchnonema phorcioides*, a new combination proposed by the authors earlier described as *Massaria phorcioides*. The genus is characterized by immersed, perithecial ascomata, short papillate ostioles, filiform pseudoparaphyses embedded in a gelatinous matrix, bitunicate cylindrical to clavate short pedicellate asci and brown ascospores with pseudosepta surrounded by a mucilaginous sheath. The asexual morph is coelomycetous. *Pseudosplanchnonema phorcioides* has been reported on dead branch of *Morus* sp. in Italy. In this study, we illustrate a new record of *Pseudosplanchnonema phorcioides* on *Morus* sp in Russia.

**Pseudosplanchnonema phorcioides** (I. Miyake) Chethana, Camporesi & K.D. Hyde, Phytotaxa 231(2): 139 (2015)

Facesoffungi number: FoF06225

*Saprobic* on dead branch of *Prunus tomentosa*. Sexual morph: *Pseudostromata* 51–102 µm diameter 56–80 µm high, immersed to semi immersed ascomata, scattered or clustered, globose, conical globose, papillate. *Ostiole* 25–27 µm high, 49–52 µm diameter (µ = 25.2 × 50.6 µm, n = 10), short, papillate, opening to exterior through bark. *Peridium* thin 43–63 µm (µ = 52.3 µm), wide at side walls, up to 85 µm wide near the apex, and 47 µm wide at the base comprising a few layers of brown compressed cells, outer layer composed of 3–4 layers of dark brown cells of *textura*
angularis, middle layer comprising 2–3 layers cells of textura angularis and inner cells comprising pale brown cells of textura prismatica. Hamathecium comprising broad, filiform 2–3 (μm = 3.2) μm wide, septate pseudoparaphyses, embedded in a gelatinous matrix. Asci 201–255 × 41–48 μm (μm = 232.3 × 43.9 μm, n = 10), 8-spored, bitunicate, fissitunicate, cylindrical to clavate, with a short pedicel with an ocular chamber. Ascospores 40–48 × 15–17 μm (μm = 51.1 × 16.6 μm, n = 15), overlapping biseriate, hyaline to pale brown when young, dark brown at maturity, fusiform to ellipsoidal, widest near the centre, with acute rounded ends, sometimes slightly curved, 1-submedian septate, constricted at the septum, 5–6-guttulate, pseudosepta between the guttules, smooth-walled, surrounded by a mucilaginous sheath. Asexual morph: Undetermined.

Culture characters – Ascospores germinating on MEA within 24 h and the germ tubes of 2.5–3 μm diameter produced near the septum. Colonies reaching 4 mm diameter after 10 days at 25 °C, entire edge, greenish black in the centre, greenish grey towards rim, white at the margin with a circular cottony mycelium on the surface and reverse black in the centre and grey towards the ends of the mycelium.

Known distribution (based on molecular data) – Russia, Austria, France, Northern Hemisphere (Chethana et al. 2015), Russia (this study).

Known hosts (based on molecular data) – Morus sp. (Chethana et al. 2015), Prunus tomentosa (this study).

Material examined – Russia, Rostov Region, Shakhty City, urban artificial forest, on dying twigs of Prunus tomentosa (Rosaceae), 26 May 2017, Timur Bulgakov (MFLU 17–2062, new host record), living cultures MFLUCC 18–1135; DSM 109792.

GenBank accession numbers LSU: MN244204, SSU: MN244181.

Notes – Our isolate MFLUCC 18–1135 clustered with the ex-type strains of Pseudosplanchnonema phorcioides (Fig. 59) originally described by Chethana et al. (2015) and collected from Italy. However, it differs from the type strain of P. phorcioides in having smaller ascomata (51–102 μm versus 159–483 μm diameter) and shorter ascospores (40–48 × 15–17 μm versus 50–66 × 14–20 μm). By considering the molecular data and the host relationship, we illustrate our collection as a new record on Prunus tomentosa from Russia.

Melanommataceae G. Winter.

We follow the latest treatment and updated account of Melanommataceae in Tian et al. (2015), Li et al. (2016d, 2017a), Hyde et al. (2018) and Wanasinghe et al. (2018a). Melanommataceae comprises 33 genera (Wijayawardene et al. 2020).

Byssosphaeria Cooke.

Byssosphaeria was introduced by Cooke (1879) and is typified by B. keithii. The genus is widespread on hosts and distribution, occurring on various hosts as a saprobe worldwide (Farr & Rossman 2020). Recently, there are 46 epithets listed in Index Fungorum (2020), however, only nine species have their phylogenetic affinities investigated within Melanommataceae (Tennakoon et al. 2018). We follow the updated accounts of Byssosphaeria in Tennakoon et al. (2018). Byssosphaeria musae is reported from decaying frond of a palm in Yunnan Province, China for the first time.

Byssosphaeria musae Phookamsak & K.D. Hyde, Fungal Diversity 72: 119 (2015) Fig. 62

Facesoffungi number: FoF00436

Saprobic on decaying frond of palm. Sexual morph: Ascomata 525–600 μm high, 560–640 μm diameter, black, superficial, solitary, gregarious, uni-loculate, subglobose, setose, with dark brown to black, filiform, filamentous setae, tapering towards the apex, apapillate, ostiole central with pore-like opening, producing yellowish to orange pigmented around the ostiole at maturity. Peridium 45–60 μm wide at side, 75–100 μm wide at base, 70–90 μm wide near ostiole, thick-walled, composed of dark brown to black pseudoparenchymatous cells, arranged in textura angularis to textura prismaticata.
Fig. 58 – *Massaria anomia* (MFLU 17–0490, new geographical record). a, b Appearance of ascomata on host surface. c Vertical section through the ascomata. d Ostiole. e Peridium f–h Asci. i–l Ascospores m Germinating ascospores n, o Culture characteristics on MEA (n: above view, o: reverse view). Scale bars: a = 400 μm, b = 500 μm, c = 50 μm, d = 30 μm, e = 50 μm, f = 40 μm, g, h = 50 μm, i–m = 20 μm.
Fig. 59 – Phylogram generated from maximum likelihood analysis based on combined LSU and SSU sequence data. Twenty-one strains are included in the combined gene analyses comprising 2327 characters after alignment (856 characters for LSU, 1058 characters for SSU and 411 characters for TUB). *Paracamarosporium hawaiiense* (CBS 120025) and *Paraconiothyrium fungicola* (CBS 113269) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -7593.818950 is presented. The matrix had 535 distinct alignment patterns, with 30.91% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.254460$, $C = 0.214516$, $G = 0.284400$, $T = 0.246625$; substitution rates $AC = 1.248767$, $AG = 2.513663$, $AT = 1.269748$, $CG = 0.839642$, $CT = 4.977442$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.917153$. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.

*Hamathecium* comprising dense, 1–2.7 μm wide, septate, filiform, anastomosed pseudoparaphyses, embedded in a gelatinous matrix. *Asci* 126–147 × 11–15.5 μm ($\bar{x} = 136.8 \times 13$ μm, $n = 20$), 8-spored, bitunicate, cylindric-clavate, rounded at the apex with ocular chamber, with a long pedicel (45–65 μm long). *Ascospores* 33–37 × 7–9 μm ($\bar{x} = 35.5 \times 7.8$ μm, $n = 20$), overlapping 1–2-seriate, initially hyaline, light brown at maturity, fusiform, 1-septate, slightly
curved, obviously constricted at the septum, guttulate, surrounded by a thin gelatinous sheath. Asexual morph: Undetermined.

Fig. 60 – *Pseudosplanchnonema phorcioides* (MFLU 17–2062, new host record). a, b Appearance of ascomata on host surface. c Vertical section through the ascoma. d Peridium. e Hamathecium. f–h Asci i–p Ascospores (p: Indian ink showing surrounding mucilaginous sheath). q Germinating ascospores. r, s Culture characteristics on MEA (q: above view; r: reverse view). Scale bars: a = 200 μm, b, c = 100 μm, d = 5 μm, e–g = 10 μm, h = 20 μm, i–m = 10 μm.
Culture characteristics – Colonies on PDA reaching 18 mm diameter after 1 week at room temperature 20–25 °C, circular, convex, effuse, velvety to fluffy, light yellow to orange from the above; yellowish-brown at the centre, yellowish brown to orange at the margin from the reverse; not producing pigments in PDA.

Known distribution (based on molecular data) – Thailand, Chiang Rai (Liu et al. 2015), China, Yunnan (this study).

Known hosts (based on molecular data) – Musa sp. (Musaceae) (Liu et al. 2015), Palm (this study).

Material examined – China, Yunnan Province, Kunming, Dianchi Lake, on decaying frond of a palm, 25 January 2019, E.F. Yang, DY1 (KUN-HKAS 102550, new host record), living culture, KUMCC 20–0003.

GenBank numbers – LSU: MT000759, SSU: MT000958, tef1: MT005775.

Notes – Byssosphaeria musae Phookamsak & K.D. Hyde was introduced by Liu et al. (2015). The taxon was collected from Musa sp. (Musaceae) in Chiang Rai Province, Thailand. Our new collection was collected from a palm nearby Lake Dianchi of Kunming, Yunnan Province, China. Morphologically of our collection is similar to B. musae. Phylogenetic analyses demonstrate our new collection is conspecific with the type of B. musae (Fig. 61). A comparison of the tef1 nucleotides of these two strains reveals less than 1.5% nucleotide differences (Jeewon & Hyde 2016). Based on known distribution and host of B. musae (Liu et al. 2015), the new collection is reported as a saprobe on palm in Yunnan, China for the first time.

**Morosphaeriaceae** Suetrong, Sakay., E.B.G. Jones & C.L. Schoch

This family comprises four saprobic genera namely *Aquilomyces, Clypeolocus, Helicascus* and *Morosphaeria* and are commonly found from both freshwater and marine habitats (Suetrong et al. 2009, Devadatha et al. 2018, Wijayawardene et al. 2018, Zeng et al. 2018). The family is characterized by subglobose to lenticular, erumpent ascomata, cylindric-clavate asci with an apical ring and fusiform to ellipsoidal, hyaline to pigmented ascospores, surrounded by thick gelatinous sheath.

**Helicascus** Kohlm.

The genus currently comprises twelve species (Kohlmeier 1969, Zeng et al. 2018, Index Fungorum 2020). Its members are saprobic on dead or decaying wood in aquatic habitats from freshwater and marine environments. They share common characters such as immersed to erumpent with uni- to multi-loculate ascomata, bitunicate asci and septate, pigmented ascospores, with or without mucilaginous sheath (Luo et al. 2016, Zeng et al. 2018). An updated tree is provided here.

**Helicascus elaterascus** (Shearer) H. Zhang & K.D. Hyde, Cryptog. Mycol. 33: 158 (2012)

Facesoffungi number: FoF02018

*Saprobic* on decaying, submerged wood in freshwater. Sexual morph: *Ascomata* 200–300 μm high × 165–180 μm diameter, immersed, solitary, aggregated, subglobose to ellipsoidal, black, periphysate ostiolar neck, papillate. *Peridium* 26–55 μm, composed of several layers of brown to dark brown cells of *textura angularis*, outer layer dark brown, inner layer of hyaline, flattened cells of *textura angularis*, 2–3 μm wide. *Asci* 88–130 × 18–23 μm (x̄ = 101 × 20, n = 15), 8-spored, bitunicate, cylindric-clavate, long pedicellate, form a long tail-like extension, apically rounded. *Ascospores* 20–27 × 8–12 μm (x̄ = 25 × 10, n = 20), 2-seriate partially overlapping, hyaline when young and brown when mature, ellipsoid-fusiform, upper cell wider, tapering towards the narrow end, 1-septate, constricted at the septum, refractively guttulate, smooth-walled, with or without mucilaginous sheath. Asexual morph: Undetermined.
Fig. 61 – Phylogram generated from maximum likelihood analysis based on a combined LSU, \textit{tef1} sequence dataset. Twenty-two strains are included in the combined gene analyses comprising 1731 total characters including gaps (LSU: 1–847 bp, \textit{tef1}: 848–1731 bp). The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring of the ML tree is selected to represent the phylogenetic relationships of taxa in \textit{Byssosphaeria}, with the final ML optimization likelihood: -4023.863653. The matrix had 225 distinct alignment patterns, with 22.61% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.239041, C = 0.253223, G = 0.291157, T = 0.216579; substitution rates AC = 18.961935, AG = 44.431399, AT = 17.633231, CG = 23.885049, CT = 290.21010184, GT = 1.000000; Tree-length = 0.230788; gamma distribution shape parameter $\alpha = 1.795248$. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.002700. Bootstrap support for maximum likelihood (ML) greater than 60% and Bayesian posterior probabilities (PP) greater than 0.90 are defined above the nodes as ML/PP. The tree is rooted to \textit{Melanomma japonicum} (CBS 142905) and \textit{M. pulvis-pyrius} (CBS 124080). The type strains are indicated in bold and the newly generated sequence is shown in blue.
**Fig. 62** – *Byssosphaeria musae* (KUN-HKAS 102550, new host record). a Ascomata on host. b Close-up of ascomata. c Section of ascoma. d Ostiole. e Peridium at the base. f Setae. g Section of peridium at side. h Pseudoparaphyses. i–l Asci (k = stained in Indian ink). m–o Ascospores. p Germinated ascospore. r Colony from above. s Colony from below. Scale bars: c = 200 μm, d–f = 100 μm, g, h = 50 μm, i = 30 μm, j–l, p = 20 μm, m–o = 10 μm.

Culture characteristics – Ascospores germinated on WA within 24 hours. Colonies on MEA, circular, flat surface, wavy margin, reaching 0.2–0.5 cm diameter in 5 days at 25 °C, at first white to pale yellow, later becoming brown, white and brown at the edge and dark brown at reverse side.
Fig. 63 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS and tef1 sequence data. Thirty-three taxa are included in the combined gene analyses comprising 3022 characters after alignment (1340 characters for LSU, 722 characters for ITS, 960 characters for tef1). Montagnula opulenta (CBS 168.34) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -9903.231098 is presented. The matrix had 861 distinct alignment patterns, with 35.34% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.246736, C = 0.232999, G = 0.271140, T = 0.249124; substitution rates AC = 0.687317, AG = 1.868367, AT = 1.195520, CG = 0.682068, CT = 5.607541, GT = 1.000000; gamma distribution shape parameter α = 0.238117. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains and/or reference specimens are in bold and black. The newly generated sequence is indicated in blue.
**Fig. 64** – *Helicasus elaterascus* (MFLU 18–1089, new geographical record). a–d Appearance of masses of ascospores on host surface. e, f Section of ascoma. g Peridium, h Ostiole section. i Pseudoparaphyses. j–p Asci. m, o Exotunica of asci showing ‘jack-in-a-box’ action. q–s Ascospores. t, u Ascospore germination. v, w Culture on MEA from surface and reverse after 3 weeks. Scale bar: a = 400 µm, b–d, f = 200 µm, e, g, i, k, l, n–p, u = 50 µm, h = 100 µm, j, m, q–t = 20 µm.

Known distribution (based on molecular data) – Hong Kong (Kodsueb et al. 2004), Japan (Tanaka et al. 2015), Thailand (this study).

Known substrates (based on molecular data) – On dead wood and submerged twigs of woody plant of unidentified hosts (Tanaka et al. 2015), on submerged wood (Kodsueb et al. 2004, this study).
Material examined – Thailand, Phayao, on submerged wood in a freshwater stream, 11 September 2017, S. Boonmee, PK98 (MFLU 18–1089, new geographical record); living culture (MFLUCC 17–2573).

GenBank numbers – ITS: MN608548, LSU: MN577417, tef1: MN612111.

Notes – *Helicascus elaterascus* (MFLUCC 17–2573) was collected from decaying submerged wood in freshwater stream in Phayao Province, Thailand. Our strain shares similarities in ascomata, asci and ascospores with *Helicascus elaterascus* strains KT 2673, KT 2682 and HKUCC 7769. Multi-gene phylogenetic analysis placed our strain among *H. elaterascus* strains with 67% ML support (Fig. 63).

**Fig. 65** – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, and RPB2 sequence data. Twenty-four strains are included in the combined gene analyses comprising 2428 characters after alignment (842 characters for LSU, 633 characters for ITS, 953 characters for RPB2). *Melanomma pulvis-pyrius* (CBS 124080) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -13896.135660 is presented. The matrix had 936 distinct alignment patterns, with 19.07% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.249814, C = 0.243135, G = 0.282173, T = 0.224878; substitution rates AC = 1.382955, AG = 3.921767, AT = 1.529942, CG = 0.881236, CT = 8.046891, GT = 1.000000; gamma distribution shape parameter α = 0.204652. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.

This species has been reported from submerged wood in a freshwater habitat from Japan and Hong Kong (Tanaka et al. 2015, Kodseeub et al. 2004). In this study, we, therefore, report our strain *Helicascus elaterascus* (MFLUCC 17–2573) as a new geographical record from northern Thailand.
Paradictyoarthriniaceae Doilom, Jian K. Liu & K.D. Hyde

Paradictyoarthriniaceae was established by Liu et al. (2015) to accommodate a single genus Paradictyoarthrinium. Wanasinghe et al. (2018a) introduced another genus Xenomassariosphaeria Jayasiri, Wanas. & K.D. Hyde which was on dead aerial spines of Rosa canina (Rosaceae).

Paradictyoarthrinium Matsush.

Paradictyoarthrinium was typified by *P. diffractum* and accommodated in Paradictyoarthriniaceae by Liu et al. (2015). Paradictyoarthrinium is a hyphomycetous genus and includes four species in Index Fungorum (2020) with molecular data confirmed. Paradictyoarthrinium aquatica and *P. diffractum* were collected on submerged wood in freshwater habitat, while *P. hydei* and *P. tectonicola* were from decaying wood in terrestrial habitats (Liu et al. 2015, 2018a). Until now, there is still no sexual morphs reported in the genus.

Paradictyoarthrinium diffractum Matsush., Matsush. Mycol. Mem. 9:18 (1996) Fig. 66

Facesoffungi number: FoF01854

*Saprobic* on dead wood of Mangifera indica. Sexual morph: Undetermined. Asexual morph: *Colonies* on natural substrate, superficial, black, gregarious, scattered, powdery. *Conidiophores* up to 25 μm long, 2–6 μm wide, macronematous, rarely micronematous, erect to slightly curved, dark brown, slightly constricted at the septa, septate, branched to unbranched, arising from hyphae. *Conidiogenous cells* holoblastic, integrated, terminal, indeterminate. *Conidia* 13–28 × 9–17 (bar = 20 × 13 μm, n = 30), muriform, initially globose to subglobose, brown to dark brown when young, ellipsoidal to irregular in shape, brown to black at maturity, solitary or developing in branched chains, with 1–2-spored short chains, verrucose, very variable in size and shape; circular to irregular with a protruding basal cell; rounded to truncate at the base.

Culture characteristics – Conidia germinating on PDA within 24 h. Germ tubes produced around conidia. Colonies on PDA edge entire, velvety, medium dense, flat or effuse, brownish grey. *Mycelium* 1.5–3.8 μm broad, partly superficial, partly immersed, pale brown to dark brown, septate. *Conidiophores* up to 19 μm long, 3.5–5.5 μm wide, micronematous, erect to slightly curved, dark brown, constricted at the septa, septate, branched to unbranched, arising from laterally and terminally on hyphae. *Conidiogenous cells* holoblastic, terminal. *Conidia* 7–24 × 7–21 (bar = 14 × 13.5 μm, n = 30), initially globose to subglobose, ellipsoidal to irregular shape, muriform, brown to dark brown, mostly with 1-spored chains, very variable in size and shape; circular to irregular with a protruding basal cell; rounded to truncate at the base.

Known distribution (based on molecular data) – India, Goa State (Prabhugaonkar and Bhat 2011), Thailand, Chiang Rai Province (Doilom et al. 2017b), China, Yunnan Province (this study).

Known hosts (based on molecular data) – Cocos nucifera (Prabhugaonkar and Bhat 2011), Tectona grandis (Doilom et al. 2017b), Mangifera indica (this study).

Material examined – China, Yunnan Province, Yuxi Prefecture, on dead wood of Mangifera indica (Anacardiaceae), 29 March 2019, M. Doilom (MFLU 19–2366, new host record and new geographical record), living culture (KUMCC 19–0111).

GenBank numbers – ITS: MN582741, LSU: MN582756, RPB2: MN643158.

Notes – Paradictyoarthrinium species are hard to distinguish based on morphological characters. This is particularly true of conidial size which is not a comparative character due to the large variation among species. Species can however, be differentiated by DNA sequence data (Liu et al. 2018). Phylogenetic analysis of combined LSU, ITS and RPB2 sequence data confirmed that our isolate (KUMCC 19–0111) is *P. diffractum* (Fig. 65). Paradictyoarthrinium diffractum has been reported on a dead twig in a stream from South Africa (holotype), a dead decaying spathe of Cocos nucifera from India, and dead stumps and stems of Tectona grandis in Thailand (Matsushima 1996, Prabhugaonkar and Bhat 2011, Doilom et al. 2017b). This is new host record and new geographical record of *P. diffractum* on Mangifera indica in China.
Fig. 66 – **Paradictyoarthrinium diffractum** MFLU 19-2366, new host record and new geographical record. a Conidia observed on host substrate. b–e Conidia on host substrate. f Conidia arising from hyphae on PDA. g–l Conidia observed on PDA. m Germinated conidium. Scale bars: b–e, h–l = 10 µm, g, m = 20 µm.

**Paradictyoarthrinium hydei** N.G. Liu & J.K. Liu, Phytotaxa 338(3): 290 (2018)

Facesoffungi number: FoF 03933

*Saprobic* on dead inner branch of *Quercus variabilis*. Sexual morph: Undetermined. Asexual morph: *Colonies* on natural substrate, superficial, black, gregarious, powdery. *Mycelium* mostly immersed, composed of pale brown to brown, septate, branched hyphae. *Conidiophores* 3–6 µm diameter, macronematous, rarely micronematous, short or reduced, dark, straight to slightly curved, branched or unbranched, slightly constricted at the septa, uneven, thick-walled. *Conidiogenous cells* holoblastic, monoblastic, integrated, determinate, terminal, dark. *Conidia* 14.5–25 × 12.5–20 µm (\( \bar{x} = 18.5 \times 16 \mu m, n = 15 \)), solitary or developing in chains, muriform, subglobose or irregular, septate, constricted at the septa, dark brown to black, verrucose.

Culture characteristics – Conidia germinating on PDA within 24 hours. Germ tubes produced around conidia. Colonies on PDA edge entire, velvety, dense, effuse, brown to dark grey. *Mycelium* 1.5–2.5 µm wide, partly superficial, pale brown to dark brown, septate. *Conidiophores* up to 70 µm long, 3–5 µm wide, micronematous, straight or flexuous, olive when mounted in water, redish brown when mounted in lactic acid, septate, constricted at the septa, mostly unbranched, rarely branched, arising from laterally and terminally on hyphae. *Conidiogenous cells* monoblastic, terminal. *Conidia* 9.5–24 × 8–20 (\( \bar{x} = 14 \times 13 \mu m, n = 20 \)), initially subglobose, becoming irregular at maturity, muriform, septate, constricted at the septa, olive when mounted in water, reddish brown when mounted in lactic acid, verrucose.
Fig. 67 – *Paradictyoarthrinium hydei* MFLU 19–2365, new host record and new geographical record. a Conidia observed on host substrate. b–d Conidia from host substrate. e Germinated conidium. f Colony on PDA (above and below views). g Conidia arising from hyphae on PDA. h–l Conidia from culture mounted in water. m–o Conidia from culture mounted in 60 % lactic acid. Scale bars: b = 15 µm, c, d, h, j–o = 20 µm, e, i = 50 µm.

Known distribution (based on molecular data) – Thailand, Chiang Mai Province (Liu et al. 2018a), China, Yunnan Province (this study).

Known hosts (based on molecular data) – *Quercus variabilis* (this study)

Material examined – China, Yunnan Province, Kunming, on dead inner branch of *Quercus variabilis* (Fagaceae), 10 June 2019, M. Doilom (MFLU 19–2365), living culture (KUMCC 19–0185).
GenBank numbers – ITS: MN582742, RPB2: MN643159.

Notes – Our isolate (KUMCC 19–0185) is identified as *Paradictyoarthrinium hydei* based on the phylogenetic analysis of combined LSU, ITS and RPB2 sequence data (Fig. 5). *Paradictyoarthrinium hydei* was reported on decaying wood from Thailand (Liu et al. 2018a). This is new host record and new geographical record of *P. hydei* on *Quercus variabilis* in China. The conidia were observed in the PDA culture and showed two different pigmentations when mounted in water (olive, Figs. h–l) and lactic acid (reddish brown, Figs. m–o).

**Phaeosphaeriaceae** M.E. Barr

We follow the latest treatment and updated accounts of Phaeosphaeriaceae in Hyde et al. (2019) and Phookamsak et al. (2019). Eighty-three genera are accepted in this family (Bakhshi et al. 2019, Hyde et al. 2019, Maharachchikumbura et al. 2019, Phookamsak et al. 2019, Yang et al. 2019, Zhang et al. 2019a). We follow the latest treatments and updated accounts of *Leptospora* in Hyde et al. (2016) and Zhang et al. (2019a). Based on molecular analysis, *L. rubella* is reported from *Dipsacus* sp. (Caprifoliaceae) in Italy for the first time.

**Leptospora** Rabenh.

*Leptospora* was introduced by Rabenhorst (1857) and is typified by *L. porphyrogona*. Rabenhorst (1857) established the genus to accommodate two species, previously described as *Sphaeria porphyrogona* and *S. rubella*. Recently, 36 species were accepted in this genus (Index Fungorum 2020) but, phylogeny has been investigated for only four species in Phaeosphaeriaceae (Hyde et al. 2016, Zhang et al. 2019a). We follow the latest treatments and updated accounts of *Leptospora* in Hyde et al. (2016) and Zhang et al. (2019a). Based on molecular analysis, *L. rubella* is reported from *Dipsacus* sp. (Caprifoliaceae) in Italy for the first time.

**Leptospora rubella** (Pers.) Rabenh., Klotzschii Herb. Viv. Mycol., Edn 2: no. 532 (1857)

Facesoffungi number: FoF02442

*Saprobic on Dipsacus* sp. Sexual morph: *Ascomata* 340–395 μm high (excluding neck), 175–275 μm diameter, scattered, solitary, visible as black spots with reddish brown area around ascomata on host surface, semi-immersed, subglobose to ampulliform, uni-loculate, glabrous, dark brown to black, with a long neck. *Neck* 170–200 μm high, 80–100 μm diameter, subcarbonaceous, easily broken, cylindrical, slightly narrower towards the apex, ostiole central, filled with hyaline periphyses, producing red pigment at the apex. *Peridium* 10–30 μm wide, of equal thickness, slightly thin, composed of 3–5 cell layers, of dark brown to black pseudoparenchymatous cells, paler towards the inner layer, arranged in *textura angularis* to *textura prismatica*. *Hamathecium* comprising dense, 2–3 μm wide, hyaline, filamentous, pseudoparaphyses, septate, constricted at the septa, anastomosed above the asci, embedded in a hyaline gelatinous matrix. *Asci* 144–184.5 × 4.5–6 μm (x̄ = 162.2 × 5.2 μm, n = 20), 8-spored, bitunicate, fissitunicate, numerous, cylindrical to subcylindric-clavate, subsessile to short pedicellate, apically rounded, with well-developed ocular chamber, clearly seen at immature state. *Ascospores* 155–182 × 1–3 μm (x̄ = 168.6 × 1.5 μm, n = 10), spiraled or twisted in the ascus, pale brown to yellowish brown, filiform, curved, indistinctly multi-septate, smooth-walled, with guttules, tapering to the lower end cell, lacking a mucilaginous sheath. Asexual morph: Undetermined.

Known distribution (based on molecular data) – Colombia (Crous et al. 2006b), UK (Hyde et al. 2016, Vu et al. 2019), France (Vu et al. 2019), Korea (Eo et al. 2014), Slovenia (Hauptman et al. 2013), Italy (this study).

Known hosts (based on molecular data) – *Eucalyptus* sp. (Myrtaceae), (Crous et al. 2006b), submerged wood in freshwater (Hyde et al. 2016), Ginseng (Eo et al. 2014), *Fraxinus excelsior* (Hauptman et al. 2013), *Dipsacus* sp. (Caprifoliaceae) (this study).
Fig. 68 – Phylogram generated from maximum likelihood analysis based on a combined LSU, SSU, tef1 and ITS sequence dataset. Thirty-one strains are included in the combined gene analyses comprising 3,248 total characters including gaps (LSU: 1–820 bp, SSU: 821–1803 bp, tef1: 1804–2677 bp, ITS: 2678–3248 bp). The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring of the ML tree is selected to represent the phylogenetic relationships of Leptospora with other related genera in Phaeosphaeriaceae, with the final ML optimization likelihood: -11371.252016. The matrix had 614 distinct alignment patterns, with 15.84% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.244665, C = 0.235963, G = 0.262539, T = 0.256833; substitution rates AC = 1.412087, AG = 3.425246, AT = 3.364101, CG = 0.789417, CT = 8.829065, GT = 1.000000; Tree Length = 1.875762; gamma distribution shape parameter α = 0.626011. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.009183. Bootstrap support for maximum likelihood (ML) greater than 70% and Bayesian posterior probabilities (PP) greater than 0.95 are defined above the nodes as ML/PP. The tree is rooted to Ophiobolus artemisiicola (MFLU 15–2140), O. disseminans (MFLUCC 17–1787) and O. rossicus (MFLU 17–1639). The type strain is indicated in bold and newly generated sequence is shown in blue.
Fig. 69 – *Leptospora rubella* (MFLU 17–1084, new host record). a Appearance of ascomata on host. b Section of ascoma. c Ostiolar neck. d, e. Peridium. f Conical ocular chamber. g, i, j Asci (j = fissitunicate). h Septate pseudoparaphysate. k–n ascospores. Scale bars: b = 100 μm, c–e = 50 μm, f =10 μm, g–n = 30 μm.

Material examined – Italy, Province of Forlì-Cesena [FC], Lago Pontini-Bagno di Romagna, on dead land stem of *Dipsacus* sp. (Caprifoliaceae), 7 June 2017, E. Camporesi, IT3376 (MFLU 17–1084, new host record).

GenBank numbers – ITS: MN994333, LSU: MN994335, SSU: MN994336, tef1: MT005778.

Notes – *Leptospora rubella* (Pers.) Rabenh. has been reported from various hosts in European and North American countries as well as India (Shoemaker 1976, Farr & Rossman 2020) and the species can be easily recognized on the host with the red-purple staining of the host substrate.
(Shoemaker 1976, Hyde et al. 2016, Zhang et al 2019a). However, these studies did not include phylogeny based on DNA sequence data. Hyde et al. (2016) designated the reference specimen of *L. rubella* from decaying submerged wood in the UK. Our collection is similar to *L. rubella* in having semi-immersed ascomata with short to long necks, producing red pigment on the host substrate, bitunicate, cylindrical asci and pale brown to yellowish brown, filiform, multi-septate ascospores twisted in the ascus (Crous et al. 2006b, Hyde et al. 2016). In the NCBI BLASTn search of ITS sequences, our strain MFLU 17–1084 matches *L. rubella* (strains CPC 11006 and MFLU 16–0965) with 99.81% similarity. Phylogenetic analyses of a combined LSU, SSU, *tef1* and ITS sequence dataset also indicated that our collection belongs to *L. rubella* (CPC 11006) with high support (100% ML and 1.00 PP, Fig. 68). We hence, identify our collection as *L. rubella*, collected from *Dipsacus* sp. in Italy for the first time.

**Pseudoophiobolus** Phookamsak, Wanas. & K.D. Hyde

*Pseudoophiobolus* was introduced to accommodate ophiobolus-like taxa (Phookamsak et al. 2017). This genus is morphologically and phylogenetically distant from *Ophiobolus sensu stricto*. Currently, there are eight species (Index Fungorum 2020).

**Pseudoophiobolus pseudoitalicus** Senan., & K.D. Hyde sp. nov.

Index Fungorum number: IF556963; Facesoffungi number: FoF06901

**Etymology** – Species epithets based on its morphological similarity to *Pseudoophiobolus italicus*.

**Holotype** – MFLU 14–0840

*Saprobic* on stem of *Galium* sp. Sexual morph: Ascomata 200–300 µm high (excluding papilla), 300–400 µm diameter, dark brown to black, scattered or rarely clustered, immersed or erumpent through host tissue, visible as raised, small, black dots on the host surface, uni-loculate, subglobose to ampulliform, coriaceous, ostiolate, papillate. *Papilla* 100–150 × 60–120 µm, central, rounded to truncate at the apex, ostiole canal periphysate. *Peridium* 25–35 µm wide, composed of dark brown to black, thick-walled, cells of *textura angularis*. *Hamathecium* comprising numerous, 2–3 µm wide, filamentous, broadly cellular pseudoparaphyses, asceptate, guttulate, embedded in a mucilaginous matrix, anastomosing at the apex. *Asci* 150–200 × 9–13 µm (x̅ = 180 × 11 µm, n = 20), 8-spored, bitunicate, cylindrical to cylindric-clavate, short pedicellate, apically rounded with indistinct ocular chamber. *Ascospores* 115–145 × 2–4 µm (x̅ = 135 × 3 µm, n = 20), fasciculate, parallel, hyaline to olivaceous, scolecosporous, filiform, with or without a swollen cell at the centre, multi-septate, not constricted at the septa, smooth, wall-rounded, with small guttules. Asexual morph: Undetermined.

**Material examined** – Italy, Province of Forlì-Cesena, Fiumicello di Premilcuore, dead aerial stem of *Galium* sp., E. Camporesi, 20 March 2014, IT1771 (MFLU 14–0840, holotype ex-type living culture MFLUCC 14–0840).

**GenBank Numbers** – ITS: MN720058, LSU: MN720057.

**Notes** – The blast results of ITS of our *Pseudoophiobolus* strain gives 90.86% similarity with *Pseudoophiobolus rosae* and 91.4% similarity with *P. gali* and blast results of LSU gives 99.55% similarity with *Pseudoophiobolus italicus*. There are 14 unique base pair differences in the ITS locus. However combined ITS, LSU, SSU and *tef1* sequence data analysis in this study (Fig. 70) shows that our *Pseudoophiobolus* strain forms an independent lineage basal to *Pseudoophiobolus italicus* and *P. mathieui* with moderate support. This taxon also exhibits some similar characters to *Pseudoophiobolus italicus* (Phookamsak et al. 2017). However it is differs from *P. italicus* in having globose to subglobose conidiomata and 14–15-septate ascospores swollen at the 6th cell near the basal septum. Therefore, based on morphology and phylogeny, we introduce and describe our strain as a new species, *Pseudoophiobolus pseudoitalicus*. 

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**Pleosporaceae** Nitschke, Verh.

A family of saprotrophic, opportunistic human and plant pathogenic fungi within Pleosporales (Ariyawansa et al. 2015). This family was introduced by Nitschke (1869) and is the largest family in Pleosporales which includes more than 16 genera (Wijayawardene et al. 2017).

**Fig. 70** – Phylogram generated from maximum likelihood analysis based on combined ITS, LSU, SSU and *tef1* sequence data. Bootstrap support for ML greater than 50% and Bayesian posterior probabilities greater than 0.90 are given near nodes. The tree is rooted with *Leptosphaeria doliolum* (CBS 541.66). Ex-type strains are in black bold and the newly generated sequence is indicated in blue bold.

**Bipolaris** Shoemaker

*Bipolaris* was introduced by Shoemaker (1959). Its sexual morph, *Cochliobolus* typified by *C. heterostrophus*, is now linked with the type species of *Bipolaris*, *B. maydis* (Rossman et al. 2013). The genus comprises 132 species.}

Index Fungorum 2020.
**Bipolaris yamadae** (Y. Nisik.) Shoemaker, Can. J. Bot. 37(5): 884 (1959)

Facesoffungi number: FoF 07377

Leaf spots on *Setaria palmiforlia* (Poaceae): Small, indefinite, dark brown, eye spots. Asexual morph: *Conidiophores* 65–170 × 6–8 μm (n = 20), macronematous, mononematous, arising singly or in small groups, simple, septate, sometimes geniculate at upper part, with a swollen basal cell, dark brown. *Conidiogenous cells* mono- or polytretic, terminal or sympodial *Conidia* 65–105 × 15–20 μm (n = 30), usually curved, sometimes straight, elliptical, sometimes obclavate, widest at or just below middle, tapering towards the ends, pale brown to dark brown, distoseptate. Sometimes end cells of conidia swell to form more or less globular vesicles from which germ tubes originate.

**Known distribution** (based on molecular data) – Japan (Manamgoda et al. 2014), Taiwan, Chia Yi (this study).

**Known hosts** (based on molecular data) – *Panicum capillare* (Manamgoda et al. 2014), *Setaria palmiforlia* (Poaceae) (this study)

Material examined – Taiwan, Chia Yi Province, Kwang Hwa, *Setaria palmiforlia* (Poaceae), 27 April 2018, A. Karunarathna AKTW 21 (MFLU 19–2697, new host and geographical record), living culture (NCYUCC 19–0368).

**GenBank numbers** – ITS – MN982855

**Notes** – Herein, we provide a new host and geographical record based on morphology and phylogeny. Our strain is morphologically similar to extant *B. yamadae* and phylogeny reveals a close association with *B. yamadae* CBS 202.29 (Fig. 72).
Fig. 72 – Phylogenetic tree generated from maximum likelihood (ML) based on ITS and TUB sequences. Bootstrap support (BS) values above 50% and Bayesian posterior probabilities equal or greater than 0.90 are shown above the branches at nodes. The tree is rooted with *Alternaria alternate* and *Alternaria infectoria*. The best RaxML tree with a final likelihood value of -5166.414243 is presented. The matrix had 415 distinct alignment patterns, with 12.79% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.244996$, $C = 0.276409$, $G = 0.226712$, $T = 0.251883$; substitution rates $AC = 0.801095$, $AG = 2.393119$, $AT = 0.892562$, $CG = 0.630462$, $CT = 4.085231$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.865731$. Newly generated sequence is indicated in blue bold.
**Curvularia** Boedijn, Bull. Jard.

*Curvularia* was introduced by Boedijn (1933) and species belong to this genus are pathogenic or saprobes on plant or humans (Manamgoda et al. 2014). Species of *Curvularia* are characterized by sympodial conidiophores, intercalary and terminal conidiogenous cell (Berbee et al. 1999, Manamgoda et al. 2014, 2015). Presently, more than 100 species are accepted in this genus (Heidari et al. 2018, Tibpromma et al. 2018b).

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**Fig. 73** – *Bipolaris yamadae* (MFLU 19–2697, new host and geographical record). a Appearance of the host tissue. b Appearance of conidial structures on host. c–f Conidiophores. g–h Conidial attachments. i–m Conidia. n Germinating conidia. Scale bars: c–f = 50 µm, g–n = 20 µm.

**Curvularia hawaiiensis** (Bugnic. ex M.B. Ellis) Manamgoda, L. Cai & K.D. Hyde, Fungal Diversity 56(1): 141 (2012)

Facesoffungi number: FoF 07051

* Saprobic or pathogenic on plant. Sexual morph: see Manamgoda et al. (2015). Asexual morph: Colonies on PDA fast growing, funiculose to cottony, grey, with a fimbriate margin. **Conidiophores** 80–160 × 2–6 µm (x̄ = 120, n = 10), single, flexuous, septate, geniculate, pale to mid brown. **Conidiogenous** nodes dark brown, polytretic, sympodial.
Fig. 74 – Phylogram generated from maximum likelihood analysis based on combined ITS and GPDH sequence data. Ninety-eight strains are included in the combined gene analyses comprising 1083 characters after alignment (544 characters for ITS, 539 characters for GPDH). *Bipolaris*
*ma*dis (CBS 136.29) is used as the outgroup taxon. The newly generated sequences are indicated in blue bold.

Conidia 19–29 × 7.5–10 μm (̄x = 22.5 × 9, n = 30), straight, oblong or cylindrical, rounded at ends, pale to mid brown, 4–7-distoseptate, end cells paler and smooth-walled.

Known distribution – Widely distributed (see Gadallah et al. 1995, Manamgoda et al. 2015, Marin-Felix et al. 2017a, Heidari et al. 2018), Oman (this study).

Known hosts – Wide host range (see Manamgoda et al. 2015), *Saccharum griffithii* (this study).

Material examined – Oman, Muscat Governorate, Wadi Hoqain, on root of *Saccharum griffithii* (Poaceae). 12 March 2017, F Al-AbdulSalam, living culture (SQUCC 13639, new host record and geographical record).

GenBank numbers – ITS: MN758886, GPDH: MN781207.

Notes – The morphology of our collection (SQUCC 13639) is similar to the type of *Curvularia hawaiiensis*. Furthermore, our collection shares a close phylogenetic affinity to *C. hawaiiensis* (BRIP 11987) in the combined ITS and GPDH sequence analyses. This is a new host record and new geographical record of *Curvularia hawaiiensis* on *Saccharum griffithii* in Oman.

Fig. 75 – *Curvularia hawaiiensis* (SQUCC 13639, new host record and geographical record). a–c Conidia and Conidiophores on PDA. d Conidia on PDA. Scale bars: a–b = 20 μm

*Curvularia muehlenbeckiae* Madrid, K.C. Cunha, Gené, Guarro & Crous, Persoonia 33: 56 (2014)

Facesoffungi number: FoF07050

Human pathogens or saprobic or pathogenic on plant. Sexual morph: Undetermined. Asexual morph: Colonies on PDA cottony, pale grey at the centre. *Hyphae* smooth, subhyaline to brown, septate, branched. *Conidiophores* mononematous, macronematous, septate, simple or rarely branched, straight or curved, geniculate near the apex, pale brown, subnodulose and nodulose intercalary swellings 4–10 μm wide. *Conidiogenous* cells integrated, mono- or polytretic, terminal, sympodial, light brown. *Conidia* straight or slightly curved, 3-distoseptate, rounded at both ends; with third cell larger than other cells, distinctly verruculose, 15–25 μm (̄x = 21 μm, n = 30) long, 8–13 μm (̄x = 10.5 μm, n = 30) wide; end cells paler and smooth-walled.

Known distribution – Australia, India, Oman, USA (Madrid et al. 2014), Oman (this study).

Known hosts – *Muehlenbeckia* sp., *Oryza* sp., culture from human chest (Madrid et al. 2014), *Citrus × aurantiifolia* (this study).
Material examined – Oman, Muscat Governorate, Al-Seeb, Al-Sahwa park, on root of *Citrus × aurantiifolia* (Rutaceae), January 2016, HHA Al-Nadabi, living culture (SQUCC 10447, new host record and geographical record).

GenBank numbers – ITS: MN758887, GPDH: MN781208

Notes – *Curvularia muehlenbeckiae* was recently introduced by Madrid et al. (2014) from the leaf of *Muehlenbeckia* sp. collected in India. Our collection from *Citrus × aurantiifolia* has a similar morphology to the holotype of *Curvularia muehlenbeckiae*. Furthermore, the multi-phylogenetic analysis based on the combined sequence data of ITS and GPDH depicts that our strain (SQUCC 10447) clusters with the ex-type of *C. muehlenbeckiae* with moderate bootstrap support (79% ML). Therefore, we confirm our collection as *C. muehlenbeckiae*, and this is the first record of *C. muehlenbeckiae* on *Citrus × aurantiifolia* and from Oman.

**Fig. 76** – *Curvularia muehlenbeckiae* (SQUCC 10447, new host record and geographical record). a Conidia and Conidiophores on PDA. b Conidia on PDA. Scale bars: a–b = 20 µm.

**Curvularia subpapendorfii** (Mouch.) Manamgoda, Rossman & K.D. Hyde, Stud. Mycol. 79: 282 (2014)

Facesoffungi number: FoF07052

*Saprobic* on plant or soil Sexual morph: Undetermined. Asexual morph: Colonies on PDA fast-growing, funiculose, grey, margin fimbriate. *Conidiophores* 160–275 × 4–8 µm (\(\bar{x} = 159 \times 15, n = 15\)), arising singly, simple or branched, flexuous, septate, geniculate at spore bearing part, dark brown, paler towards apex. *Conidiogenous nodes* polytretic, sympodial, dark brown, smooth, with cicatrized pores. *Conidia* 17–25 × 10–14 (\(\bar{x} = 21 \times 12, n = 25\)), smooth-walled, obpyriform, tapering towards rounded ends, pale brown to dark reddish brown, 3-septate. *Hilum* usually inconspicuous or sometimes slightly protuberant.

Known distribution – India (Manamgoda et al. 2015), Oman (this study).

Known hosts – Desert soil, *Phaseolus aconitifolius* (Manamgoda et al. 2015), *Citrus × aurantiifolia* (this study).

Material examined – Oman, Muscat Governorate, Al-Seeb, Al-Sahwa park, on root of *Citrus × aurantiifolia* (Rutaceae), January 2016, HHA Al-Nadabi, living culture (SQUCC 10447, new host record and geographical record).

GenBank numbers – ITS: MN758885, GPDH: MN781206.

Notes – Manamgoda et al. (2014) designated a lectotype for *Curvularia subpapendorfii* for a sample reported from desert soil in Egypt (ex-type culture CBS 656.74). Our collection from the
root of *Citrus × aurantiifolia* (SQUCC 10443) is closely related to the ex-type culture (CBS 656.74) of *C. subpapendorfii*. In addition, conidiophores and conidial size of our specimen are similar to the description of the type material. Thus we identified our collection as *C. subpapendorfii*, and this is the first report of *C. subpapendorfii* on *Citrus × aurantiifolia* in Oman.

![Figure 77](image)

**Fig. 77** – *Curvularia subpapendorfii* (SQUCC 10443, new host record and geographical record). a–c Conidia and Conidiophores on PDA. d Conidia on PDA. Scale bars: a–b = 20 μm.

*Curvularia verruculosa* Tandon & Bilgrami ex M.B. Ellis, Mycol. Pap. 106: 20 (1966)

| Facesoffungi number: FoF 00571 |
|--------------------------------|
| Human pathogens or saprobic or pathogenic on Poaceae. Sexual morph: Undetermined. Asexual morph: Colonies on PDA superficial, effuse, scattered, dark brown. *Hyphae* smooth, subhyaline, septate, branched. *Conidiophores* mononematous, macronematous, septate, simple or rarely branched, straight or curved, geniculate near the apex, pale brown, 7–9-septate, nodose, 115–230 μm (x̄ = 150 μm, n = 15) long, 4–8 μm (x̄ = 6 μm, n = 15) wide. *Conidiogenous* cells terminal or intercalary, polytretic, sympodial, light brown, *Conidia* straight or slightly curved, 3-distoseptate, rounded at both ends; with third cell larger than other cells, distinctly verruculose, 20–32 μm (x̄ = 25 μm, n = 30) long, 10–16 μm (x̄ = 13.5 μm, n = 30) wide, pale to dark brown.

Known distribution – Australia, China, India, Nigeria, Oman, Pakistan, Thailand (Madrid et al. 2014, Manamgoda et al. 2015, Su et al. 2015), Oman (this study).

Known hosts – Various monocotyledons and dicotyledons plants (Manamgoda et al. 2015), grass (this study).

Material examined – Oman, Muscat Governorate, Al-Seeb, Al-Sahwa park, on unidentified dead grass (Poaceae), 12 March, 2017, SSN Maharachchikumbura, living culture (SQUCC 13614, new geographical record).

GenBank numbers – ITS: MN758884, GPDH: MN781205.

Notes – Our collection SQUCC 13614 morphologically resembles the type description of the *Curvularia verruculosa*. Furthermore, phylogenetically our collection forms a strongly supported clade with the epitype CBS 150.63 of *C. verruculosa* (Fig. 74), which was collected from leaves of *Punica granatum* in India. This is the first report of *C. verruculosa* from the Arabian Peninsula.

*Pyrenophora* Fr.

*Pyrenophora* typified by *Pyrenophora phaeocomes* was introduced by Rebentisch (1804). The absence of pseudoparaphyses and smaller ascospore dimensions distinguishes *Pyrenophora* from other genera of Pleosporaceae (Sivanesan 1984, Ariyawansa et al. 2014b). The asexual morph is *Drechslera* (Zhang & Berbee 2001, Ariyawansa et al. 2014b, Marin-Felix et al. 2019). Several *Pyrenophora* species are considered severe pathogens (Tekauz 1983, Lamari & Bernier 1989,
Kingsland 1991, Gupta & Loughman 2001, Ariyawansa et al. 2014b). Owing to its widespread pathogenicity, the development of molecular genetic markers for rapid identification of *Pyrenophora* species have been detailed in Moreno et al. (2011). Most recently Marin-Felix et al. (2019) accepted 27 *Pyrenophora* species with molecular data based on a combined ITS, LSU, GAPDH and RPB2 gene sequence data analysis.

Fig. 78 – *Curvularia verruculosa* (SQUCC 13614, new geographical record( a–b Conidia and Conidiophores on PDA, c Conidia on PDA. Scale bars: a–c = 20 µm.

*Pyrenophora trichostoma* (Fr.) Fr., Jb. nassau. Ver. Naturk. 23–24: 215 (1870) [1869–70]

Fig. 80

FACESOFFUNGI number: FoF06984

*Saprobic* or biotrophic on dead stems and leaves of *Bothriochloa ischaemum*. Sexual morph: *Ascomata* 350–450 µm high 250–300 µm diameter (*n* = 10), black, immersed to semi-immersed or superficial, solitary or in small groups, uniloculate, globose to subglobose, visible as black dome shaped structures on the host surface, conspicuous, surrounded by dark brown setae, glabrous, without papillate, black. *Ostiole* short, rounded, immersed in ascomata, with a pore-like opening. *Peridium* 10–17 µm wide, thin, 3–5-layered of cells of outer dark brown *textura angularis*, and inner pale brown flattened *textura angularis* to *textura prismatica*. *Hamathecium* consisting of irregular-shaped cellular matter, pseudoparaphyses absent. *Asci* 40–102 × 6–15 µm (*x̄* = 70 × 10 µm, *n* = 15), 8-spored, bitunicate, fissitunicate, broadly cylindric-clavate, slightly curved, with a bilobed, short pedicel, apex rounded with a well-distinct ocular chamber, smooth-walled. *Ascospores* 45–85 × 15–35 µm, (*x̄* = 55 × 23 µm, *n* = 40), overlapping biseriate, initially hyaline, becoming pale to golden brown at maturity, muriform, mostly ellipsoidal, with 3 transverse septa, with 1–2 longitudinal septa, constricted at the septa, asymmetrical, end cells slightly lighter, conical and broadly to narrowly rounded at the ends, surrounded by a thick mucilaginous sheath. Asexual morph: Undetermined.

Culture characteristics – Colonies on PDA reaching approximately 2.5 cm diameter after 7 days at 16–18 °C, pinkish-white, dense mycelium, circular, flat, with smooth margins, becoming orangish-red and reverse brown.
Fig. 79 – Phylogram generated from maximum likelihood analysis based on combined ITS and LSU sequence data. Forty-seven strains are included in the combined gene analyses comprising 1779 characters after alignment (868 characters for ITS, 911 characters for LSU). Bipolaris panici-miliacei (CBS 202.29) and Bipolaris yamadae (CBS 199.29) are used as outgroup taxa. The best RaxML tree with a final likelihood value of -7099.358315 is presented. The matrix had 466 distinct alignment patterns, with 13.76 % undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.245308, C = 0.229294, G = 0.279135, T = 0.246263; substitution rates AC = 1.382376, AG = 1.398715, AT = 1.182859, CG = 0.745659, CT = 3.139154, GT = 1.000000; gamma distribution shape parameter α = 0.108634. Bootstrap values for ML and MP equal to or greater than 50 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.
Known distribution (based on molecular data) – USA, Kansas State (Vu et al. 2019), Russia (this study).

Known hosts (based on molecular data) – Bothriochloa ischaemum (this study).

Material examined – Russia, Rostov region, Shakhty City, 20th anniversary of Red Army microdistrict, Solyonaya Balka (Salty gully), on dying stems and leaves of Bothriochloa ischaemum (L.) Keng (Poaceae), 21 May 2015, Timur S. Bulgakov, T-336 (MFLU 15–2040, new host record), living culture KUMCC 16–0120.

GenBank numbers – LSU MN733192, ITS MN736436.

Notes – Pyrenophora species are characterized by immersed to semi-immersed ascomata with necks surrounded by brown to reddish-brown setae, absence of pseudoparaphyses, clavate to saccate asci with large apical rings, and muriform ascospores (Ariyawansa et al. 2014b, Marin-Felix et al. 2019). Fuckel (1870) described P. trichostoma as having 330 μm long, 40 μm wide, 8-spored, asci, with broad-oblong, yellow, asymmetric, muriform ascospores that are 4–6 septate, constricted at the mid septum, 52 × 20 μm (translated from Latin). In the specimen illustrated here, we observed 300 × 54 μm, broadly cylindric-clavate and slightly curved asci with asymmetric, broad, muriform, 55 × 23 μm ascospores. Therefore, we conclude that our species is morphologically similar to P. trichostoma. Pyrenophora trichostoma was once synonymized under P. tritici-repentis based on their morphological similarities (Ciuffetti & Tuori 1999) such as producing ascospores with mucilaginous sheaths (Sivanesan 1984). However, in their phylogenetic analysis, Kodueb et al. (2006) discussed that despite their phylogenetic affinities, there were several base pair differences in the sequences of these two species. In our phylogenetic analysis we have included three strains CBS. 328.53, CBS 391.54 and CBS 392.54 identified as P. trichostoma following Marin-Felix et al. (2019). Our isolate clustered together with these strains with high bootstrap support (99% ML, 100% MP) (Fig. 79).

Hosford (1970) reported a form of P. trichostoma which severely infected wheat in North Dakota in 1968 and 1969. Disease symptoms on leaves included light-brown lesions with yellow halos and the presence of many ascomata on wheat stubble. The pathogenicity of the taxon varied, as well as the resistance of different wheat cultivars towards the pathogen varied, depending on the duration of free moisture on leaves (Hosford 1970). Pyrenophora trichostoma has been reported worldwide on a range of cereal crops, from Bolivia, Brazil, China, Europe (Austria, Italy, England, Poland, Portugal, Spain) to North America (Farr & Rossman 2020), however, no molecular data in GenBank are given in these reports.

Roussoellaceae Jian K. Liu, Phook., D.Q. Dai & K.D. Hyde

Liu et al. (2014) erected the Roussoellaceae to accommodate Neoroussoella, Roussoella and Roussoellopsis. Jaklitsch & Voglmayr (2016b) treated Roussoellaceae as a synonym of Thyridariaceae. Tibpromma et al. (2017), Wanasinghe et al. (2018b), Wijayawardene et al. (2018) and Phookamsak et al. (2019) accepted Roussoellaceae as a stable family.

Pseudoneoconiothyrium Wanas., Phukhams., Camoispersi & K.D. Hyde

Wanasisinghe et al. (2018b) introduced Pseudoneoconiothyrium Wanas et al. to accommodate Roussoella-like taxa in Thyridariaceae (Phookamsak et al. 2019). However, in multi gene (LSU, SSU, tef1, ITS and RPB2) analysis conducted by (Phookamsak et al. 2019) Pseudoneoconiothyrium clustered within Roussoellaceae.

Pseudoneoconiothyrium rosae (Phukhams., Camoispersi & K.D. Hyde) Phukhams., Camoipersi & K.D. Hyde, Index Fungorum 357: 1 (2018) Fig. 82

Facesoffungi number: FoF 04055

Biotrophic on living branch of Lonicera sp. Sexual morph: Undetermined. Asexual morph: Conidiomata 200–320 μm diameter × 280–350 μm μm high (̄x = 265.74 × 341.3 μm, n = 5), pycnidial, semi-immersed, erumpent from the substrate, solitary-scattered, globose to subglobose, dark brown to black, unilocular, ostiolate. Ostiole central, single, apapillate, filled with dark brown
to light brown cells. **Pycnidial wall** 20–35 μm thick, 5–7 layered, composed of cells of *textura angularis*, cells in outer layer brown-walled, cells in inner layer light brown to hyaline. **Conidiogenous cells** ampulliform, enteroblastic, anellidic, with distinct percurrent proliferations, integrated, discrete, smooth, and hyaline. **Conidia** 8–12 × 6–9 μm (\( \bar{x} = 10.41 \times 7.98 \) μm, n = 35), initially hyaline, smooth and thin-walled, light brown to dark brown and rough-walled when mature, globose to ellipsoid, aseptate, guttulate.

Culture characteristics – Conidia germinating on WA within two days. Colonies on PDA reaching 5–10 mm diameter after one week at 16 °C, entire edge, flat or effuse, greyish white, with dense, flat mycelium on the surface, dark green in reverse.

**Fig. 80** – *Pyrenophora trichostoma* (MFLU 15–2040, new host record and geographical record). a Appearance of ascomata on host. b Close up of ascoma. c Section of ascoma. d Setae. e Peridium. f Hypha-like matter in hamathecium. g Ocular chambers in asci. h–k Immature to mature asci. l–o Ascospores. p Sheath surrounding the ascospores. Scale bars: a = 500 μm, b = 200 μm, c = 100 μm, d, f, g = 20 μm, e, h–m = 10 μm.
Fig. 81 – Phylogram generated from maximum likelihood analysis (RAxML) of Roussellaceae based on ITS, LSU, SSU, tef1 and RPB2 sequence data. The tree is rooted to Torula herbarum (CBS 140066, CBS 111855) and Torula hollanica (CBS 220.69). The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The final optimization likelihood value is -22735.444576 of the best RaxML tree with 1227 distinct alignment patterns and 37.75 % undetermined characters or gaps. Estimated base frequencies are as
follows: A = 0.245625, C = 0.250937, G = 0.271866, T = 0.231572; substitution rates AC = 1.253105, AG = 3.802122, AT = 1.885811, CG = 0.990899, CT = 8.801862, GT = 1.000000; gamma distribution shape parameter α = 0.158691. Maximum likelihood bootstrap values equal or above 60% and the Bayesian posterior probabilities equal or above 0.90 (ML/PP) are given at the nodes. The ex-type strains are indicated in black-bold and the newly generated strains are in blue-bold.

Fig. 82 – *Pseudoneoconiothyrium rosae* (MFLU 18–0930, new host record). a Conidiomata on living host surface of *Lonicera* sp. b Enlarged conidiomata on host. c Longitudinal section of conidioma. d Longitudinal section of ostiole. e Longitudinal section of a pycnidial wall. f–h Conidiogenous cells with developing conidia. i–m Conidia. n Germinating conidium. o–p Colonies on PDA (o upper, o lower). Scale bars: a = 200 μm, b–c = 100 μm, d, n = 50 μm, e–f, h = 20 μm, g = 10 μm, i–m = 5 μm.
Known distribution (based on molecular data) – Italy, Province of Forlì-Cesena (Wanasinghe 2018b, Phookamsak et al. 2019, this study).

Known hosts (based on molecular data) – dead aerial spines of *Rosa canina* (Rosaceae) (Wanasinghe 2018b, Phookamsak et al. 2019), lives on a live stem of *Lonicera* sp. (Caprifoliaceae) (this study).

Material examined – Italy, Province of Forlì-Cesena, Rocca delle Caminate - Predappio, on a live stem of *Lonicera* sp. (Caprifoliaceae), 4 April 2018, E. Camporesi (MFLU 18–0930, new host record), culture (MFLUCC 18–1353).

GenBank numbers – ITS: MN783333, LSU: MN783330, RPB2: MN814846, SSU: MN783332.

Notes – In the multi-gene phylogeny (LSU, ITS, SSU and RPB2) of our study, our novel strain (MFLUCC 18–1353) and the ex-type strain of *Pseudoneoconiothyrium rosae* (MFLUCC 15–0052) clustered with relatively high support (100% ML/1.00 PP) (Fig. 81). These isolates were collected from different hosts, but in the same locality. The characters of our species overlap with the holotype (Wanasinghe et al. 2018b, Phookamsak et al. 2019). *Neoconiothyrium rosae* (MFLU 18–0117, holotype) was synonymized under *Pseudoneoconiothyrium rosae* (Phookamsak et al. 2019). Both species have many similar characters with solitary-scattered and erumpent, globose to subglobose, dark brown to black, unilocular ascomata and single, central, ostioles. The pycnidial wall is composed of cells of *textura angularis* and cells in the outer layer are brown-walled, cells in inner layer light brown to hyaline. Conidia are also globose to ellipsoid shape aspate and guttulate. However, the size of ascomata, conidia and the number of pycnidial wall cell layers of our new collection is comparatively larger than the ex-type strain. According to the guidelines of Jeewon and Hyde (2016), we have analyzed nucleotide differences within the rRNA gene region for further clarifications. When comparing the ITS region (ITS1–5.8S–ITS2) from 436 nucleotides and LSU rDNA region from 799 bp, there are zero bp (0 %) differences between MFLUCC 18–1353 and MFLUCC 15–00520 strains. Considering the morpho-molecular data analysis, we conclude that our new collection is another record of *Pseudoneoconiothyrium rosae* and a new host record on *Lonicera* sp. (Caprifoliaceae) in Italy.

*Pseudoneoconiothyrium rosae* (Phukhams., Camporesi & K.D. Hyde) Phukhams., Camporesi & K.D. Hyde, Index Fungorum 357: 1 (2018)  
Facesoffungi number: FoF 04055

*Saprobic* on dead and hanging branch of *Galium* sp. Sexual morph: Undetermined. Asexual morph: *Conidiomata* 300–320 μm diameter × 240–360 μm high (̄ = 311.6 × 295 μm, n = 5), pycnidial, semi immersed, erumpent from the substrate, solitary-scattered, globose to subglobose, dark brown to black, unilocular, ostiolate. *Ostiole* single, circular. *Pycnidial wall* 30–50 μm thick, 5–8 layered, composed of cells of *textura angularis* and cells in outer layer brown walled, cells in inner layer light brown to hyaline. *Conidiogenous cells* ampulliform, enteroblastic, annellidic with distinct percurrent proliferations, sometimes cylindrical, elongate neck, integrated, discrete, smooth and hyaline. *Conidia* 7–10 × 4–8 μm (̄ = 8.45 × 6.22 μm, n = 20), initially hyaline, smooth and thin-walled, light brown to dark brown and rough-walled when mature, globose to ellipsoid, aspate, guttulate.

Culture characteristics – Conidia germinating on MEA within three days. Colonies on MEA reaching 5-10 mm diameter after one week at 16 °C, undulate, white, with dense, flat mycelium on the surface, pale yellow in reverse.

Known distribution (based on molecular data) – Italy, Province of Forlì-Cesena (Wanasinghe et al. 2018b, Phookamsak et al. 2019, this study).

Known hosts (based on molecular data) – dead aerial spines of *Rosa canina* (Rosaceae) (Wanasinghe et al. 2018b, Phookamsak et al. 2019), lives on a dead hanging stem of *Galium* sp. (Rubiaceae) (this study).
Material examined – Italy, Province of Forlì-Cesena, Rocca delle Caminate – Predappio, on a dead hanging stem of *Galium* sp. (Rubiaceae), 4 April 2018, E. Camporesi (MFLU 18–0927, new host record), culture (MFLUCC19-0494). GenBank numbers – ITS: MN783331, LSU, MN783329, *tef1*: MN794051.

![Image](image.png)

**Fig. 83** – *Pseudoneoconiothyrium rosae* (MFLU 18–0927, new host record). a Conidiomata on dead branch of *Galium* sp. b–c Enlarged conidiomata on host surface. d Longitudinal section of a conidioma with ostiole. e Longitudinal section of ostiole. f Longitudinal section of a conidioma wall. g–k Conidiogenous cells with developing conidia; l–p Conidia. q Germinating conidium. r–s Colonies on PDA (r upper, s lower). Scale bars: a, e, l = 100 μm, q = 50 μm, j–k = 30 μm, f–i = 20 μm, g–h =10 μm, m–p = 5 μm.
In the multi-gene phylogeny (LSU, ITS, SSU, RPB2 and tef1) of our study, our novel strain (MFLUCC 19–0494) and the ex-type strain of *Pseudoneoconiothyrium rosae* (MFLUCC 15–0052) clustered with relatively high support (100% ML/1.00 PP) (Fig. 81). These collections are from different hosts, but in the same locality. When morphological characters of our species were examined, they overlap with the holotype from dead spines of *Rosa canina* (Rosaceae) in Italy which was described by Wanasinghe et al. (2018b). *Neoconiothyrium rosae* (MFLU 18–0117, holotype) was synonymized under *Pseudoneoconiothyrium rosae* (Phookamsak et al. 2019). Both species have similar morphological characters with solitary-scattered, globose to subglobose, dark brown to black, uniloculate and erumpent ascomata, with a single, central, ostiole. Conidia are also globose to ellipsoid, aseptate and guttulate. However, the size of ascomata and thickness of pycnidial wall of our new collection are comparatively larger than the ex-type strain. Also, some conidiogenous cells have cylindrical and elongate neck. According to the guidelines of Jeewon & Hyde (2016) we have analyzed nucleotide differences within the rRNA gene region for further clarifications. In comparison of ITS regions (ITS1-5.8S-ITS2) from 436 nucleotides and LSU rDNA region from 799 bp, there are zero bp (0%) differences between MFLUCC 18–1353 and MFLUCC 15–00520 strains. Considering the morpho-molecular data analysis, we conclude that our new collection is a record of *Pseudoneoconiothyrium rosae* and also a new host record on *Galium* sp. (Rubiaceae) in Italy.

**Roussoella** Sacc.

*Roussoella* was introduced with the type species *R. nitidula* Sacc. & Paol. The ambiguous placement of *Roussoella* was later on resolved by Liu et al. (2014). The latest treatment for *Roussoella* was by Phookamsak et al. (2019).

**Roussoella siamensis** Phook., Phook., Jian K. Liu & K.D. Hyde, Phytotaxa 181(1): 18 (2014)

Facesoffungi number: FoF01984

*Saprobic on Phragmites australis*. Asexual morph: Coelomycetous. *Conidiomata* 140–150 μm high × 220–225 μm diameter (\( \bar{x} = 145 \times 223 \) μm, n = 5), pycnidial, immersed in host epidermis, raised, hemisphaerical to subconical, coriaceous, scattered or clustered, solitary or gregarious, uniloculate, glabrous, centrally ostiolate, with minutely pore-like opening. *Pycnidial wall* 6–8 μm wide, composed of several layers of brown to dark brown pseudoparenchymatous cells of *textura angularis*, flattened at the base, thick at side towards the apex, outer layer fused with the host tissue. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* 8–13.5 × 1.5–3 μm (\( \bar{x} = 11 \) μm × 2 μm, n = 20), enteroblastic, phialidic, unbranched, cylindrical to cylindric-clavate, or ampulliform, hyaline, aseptate, thin-walled, smooth-walled, arising from the basal cavity around the conidiomata. *Conidia* 3–4 × 1.5–2.5 μm (\( \bar{x} = 3.7 \times 2 \) μm, n = 40), oblong to ellipsoidal, hyaline, aseptate with two guttules, smooth-walled.

Culture characteristics – Conidia germinated on PDA within 12h at 20–25 °C, with germ tubes produced from a one end. Colonies on PDA moderate growing, dense, irregular in shape, flattened to raised, slightly rough-walled with lobate edge, and uneven margin, velvety, colony from above dark green to dark brown, from below black; sometimes white to pale pinkish, radiated with yellowish-brown to dark brown brown at the center and white at the margin, not producing pigmentation on agar medium.

Known distribution (based on molecular data) – Thailand, Chiang Rai Province (Liu et al. 2014, China, Dali (this study).

Known hosts (based on molecular data) – *Bambusa* (Liu et al. 2014), *Phragmites australis* (this study)

Material examined – China, Kunming, Lake Dianchi, on decaying stems of *Phragmites australis* (Poaceae), 5 November 2016, K.D. Hyde, AKD 04 (MFLU 17–0351, new host and geographic record); ex-type living culture, MFLUCC 17–1352; KUMCC 16–0207.

GenBank number – LSU: MN989183.

**Notes** –
Fig. 84 – Phylogenetic tree generated from maximum likelihood (ML) based on LSU, SSU, ITS, tef1 and RPB2 sequences. Bootstrap support (BS) values above 50% and Bayesian posterior probabilities equal or greater than 0.90 are shown above the branches at nodes. The tree is rooted with Neoroussoella bambusae and N. alishanense. The best RaxML tree with a final likelihood value of -13900.126681 is presented. The matrix had 823 distinct alignment patterns, with 44.82% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.244559, C = 0.251803, G = 0.271754, T = 0.231883; substitution rates AC = 1.179055, AG = 4.071030, AT = 1.977458, CG = 0.917006, CT = 10.347006, GT = 1.000000; gamma distribution shape parameter α = 0.763580. The newly generated sequence is indicated in blue.

Notes – This is the first record of the asexual morph of Roussoella siamensis. The species is supported by phylogeny. Herein, we provide the new host and geographic record for the Roussoella siamensis. In our phylogenetic analysis, the strain MFLUCC 17–1352 clusters with MFLU 11-0149 (Fig. 84).

Sporormiaceae Munk

Sporormiaceae is classified in Pleosporales, in Dothideomycetes and was established by Munk (1957) and is typified by Sporormia (De Notaris 1844), with the type species Sporormia fimetaria (Phukhamsakda et al. 2016). This type genus is characterized by ostiolate, globose
pseudothecia and spores joined in bundles by a common gelatinous sheath (Ahmed & Cain 1972, Dissing 1992). Even though absent in the type species, germ slits are also considered as a common character in this family (Kruys et al. 2006). Sporormiaceae is the largest coprophilous ascomycetes family and most members are cosmopolitan on a vast variety of animal dung types and have also been reported on other substrates like decaying plant debris, decaying textiles, rotten vegetation, soil and wood, or as endophytes and saprobes in plants (Kruys et al. 2006, Sue et al. 2014). Sporormiaceae comprises approximately 100 species classified in ten genera, including the recently described genus, *Sporormurispora* by Wanasinghe et al. (2018b) and *Chaetopreussia, Forliomyces, Pleophragmia, Preussia, Sparticola, Sporormia, Sporormiella, Spororminula,* and *Westerdykella* (Karunarathna et al. 2017, Phukhamsakda et al. 2016).

**Fig. 85** – *Roussoella siamensis* (MFLU 17–0351, new host and geographic record). a Appearance of the host. b–c Colonies on host substrate. d Longitudinal section of conidioma. e Longitudinal section of ostiole. f Peridium. g–h Conidiogenous cells and developing conidia. i–m Conidia. n Germinated conidium. o–p Culture characteristics. Scale bars: d = 50 µm, e = 10 µm, f = 10 µm, g–h = 10 µm, i–n = 5 µm.
**Forliomyces** Phukhams., Camporesi & K.D. Hyde

*Forliomyces* is a monotypic genus in Sporormiaceae introduced by Phukhamsakda et al. (2016) with *Forliomyces uniseptatus* as the type species. *Forliomyces* is characterized by immersed, pycnidial, subglobose, unilocular conidiomata with oblong to subobovoid, slightly curved, septate, conidia with abscission scars (Phukhamsakda et al. 2016). Sexual connections are still unclear for *Forliomyces*.

**Forliomyces uniseptatus** Phukhams., Camporesi & K.D. Hyde, Cryptog. Mycol. 37(1): 84 (2016)

Fig. 87

Facesoffungi number: FoF07381

Saprobic on dead aerial branches of *Hippocrepis emerus*. Conidiomata visible as round dark brown to black dots on the host surface. Sexual morph Undetermined. Asexual morph Coelomycetous. Conidiomata 120–155 μm high, 160–250μm diameter, pycnidial and solitary, scattered, globose, semi-immersed to immersed, unilocular, thin walled. Conidioma wall composed of 3–4 layers, similarly dense 15–21 μm wide, outer and inner layers light brown to brown, with thin walled cells of textura angularis. Conidiophores reduced to conidiogenous cells. Conidiogenous cells formed from the cells lining the inner wall of conidiomata 2–8 μm long × 1–3 μm wide, hyaline, enteroblastic, phialidic, determinate, discrete, and oblong to clavate. Conidia 8–13 × 4–7 μm wide (x̄ = 10 × 5 μm, n = 20), oblong, sub obovoid and some are ellipsoidal to cylindrical, initially hyaline to light brown, brown to dark brown at maturity, slightly bent at the middle, septate, thick, smooth-walled, guttulate.

Culture characteristics – Colonies on PDA reaching 15mm diameter after 2 weeks at 25 °C, circular colony, concentrically zonate, flat, smooth, with pinkish to orange diffusible pigments into agar. Colonies from above light creamy orange at the margin and pinkish red with whitish in middle, reverse colony color deep red in center, pinkish to red in middle and creamy orange at the edge.

Known distribution (based on molecular data) – Italy (Phukhamsakda et al. 2016) and (this study).

Known hosts (based on molecular data) – On *Salvia* (Phukhamsakda et al. 2016), on dead aerial branch of *Hippocrepis emerus* (this study)

Material examined – Italy, Province of Forlì-Cesena [FC], Galeata, Strada San Zeno, on dead aerial branch of *Hippocrepis emerus* (Fabaceae), 25 April 2017, E. Camporesi, IT 3331 (JZBH 3390001, new host record), living culture: JZB: 3390001.

GenBank numbers – LSU: MK066913, SSU: MK066909, ITS: MK066911.

Notes – The collection obtained from dead aerial branches of *Hippocrepis emerus* was identified as *Forliomyces uniseptatus* with support from morphology and phylogeny. Our isolate clustered with the type species, *Forliomyces uniseptatus* (MFLUCC 15-0765), in the combined LSU, SSU, ITS, tef1 and RPB2 sequence phylogeny (Fig. 86). In addition, our isolate showed 100%, 98% and 100% base pair similarity with *Forliomyces uniseptatus* (MFLUCC 15-0765) in LSU, ITS and tef1 gene regions. This is the first record of *Forliomyces uniseptatus* species reported from *Hippocrepis emerus* from Italy.

**Tetraplosphaeriaceae** Kaz. Tanaka & K. Hiray.

Tetraplosphaeriaceae was introduced by Tanaka et al. (2009) and typified by *Tetraploa* with *T. aristata* as the type species (Berkeley & Broome 1850, Hyde et al. 2013). Recently, seven genera were accommodated in this family viz. *Ernakulamia*, *Polylosphaeria*, *Pseudotetraploa*, *Quadricrura*, *Shrungabeeja*, *Tetraploa* and *Triplosphaeria* (Berkeley & Broome 1850, Rao & Reddy 1981, Subramanian 1994, Tanaka et al. 2009, Hyde et al. 2013, Wijayawardene et al. 2018).
Fig. 86 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU, ITS, tef1 and RPB2 sequence data. Twenty-four strains are included in the combined gene analyses and *Melanomma pulvis-pyrius* (CBS 124080) is used as the out-group taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -13932.036268 is presented. The matrix had 1020 distinct alignment patterns, with 45.20% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.246751, C = 0.248031, G = 0.276957, T = 0.228261; substitution rates AC = 1.342154, AG = 2.568856, AT = 1.620900, CG = 1.189329, CT = 7.343609, GT = 1.000000; gamma distribution shape parameter $\alpha = 0.669698$. Bootstrap values for maximum likelihood equal to or greater than 60 and Bayesian posterior probabilities equal or greater than 0.90 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.
**Polyplosphaeria** Kaz. Tanaka & K. Hiray.

*Polyplosphaeria* was introduced by Tanaka et al. (2009), with *P. fusca* as the type. Until now, there are five species accommodated in the genus viz. *P. fusca*, *P. nabanheensis*, *P. pandanicola*, *P. thailandica*, *P. xishuangbannaensis*, however, only *P. fusca* has been reported as a holomorph, while in the other four species only the asexual morph is known (Tanaka et al. 2009, Li et al. 2016a, Tibpromma et al. 2018b). The sexual morph of the genus is characterized by globose, black to sometimes reddish-brown ascomata, 8-spored, fissitunicate, clavate asci with 1(−3)-septate, hyaline to pale olive-brown, fusiform ascospores (Tanaka et al. 2009). The asexual morph produces globose to subglobose, brown to dark brown conidia with setose appendages (Tanaka et al. 2009, Li et al. 2016a, Tibpromma et al. 2018b).

**Polyplosphaeria fusca** Kaz. Tanaka & K. Hiray., Studies in Mycology 64: 193 (2009)  

Facesoffungi number: FoF 06806

*Saprobic* on stem of dead bamboo branches. Sexual morph: *Ascomata* 200–300 μm high, 300–500 μm diameter, immersed under the host, solitary, scattered, 2–3-gregarious, dark brown, conical to subglobose, with ostiole vertically lined with hyaline periphyses, uni-loculate, mostly associated with reddish pigments. *Peridium* 20–30 μm wide, comprising several layers, with dark brown to brown parenchymatous cells arranged in *textura angularis*. *Hamathecium* of hyaline, septate, branched, 1–1.7 μm broad pseudoparaphyses, anastomosing at the top. *Asci* 115–140 × 23–32 μm ( *= 127.5 × 27.5 μm, n = 20*), 8-spored, fissitunicate, broadly cylindric-clavate, pedicellate with slightly furcate ends, apically rounded. *Ascospores* 46–58 × 9–10 μm ( *= 52 × 9.5 μm, n = 20*), 1–3-seriate, hyaline to pale olive-brown, fusiform, slightly curved, 1-septate, constricted at the septum, smooth-walled, with guttules. Asexual morph: Refer to Tanaka et al. (2009).

Culture characteristics – *Ascospores* germinating on PDA within 24 h. Colonies on PDA reaching 17–20 mm diameter after 10 days at room temperature (20–30 °C); from above, raised, convex, brown to dark brown at the centre, white at the margin, circular, velvety; from reverse, yellowish.

Known distribution (based on molecular data) – Japan, Aomori, Nagasaki, Shizuoka, Tochigi (Tanaka et al. 2009), China, Yunnan (this study).

Known hosts (based on molecular data) – *Chimonobambusa marmorea* (Gramineae), *Phyllostachys bambusoides* (Gramineae), *Pleioblastus chino* (Gramineae), *Sasa kurilensis* (Gramineae) (Tanaka et al. 2009), bamboo (Tanaka et al. 2009; this study).

Material examined – China, Yunnan Province, Honghe Prefecture, Pingbian County, nearby Tuanpo Reservoir, on stem of dead bamboo (Poaceae), 21 September 2017, H.B. Jiang, Pb002 (KUN-HKAS 101757, new geographic record), living culture KUMCC 18–0110.

GenBank numbers – ITS: MN629458, LSU: MN629359, SSU: MN629360.

Notes – The new collection is similar to *Polyplosphaeria fusca* and based on multi-gene phylogenetic analyses, it clustered with *P. fusca* (KT1616 and KT2124), but separated from *P. fusca* (KT1043 and KT1640) (Fig. 88). These two groups can be distinguished based on the characteristics, such as ascomata in these specimens are “almost superficial without associated pigmentation” and “immersed with reddish pigments”, respectively, as reported by Tanaka et al. (2009). A comparison of the ITS, LSU and SSU nucleotides of the type strain of *P. fusca* (KT1616) and the new strain revealed ≤ 1.5% nucleotide differences which confirm our new strain as *P. fusca* (Jeewon & Hyde 2016). *Polyplosphaeria fusca* was reported from Japan (Tanaka et al. 2009); our collection is a new geographic record for China.

**Tetraploa** Berk. & Broome

*Tetraploa* was given preference over *Tetraplosphaeria* as discussed by Hyde et al. (2013) and Wijayawardene et al. (2014). Among the 20 epithets in Index Fungorum (2020), molecular data is available for only for seven species. *Tetraploa* species are mostly found on monocotyledons (Tanaka et al. 2009, Farr & Rossman 2020).
Fig. 87 – *Forliomyces uniseptatus* (JZB 3390001, new host record). a Appearance of conidiomata on the host. b Close up view of conidiomata. c Section through the conidioma. d Conidioma wall. e Immature conidia attached to conidiogenous cell. f Mature conidia. g–h Colony on PDA after 2 weeks (g = from above, h = from below). i Pigmented mycelium. j–k Clamydospores formation from hyphae. l Conidiomata in culture. m Conidiogenous cell. n Conidia. Scale bars: a, b = 200 μm, e, f, k = 20 μm, n = 10 μm.

*Tetraploa sasicola* (Kaz. Tanaka & K. Hiray.) Kaz. Tanaka & K. Hiray.  
Facesoffungi number: FoF 01984

*Saprobic* on decaying culms of *Phragmites australis*. *Mycelia* dark brown to black, effuse, branched, septate, partly superficial. Asexual morph: *Conidiogenous cells* micronematous, dark brown, monoblastic, integrated, short, usually undistinguishable from superficial hyphae. *Conidia* short cylindrical, conidial body four columnar, coarsely verruculose, 35–45 × 25–30 μm, with four
apical setose appendages, 90–120 µm, partly split conidia, where the two columns of the upper part of the main conidial body is totally split along with the two apical appendage but connected at the base.

**Fig. 88** – Phylogram generated from maximum likelihood analysis based on a combined ITS, LSU and SSU sequence dataset. Thirty strains are included in the combined gene analyses comprising 3,219 total characters including gaps (ITS: 1–579 bp, LSU: 580–1877 bp, SSU: 1878–3219 bp). The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring of the ML tree is selected to represent the phylogenetic relationships of *Polyplosphaeria* species, with the final ML optimization likelihood: -8728.061069. The matrix had 434 distinct alignment patterns, with 16.08% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.251610, C = 0.234539, G = 0.277180, T = 0.236670; substitution rates AC = 4.456817, AG = 4.305638, AT = 2.601496, CG = 1.500369, CT = 14.507453, GT = 1.000000; Tree-Length = 0.450936; gamma distribution shape parameter $\alpha = 0.497321$; The proportion of invariable sites $I = 0.712404$. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.009756. Bootstrap support for maximum likelihood (MLBP) greater than 70% and Bayesian posterior probabilities (BYPP) greater than 0.95 are defined above the nodes as MLBP/BYPP. The tree is rooted to *Lophiotrema neoarundinaria* (KT 856). The type strain is indicated in bold and newly generated sequence is shown in blue.
Fig. 89 – Polyplosphaeria fusca (HKAS 101757, new geographic record). a–c Ascomata on bamboo host. d Ascoma from lateral view. e Ostiole. f Periphyses. g Peridium. h Pseudoparaphyses. i–l Asci. m Culture characteristics. n–p Ascospores. q Germinating ascospore. Scale bars: e, q = 30 μm, g–l, n–p = 20 μm, f = 10 μm.

Culture characteristics – Conidia germinated on PDA within 12h at 20–25 °C, with germ tubes produced from a one end. Colonies on PDA moderate growing, dense, irregular in shape, flattened to raised, rough with entire edge, velutinous to floccose, greenish grey in the center and to dull green towards margin, mycelia white, thin, regular. Colony reverse grey to brownish grey. Soluble pigments and exudates are absent.

Known distribution (based on molecular data) – JAPAN, Hokkaido, Yoichi, Sawamachi (Tanaka et al. 2009), China, Kunming (this study).
Known hosts (based on molecular data) – *Sasa senanensis* (Tanaka et al. 2009), *Phragmites australis* (this study).

Material examined – China, Kunming, Lake Dianchi, on decaying stems of *Phragmites australis* (Poaceae), 28 Nov 2016, A. Karunarathna, AKKIB 57 (MFLU 17–0377, new host and geographical record); living culture, MFLUCC 17–1387; KUMCC 16–0240.

GenBank number – LSU: MN989185.

Notes – Here we report our collection from a new host and geographical record for *Tetraploa sasicola* based on phylogeny and taxonomy (Fig. 90).

**Fig. 90** – Phylogenetic tree generated from maximum likelihood (ML) based on LSU, ITS and RPB2 sequences. Bootstrap support (BS) values above 50% and Bayesian posterior probabilities equal or greater than 0.90 are shown above the branches at nodes. The tree is rooted with *Pseudotetraploa javanica* and *P. curviappendiculata*. The best RaxML tree with a final likelihood value of -6537.602092 is presented. The matrix had 461 distinct alignment patterns, with 24.41% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.238681, C = 0.271326, G = 0.277534, T = 0.212459; substitution rates AC = 3.351614, AG = 2.810733, AT = 2.048449, CG = 1.295686, CT = 8.198345, GT = 1.000000; gamma distribution shape parameter α = 0.563647. Newly generated sequence is in blue-bold.

**Torulaceae** Corda

Torulaceae was established by Sturm (1829) based on the type genus *Torula* Pers. Sexual morphs are undetermined in this family. Currently, six genera are accepted in Torulaceae (Li et al. 2016c, Su et al. 2016b, Su et al. 2018, Hyde et al. 2020).

**Dendryphion** Wallr.

*Dendryphion* was introduced by Wallroth (1833), typified by *D. comosum*. Seventy seven
epithets and eight varieties are listed in Index Fungorum (2020), but sequence data are available for only few species.

Fig. 91 – *Tetraploa sasicola* (MFLU 17–0377, new host and geographical record). a Appearance of the host. b Colonies on host substrate. c Attachment of conidia to the conidiophore. d–g Conidia. h Germinated conidium. i–j Culture characteristics. Scale bars: c = 50 µm d–h = 50 µm.

*Fig. 93 – Dendryphion nanum* (Nees) S. Hughes, Canadian Journal of Botany 36 (6): 761 (1958) Facesoffungi number: FoF06714

*Saprobic* on dead stems of *Sambucus ebulus*. Sexual morph: Undetermined. Asexual morph: Hyphomycetous. *Colonies* on natural substrate effuse, dark brown or black, velvety. *Mycelium* immersed, composed of hyaline, septate, branched hyphae. *Conidiophores* mononematous, erect, straight to broadly curved, septate, thick-walled, smooth, dark brown, branched in the upper half. *Conidiogenous cells* polytretic, terminal and intercalary, integrated, discrete, cylindrical, with raised brown large scars appear when the conidia detached. *Conidia* 40–70 µm long, 8–14 µm in the broadest part, solitary or in chains, dry, obclavate always tapered somewhat at the ends, rounded or with a scar at the apex; truncate with a well-defined dark brown scar at the base, septate, slightly constricted at the septa, median brown when immature, dark brown except for the end cells which are median brown, verrucose.

Culture characteristics – Conidia germinating on water agar within 24 h. Germ tubes produced from one or both ends. Mycelia superficial, circular, sparse, decumbent, with entire edge, white from above and below.

Known distribution (based on molecular data) – Germany (Farr et al. 2000), Canada (Corredor 2011), Sweden (Grudzinska-Sterno et al. 2016) China, Yunnan Province (Su et al. 2016b), Su et al. 2018), Italy (this study).

Known hosts (based on molecular data) – Soil (Farr et al. 2000), Salix sp. (Corredor 2011), Winter wheat (Grudzinska-Sterno et al. 2016), Submerged decaying wood (Su et al. 2016b, Su et
al. 2018). *Sambucus ebulus* (this study).

Material examined – Italy, on a dead stem of *Sambucus ebulus*, E. Camporesi, I3077 (GZAAS 20–0004, new geographicol record), living culture, GZCC 20–0003.

GenBank numbers – ITS: MN999921; LSU: MN999926.

Notes – *Dendryphion nanum* was introduced by Hughes (1958). These species are common on dead stems of herbaceous plants, also on roots and cut tree stumps (Ellis 1971). Su et al. (2016b) reported *D. nanum* on submerged decaying wood from Yunnan Province, China and designated the reference specimen (HKAS 84012). Our collection has slightly smaller conidia (40–70 μm long) compared with those of reference specimen (56.5–74.5 μm long). Molecular data confirmed that our collection is *D. nanum* (Fig. 92). This is a new geographical record of *D. nanum* in Italy.

**Torula** Pers.

*Torula* is an asexual genus, which is characterized by dark brown to black, effuse, powdery colonies, with terminal or lateral, monoblastic or polyblastic conidiogenous cells. The conidiogenous cells are thickened, with heavily melanized walls at the base, becoming coronate at the apex, cupulate, brown, smooth to verruculose, with the terminal cell of conidia often becoming a conidiogenous cell. Conidia are acrogenous, in short to long branched chains, pigmented, phragmosporous to scolecosporous, moniliform, smooth to verrucose (Su et al. 2016b, Li et al. 2017b, 2020, Hyde et al. 2019, 2020). Currently, more than 500 epithets are listed in Index Fungorum (2020), although, only 16 species have molecular data to confirm their phylogenetic affinities in Torulaceae (Crous et al. 2015a, Su et al. 2016b, 2018, Li et al. 2017b, Hyde et al. 2017, 2019, 2020, Tibpromma et al. 2017). We followed the latest treatments and updated accounts of *Torula* in Hyde et al. (2019, 2020). In this study, *Torula chromolaenae* is reported from dead branch of a herbaceous plant in Yunnan, China for the first time.

**Torula chromolaenae** J.F. Li, Phookamsak, A. Mapook & K.D. Hyde, Mycol Progress 16:447–461 (2017) Fig. 94

Facesoffungi number: FoF 02713

*Saprobic* on herbaceous litter. Sexual morph: Undetermined. Asexual morph: Hyphomycetous, visible as black, powdery, dense to effuse colony on the natural substrate. *Mycelium* 1.8–3.5 μm wide, immersed to superficial, composed of septate, branched, smooth, pale brown hyphae, becoming black closer to fertile region. *Conidiophores* (2–)5–8(–13) × 2–5 μm (x̄ = 7.2 × 3.7 μm, n = 30) macronematous, mononematous, erect, solitary, brown, oblong to cylindrical, smooth, straight or slightly flexuous, 0–1-septate, without apical branches, occasionally reduced to conidiogenous cells. *Conidiogenous cells* 4–8(–11) × 4–6.5 μm (x̄ = 5.5 × 5.5 μm, n = 30), polyblastic, integrated, terminal, brown to dark brown, cupulate to subglobose, with terminal cell of conidia become conidiogenous cell, smooth, thick-walled, doliiform. *Conidia* (9–)10–16(–22) × 5–7 μm (x̄ = 13 × 6.1 μm, n = 50), catenate, in branch chains, dry, acrogenous, brown to dark brown, oblong to cylindrical, 2–3-septate, constricted at septa, rounded at apex, with truncate base, smooth to minutely verruculose.

Known distribution (based on molecular data) – Thailand (Li et al. 2017b), Yunnan, China (Tibpromma et al. 2018b, this study).

Known hosts (based on molecular data) – *Chromolaena odorata* (Asteraceae; Li et al. 2017b), on fallen dead and decaying leaves of *Pandanus tectorius* (Tibpromma et al. 2018b), on herbaceous litter (this study).

Material examined – China, Yunnan Province, Xishaungbanna, Jinghong, the Nabanhe National Nature Reserve, on herbaceous litter, 25 November 2015, R. Phookamsak, XB012 (MFLU 20–0140, new host record).
Fig. 92 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, SSU, tef1 and RPB2 sequence data. Forty-nine strains are included in the combined gene analyses comprising 4003 characters after alignment (841 characters for LSU, 592 characters for ITS, 934 characters for SSU, 787 characters for tef1, 849 characters for RPB2). Neooccultibambusa chiangraiensis (MFLUCC 12–0559) and Occultibambusa bambusae (MFLUCC 13–0855) are used as the outgroup taxa. The tree topologies derived from the Bayesian analysis and maximum parsimony analysis were similar to that derived from the maximum likelihood analysis. The best RAxML tree with a final likelihood value of −18261.964665 is presented. The matrix had 1151 distinct alignment patterns, with 33.41 % undetermined characters or gaps. Estimated base frequencies were as follows: \( A = 0.245872, \ C = 0.261687, \ G = 0.270980, \ T = 0.221461; \)
substitution rates $AC = 1.850933$, $AG = 4.358262$, $AT = 1.744260$, $CG = 1.264921$, $CT = 9.195943$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.137863$. Bootstrap values for maximum likelihood equal to or greater than 75 and Bayesian posterior probabilities equal or greater than 0.95 are placed near the nodes. Ex-type or ex-epitype strains are in bold and black. The newly generated sequences are indicated in bold and blue.

**Fig. 93** – *Dendryphion nanum* (GZAAS 20–0004, new geographical record). a, b Colonies on substrate. c, d Conidiophores. e–g Conidiogenous cells. h–n Conidia. o Geminated conidium. Scale bar: a = 500 μm, b = 200 μm, c = 20 μm, d = 50 μm, e–g = 10 μm, h–m = 15 μm.
**Fig. 94** – *Torula chromolaenae* (MFLU 20–0140, new host record). a Appearance of colonies on host substrate. b–e Colonies on host substrate showing partly immersed mycelium and its erect conidiophores attached with conidia. f–h Conidiophores and conidiogenous cells. i–n Conidia. Scale bars: b = 50 μm, c–f, i = 20 μm, g, h, j–n = 10 μm.

GenBank numbers – ITS: MT007586, LSU: MT007599, SSU: MT007598.

Notes – In NCBI BLASTn search based on ITS sequences, our strain XB012 is most similar to *Torula chromolaenae* strain NBHC3-4 (99.18% similarity) and the type strain KUMCC 16-0036 (99.05% similarity). Multi-gene phylogenetic analyses showed that our strain is sister to the type strain of *T. chromolaenae* with high support (99% ML, 1.00 BYPP; Fig. 92). A comparison of ITS and *tef1* also showed that our strain is identical to *T. chromolaenae*. Our collection shares the size range of the conidiophores, conidiogenous cells and conidia with the type of *T. chromolaenae* (Li et al. 2017b). Furthermore, our collection resembles the type of *T. chromolaenae* in having brown, oblong to cylindrical, 2–3-septate, smooth to minutely verruculose conidia, although, our collection has a truncate base to the conidia and lacks a paler apex (Li et al. 2017b). Therefore, we identify
our collection as *T. chromolaenae, Torula chromolaenae* was introduced by Li et al. (2017b) from *Chromolaena odorata* (Asteraceae) in Thailand. Tibpromma et al. (2018b) reported the species occurring on decaying leaves of *Pandanus tectorius* in Yunnan, China. Our collection is also collected from Yunnan, China. However, it was found on herbaceous litter.

**Asterinales** M.E. Barr ex D. Hawksw. & O.E. Erikss.

**Asterinaceae** Hansf.

Phylogenetic studies have found that phylogenetic placement of Asterinaceae is unresolved with taxa occurring in two separate lineages (Zeng et al. 2019).

**Asterostomella** Speg.

*Asterostomella* is the asexual morph of Asterinaceae characterized by brown, ovoid to oblong conidia, sometimes with a hyaline band in the center (Hongsanan et al. 2014). Species distinction in the genus has largely been based on fruiting body diameter and conidia measurements (Reynolds & Gilbert 2005). We provide the first sequence data for this genus herein and phylogenetically confirm its placement as a member of Asterinaceae.

**Asterostomella grewiae** Petr., Sydowia 12: 485 (1959) [1958]  
Facesoffungi number: FoF06899

*Parasitic* on living leaves. Sexual morph: Undetermined. Asexual morph: Colonies epiphyllous, densely scattered, confluent. *Hyphae* straight to substraight, loosely reticulate. *Hyphopodia* 8–11 × 6–9 μm (x̄ = 9 × 7 μm, n = 5), 2-celled, mostly lobed. *Pycnothyria* 71–93 μm (x̄ = 80 μm, n = 20) in diameter, superficial, dark brown, seated on a thin, hyaline basal stroma with a central X- or Y-shaped dehiscence. *Upper wall* comprising radial arranged square cells, with meandrous marginal cells. *Pycnothyriospores* 12–15 × 10–11 μm (x̄ = 14 × 10.5 μm, n = 30), aseptate, ovoid to pyriform, 2-layered.

![Phylogram generated from maximum likelihood analysis based on LSU sequence data.](image)

Thirty-nine strains are included in the analyses comprising 832 characters after alignment. *Cladoriella rubrigena* (CBS 124760) and *C. eucalypti* (CPC 10953) are used as out group taxa. The best RAxML tree with a final likelihood value of - 6491.621315 is presented. The matrix had 453 distinct alignment patterns, with 11 % undetermined characters or gaps. Estimated base frequencies
were as follows: $A = 0.231783$, $C = 0.252162$, $G = 0.326987$, $T = 0.189068$; substitution rates $AC = 0.895393$, $AG = 2.984753$, $AT = 0.711520$, $CG = 1.133511$, $CT = 9.597818$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.587263$. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above the branches. The newly generated sequence is indicated in blue.

Fig. 96 – Asterostomella grewiae (MFLU 13–0629, geographical record). a Host leaves. b Colonies on leaf surface. c, d Ascomata. e Basal layer of the pycnothyrium wall. f Hyphae with hyphopodia. g–i Conidia. Scale bars: b = 100 μm, c, d = 50 μm, e, f = 20 μm, g–i = 10 μm.

Known distribution (based on molecular data) – Philippines (Petrak 1983), Thailand (this study).

Known hosts (based on molecular data) – Grewia multiflora (Petrak 1983), Malvaceae (this study).
Material examined – Thailand, Chiang Mai, Mae Taeng, Pa Pae, Bahn Pa Deng, 128 Moo 3, Mushroom Research Centre, on the living leaves of Malvaceae, 22 November 2013, X.Y. Zeng (MFLU13-0629, new sequence data and geographical record, reference specimen designated here).

GenBank numbers – LSU: MN364645; SSU: MN364416.

Notes – Asterostomella species reported from Malvaceae include A. diplocarpa, A. grewiae, A. helicteris and A. isotheca. Our new collection is similar to Asterostomella grewiae in the size of pycnothyria (71–93 µm vs 40–90 µm in diameter), hyphopodia (8–11 × 6–9 µm vs 5–8 × 6–11 µm) and pycnothriospores (12–15 × 10–11 µm vs 10–17 × 6–10 µm), as well as 4-lobed hyphopodia and the host leaves. Therefore, we designate a reference specimen herein with illustrations and sequence data (Fig. 95).

**Botryosphaeriales**

**Aplosporellaceae** Slippers, Boissin & Crous

Aplosporellaceae was introduced by Slippers et al. (2013) and is typified by *Aplosporella* with *A. chlorostroma* as the type species. Recently, three genera were accommodated in this family viz. *Alanomyces*, *Aplosporella* and *Bagniella* (Dissanayake et al. 2016, Sharma et al. 2017, Wijayawardene et al. 2018).

**Aplosporella** Speg.

*Aplosporella* was introduced by Spegazzini (1880) and is characterized by uni- to multi-loculate, pycnidial, stromatic conidiomata, with a communal ostiole, and aseptate, ellipsoid to subcylindrical conidia, brown to dark brown and spinulose when mature (Hyde et al. 2013, Slippers et al. 2013, Ekanayaka et al. 2016, Sharma et al. 2017). We follow the latest treatments and updated accounts of *Aplosporella* in Fan et al. (2015b), Dissanayake et al. (2016), Ekanayaka et al. (2016), Du et al. (2017), Dou et al. (2017a), Jia et al. (2019) and Phillips et al. (2019). *Aplosporella javeedii* is reported from *Acer palmatum* in Yunnan, China for the first time.

**Aplosporella javeedii** Jami, Gryzenh., Slippers & M.J. Wingf., Fungal Biology 118(2): 174 (2013) Fig. 98

Facesoffungi number: FoF07383

*Saprobic* on dead hanging twigs of *Acer palmatum*. Sexual morph: Undetermined. Asexual morph: *Conidiomata* 210–290 µm high, 270–680 µm diameter, pycnidial, rounded or valsoïd, stromatic, immersed to erumpent through host epidermis, becoming raised, superficial, dark brown to black, subglobose to dome-shaped, discoid or irregular in shape, 1–3-loculate, ostiolate, with a single ostiole throughout stroma. *Locules* quadrilateral to subglobose, or irregular in shape, 2–3 subdivided chamber separated by invaginations with common walls, the developing invaginations frequently present. *Pycnidial walls* 25–60 µm wide, thick-walled, composed of two type cells, outer layers comprising several cell layers, of large, broad, thick-walled, dark brown to black cells, arranged in a *textura angularis* to *textura globulosa*, inner layers comprising 1–3 layers of small, flattened, light brown pseudoparenchymatous cells paler towards the inner region, arranged in a *textura prismatica* to *textura angularis*; interstitial walls of a locule composed of several cell layers of flattened, light brown to brown pseudoparenchymatous cells, arranged in *textura prismatica* or palisade-like. *Paraphyses* 28–111 µm long, 1.4–3.2 µm (x̄ = 68.4 × 2.1 µm, n = 40) wide, hyaline, indistinctly septate, thin-walled, filiform, with rounded tip, filamentous, associated with blastic conidiogenesis. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* (5.5–)15–28(–31.5) × (2.5–)3–5.7(–7.7) µm (x̄ = 19.7 × 4.1 µm, n = 30), raising from the inner cavity, holoblastic, monophialidic, hyaline, smooth, short cylindrical to flask-shaped. *Conidia* 15–27 × 7–12 µm (x̄ = 21.3 × 10 µm, n = 100), initial hyaline, becoming brown to dark brown when mature, ellipsoidal to oblong with rounded ends, or subcylindrical, aseptate, reticulate.

Culture characteristics – Colonies on PDA fast growing, reaching 90 mm diameter, after 1 week at 20–25 °C, colonies medium sparse to dense, with aerial mycelia, circular, flat, surface slightly rough with edge entire, margin well defined, cottony to fairly fluffy, with standing tufts;
colony from above and below, white to cream at the beginning, becoming black after 4 weeks; not producing pigments on PDA.

Known distribution (based on molecular data) – China, Beijing, Gansu, Henan, Heilongjiang, Shanxi, Yunnan (Fan et al. 2015b, Dou et al. 2017a, Jia et al. 2019), South Africa, Gauteng (Jami et al. 2013).

Known hosts (based on molecular data) – Acer buergerianum (Aceraceae), Albizia julibrissin (Fabaceae), Broussonetia papyrifera (Moraceae), Celtis africana (Cannabaceae), Chaenomeles sinensis (Rosaceae), Gleditsia sinensis (Fabaceae), Juglans regia (Juglandaceae), Juniperus chinensis (Cupressaceae), Morus alba (Moraceae), Searsia lancea (Anacardiaceae), Styphnolobium japonicum (Fabaceae), Ziziphus jujuba (Rhamnaceae) (Jami et al. 2013, Fan et al. 2015b, Dou et al. 2017a, Jia et al. 2019).

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**Fig. 97** – Phylogram generated from maximum likelihood analysis based on combined LSU-ITS and tef1 sequence data. Thirty strains are included in the combined gene analyses comprising 1631 characters after alignment (844 characters for LSU, 621 characters for ITS, 164 characters for tef1). Melanops tulasnei (CBS 116805) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RAxML tree with a final likelihood value of -4527.083852 is presented. The matrix had 342 distinct alignment patterns, with 22.06 % undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.235117, C = 0.239252, G = 0.286504, T = 0.239127; substitution rates AC = 2.896203, AG = 3.311309, AT = 1.557883, CG = 2.969131, CT = 7.107161, GT = 1.000000; gamma distribution shape parameter α = 0.699794. Bootstrap values for maximum parsimony and maximum likelihood equal to or greater than 50% and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold. The newly generated sequences are indicated in blue.
Fig. 98 – *Aplosporella javeedii* (KUN-HKAS 102211, new host and geographical record). a, b Appearance of conidiostromata on host substrate. c, d Section through conidiostroma. e, f Section through pycnidial wall. g Paraphyses. h–j Conidiogenous cells embedded in paraphyses. k–q Variation in shape of conidia. r Conidia showing reticulate, rough-walled. Scale bars: a = 1000 µm, b = 500 µm, c, d = 100 µm, e–g = 50 µm, h–r = 10 µm.

Material examined – China, Yunnan Province, Kunming, Panlong, Ciba, on dead hanging twigs of *Acer palmatum* (Aceraceae), 28 November 2015, R. Phookamsak, COE008, KUN-HKAS 102211, new host and geographical record, living culture, KUMCC 18–0134.

GenBank numbers – ITS: MH974687, *tef1*: MT005776.

Notes – An NCBI BLASTn search of ITS sequence data shows that our strain (KUMCC 18-0134) is identical to *Aplosporella javeedii* with 100% similarity. Phylogenetic analyses of concatenated ITS and *tef1* sequence dataset also indicated that our taxon has a close affinity with *A. javeedii* with strong support (Fig. 97, 100% ML and 1.00 PP). Furthermore, our collection (KUN-HKAS 102211) shares a size range of conidia and conidial characters with the type and other representative specimens (Jami et al. 2013, Fan et al. 2015b). Therefore, we identify our collection as *A. javeedii*. This species is collected from dead hanging twigs of *Acer palmatum* in Yunnan, China for the first time.
**Aplosporella prunicola** Damm & Crous, Fungal Diversity 27(1): 39 (2007)

Facesoffungi number: FoF 04955

*Saprobic* on a dead branch of *Prunus serrulata* (Rosaceae). Sexual morph: Undetermined.

Asexual morph: *Conidiomata* 650–800 × 350–420 μm, pycnidial, solitary, dark brown, immersed to semi immersed, erumpent, multi-loculate, central ostiole. *Conidioma wall* 55–90 μm wide at the side, 35–46 μm wide at the base, composed of 6–12 layers of brown, thick-walled cells, arrange in a *textura angularis*. *Paraphyses* 26.5–60 μm (x̅ = 40.2 μm, n = 10) long, 4–6.3 μm (x̅ = 4.5 μm, n = 10) wide at base, 1.6–2.7 μm (x̅ = 2 μm, n = 10) wide at upper part, hyaline, smooth-walled, septate, blunt ended. *Conidiophores* 7.4–13.2 × 4.7–6.7 μm (x̅ = 10.5 × 5.6 μm, n = 15), reduced to conidiogenous cells, monophialidic, hyaline, cylindrical to doliform, smooth-walled. *Conidia* 17.5–21 × 9.7–12.2 μm (x̅ = 19.5 × 10.9 μm, n = 30), L/W 1.8, aseptate, initially hyaline, smooth-walled, broadly ellipsoid to subcylindrical, with rounded ends, becoming dark brown.

Culture characteristics – Conidia germinating on PDA within 16 hr. Colonies on PDA, reaching 41.5 mm diameter after one week at 25–30 °C, circular, flat or effuse, dense, upper surface initially greenish, becoming blackish-green from the center within 7 days. Reverse dark green to black.

Known distribution (based on molecular data) – South Africa, Limpopo, China, Henan, Yunnan (Damm et al. 2007, Dou et al. 2017a).

Known hosts (based on molecular data) – *Acacia cochlearia*, *Cercis chinensis f. chinensis*, *Ficus sp.*, *Prunus persica var. nucipersica* (Damm et al. 2007, Dou et al. 2017a).

Material examined – China, Guizhou Academy of Agricultural Sciences (GZAAS), Guiyang, Guizhou, on a dead branch of *Prunus serrulata* (Rosaceae), 12 June 2018, MC. Samarakoon, SAMC156 (MFLU 18–0774, HKAS 102387, new host record), living culture MFLUCC 19–0490.

GenBank numbers – ITS: MN473059, LSU: MN473053, SSU: MN473047, TUB: MN987001.

Notes – Our strain (MFLUCC 19–0490) collected on *Prunus serrulata* is similar to *Aplosporella prunicola* (Damm et al. 2007), which was collected from the bark of a small dead tree of *P. persica var. nucipersica*. BLASTn searches of ITS and LSU gave results of 100 % similarity of bp comparisons with CBS 121167 and STE U 6326 strains of *Aplosporella prunicola*. The combined LSU, ITS, tef1 phylogenetic analyses show the clustering of MFLUCC 19–0490 in the *A. prunicola* clade. Based on similar morphology and molecular data, here, we report our strain as a new host record from *Prunus serrulata*.

**Botryosphaeriaceae** Theiss. & Syd.

The most recent treatments of Botryosphaeriaceae are by Phillips et al. (2019) and Burgess et al. (2019), and species are widespread in tropical and temperate regions, and are forest pathogens, while some may be saprobes or endophytes. Currently, 28 genera are accepted in Botryosphaeriaceae (Wijayawardene et al. 2018).

**Barriopsis** A.J.L. Phillips, A. Alves & Crous

**Barriopsis** was introduced by Phillips et al. (2008) with *B. fusca* as the type species. The peculiar taxon is characterized by brown, aseptate ascospores without apiculi (Phillips et al. 2008). Currently, *Barriopsis archontophoenics*; *B. iraniana*, *B. tectonae*, *B. stevensiana* and *B. thailandica* are accepted (Abdollahzadeh et al. 2009, Doilom et al. 2014, Konta et al. 2016, Tiberomma et al. 2017, Wijayawardene et al. 2017). In this study, we illustrate and describe *B. stevensiana* on *Cassia* sp. from Thailand, a new geographical record.

**Barriopsis stevensiana** A.J.L. Phillips & Pennycook, Fungal Diversity, 86: 56 (2017)

Facesoffungi number: FoF07061

*Saprobic* on dead twigs of *Cassia* sp. Sexual morph: *Ascostromata* individual locule 150–280 μm high × 215–300 μm diameter (ascostromata with papilla), black, convex on host tissue, appearing through cracks in bark, solitary, initially immersed, becoming erumpent, when cut...
horizontally locules visible as white contents and dark ascospore dots, uni to multi-loculate, globose to subglobose or flask-shaped, with central papilla, ostiole with periphyses. Peridium composed of two layers, outer layer composed of black to reddish brown, thick-walled cells of textura angularis, inner layer composed of hyaline, thin-walled cells of textura angularis. Hamathecium comprising 3–5.5 μm wide, hyaline, hyphae-like, numerous, septate, pseudoparaphyses, constricted at the septa. Asci 100–180 × 28–35 μm (x̄ = 123 × 32 μm, n = 15), 8-spored, bitunicate, fissitunicate, cylindro-clavate or clavate, with a short or long pedicel, apically rounded with an ocular chamber. Ascospores 21–34 × 11.5–15 μm (x̄ = 33 × 14 μm, n = 30), uni-seriate at the base, 2–3-seriate at the centre and end, hyaline when immature, gradually pale brown, becoming brown when mature, ellipsoidal, aseptate, straight or slightly curved, widest in the middle, thick-walled, smooth. Asexual morph: Undetermined.

Fig. 99 – Aplosporella prunicola (MFLU 18–0774, new host record). a Host. b–d Conidiomata on the substrate. e Vertical section of conidioma. f Peridium. g Paraphyses. h, i Conidiogenous cells. j–o Conidia. p Upper view of the colony. q Reverse view of the colony. Scale bars: b = 1 cm, c = 1000 μm, d, e = 200 μm, f = 50 μm, g–o = 10 μm.
**Fig. 100** – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS and tef1 sequence data. Sixteen strains are included in the combined gene analyses comprising 1770 characters after alignment (866 characters for LSU, 539 characters for ITS, 365 characters for tef1). *Diplodia mutila* (strains CBS 230.30 and CBS 112553) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -4240.541158 is presented. The matrix had 264 distinct alignment patterns, with 18.91% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.229339, C = 0.254463, G = 0.291126, T = 0.225072; substitution rates AC = 1.116498, AG = 2.893632, AT = 0.794870, CG = 1.586169, CT = 5.750768, GT = 1.000000; gamma distribution shape parameter \( \alpha = 0.140521 \). Bootstrap values for maximum likelihood equal to or greater than 70 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.

Known distribution (based on molecular data) – Cuba (Phillips et al. 2008)

Known hosts (based on molecular data) – *Citrus* sp. (Phillips et al. 2008), *Cassia* sp. (this study).

Material examined – Thailand, Chiang Rai Province, Muang District, Tha Sut Subdistrict, on dead twigs of *Cassia* sp. (Fabaceae), 30 September 2011, M. Doilom (MFLU 19–1560), living culture (MFLUCC 11–0420).

GenBank numbers – ITS: MN582740, LSU: MN582755, SSU: MN582689.

Notes – Our specimen is similar to the holotype of *Barriopsis stevensiana* (= *Barriopsis fusca*) which was re-examined by Liu et al. (2012b) and Phillips et al. (2013). Based on phylogenetic analysis of combined LSU, ITS and tef1 sequence data our strain (MFLUCC 11–0420) clusters with the ex-type strain of *B. stevensiana* (CBS 174.26) with high support (Fig. 100, 100% ML/ 0.98 PP). The holotype was collected on twigs of *Citrus* in Cuba (Stevens 1926, Phillips et al. 2008, Liu et al. 2012b). This is new geographical record of *B. stevensiana* on *Cassia* sp. in Thailand.
Fig. 101 – *Barriopsis stevensiana* MFLU 19–1560, new geographical record. a Ascomata on dead twigs of *Cassia* sp. b Ascomata cut through horizontally showing the white contents with dark spots. c, d Sections through ascomata. e Peridium. f–k Asci. l–o Ascospores. p Germinated ascospore. Scale bars: c, d = 200 µm, e, g, h, j, k = 20 µm, f, i = 50 µm, l–o = 10 µm, p = 100 µm.
**Botryosphaeria** Ces. & De Not.

*Botryosphaeria* is typified with *Botryosphaeria dothidea*. This genus has undergone various revisions and updates over the years and the most recent taxonomic treatment is Jayawardena et al. (2019a). In this study, we collected *Botryosphaeria dothidea* from *Torilis arvensis*.

**Botryosphaeria dothidea** (Moug.) Ces. & De Not., Comm. Soc. crittog. Ital. 1(fasc. 4): 212 (1863)

Fig. 103

Facesoffungi number: FoF03512

Saprotrophic on dead and aerial branches of *Torilis arvensis*. Sexual morph: Undetermined. Asexual morph: Colonies on natural substrate stromatic, *Conidiophores* hyaline, cylindrical, smooth. *Conidiogenous cells* phialidic, hyaline, subcylindrical. *Conidia* 15–10 × 7–5 μm (μ = 13 × 6 μm n = 20), hyaline, 1-celled, narrowly fusiform, with a subtruncate to bluntly rounded base, forming a septum before germination, smooth-walled with a granular content.

Culture characteristics – Colonies on PDA reaching 50 mm diameter after 4 d at 28 °C. Mycelial mat moderately dense, margin smooth, olivaceous, becoming grey.

Known distribution (based on molecular data) – worldwide (Phillips et al. 2013, Manawasinghe et al. 2018), Italy (this study).

Known hosts (based on molecular data) – wide range of hosts from monocotyledons to dicotyledons (Phillips et al. 2013, Dissanayake et al. 2017), *Torilis arvensis* (this study).

Material examined – Italy, Province of Forlì-Cesena, Teodorano – Meldola, on dead branch of *Torilis arvensis* (Apiaceae), 30 January 2018, E. Camporesi IT 3699 (MFLU 18-0177, new host record), living culture, JZB310196.

GenBank number – ITS: MN533806.

Notes – *Botryosphaeria dothidea* is the type species. In this study, we isolated a *Botryosphaeria* strain from Italy associated with branches of *Torilis arvensis*. In the phylogenetic analysis, this strain clustered with the *Botryosphaeria dothidea* isolates with <50% MI bootstrap and <0.95 BYPP values (Fig. 102). However, the characters, such as conidial shape, size and colour are similar to the type (Phillips et al. 2005, Phillips et al. 2013). Therefore, we identified our current taxon as *Botryosphaeria dothidea*. This species has a worldwide distribution on many economical important trees (Phillips et al. 2005, Phillips et al. 2013, Dissanayake et al. 2017). This is the first report of *B. dothidea* on *Torilis arvensis* (Farr & Rossman 2020).

**Diplodia**

*Diplodia* was introduced by Montagne and is typified by *D. mutila*. *Diplodia* has a worldwide distribution and are known to be pathogens, endophytes and saprobes on a wide range of woody hosts (Crous et al. 2006a, Slippers & Wingfield 2007, Phillips et al. 2008, Phillips et al. 2013, Ariyawansa et al. 2015). The genus comprises 30 species with molecular data (Hyde et al. 2018). Two types of distinct conidial morphologies are observed in the genus. The first type of conidia is initially hyaline, aseptate and later becomes pale to dark brown and 1-septate. In this type, delayed pigmentation is observed and in some species dark conidia are never observed. The second type of conidia becomes pigmented at an early stage of development where they are still enclosed inside the pycnidia and rarely become septate (Phillips et al. 2013).

**Diplodia galiicola** Dissanayake, Camporesi & K.D. Hyde, Fungal Diversity, 75: 54 (2015)

Fig. 105

Facesoffungi number: FoF00884

Saprobic on dead aerial stem of *Knautia* sp. Sexual morph: Undetermined. Asexual morph: *Conidiomata* 220–225 μm high × 372–380 μm diameter (μ = 219 × 376 μm, n=10), stromatic, solitary, semi immersed in host epithelium, dark brown to black, uniloculate or biloculate, globose to subglobose. *Peridium* 36–54μm wide (μ = 44 μm, n=10), outer layer composed of dark brown cells of *textura angularis*, inner layers of thin-walled hyaline cells of *textura angularis*. *Conidiogenous cells* 7–12 μm high × 3–7 μm wide (μ = 10 × 4 μm, n=15), holoblastic, integrated.
hyaline, thin-walled, smooth, cylindrical, swollen at the base, producing a single conidium at the apex. *Conidia* 18–26 × 8–18 μm (x̄ = 21×11 μm, n = 30), initially hyaline, turning dark brown with maturity, moderately thick-walled, aseptate, ovoid, widest in the center, obtuse apex, verruculose.

**Fig. 102** – Phylogram generated from maximum likelihood analysis based on combined ITS, *tef1*, and TUB sequence data. Twenty-seven strains are included in the combined gene analyses comprising characters after alignment. *Macrophoma phaseolina* (strains CBS 162.25 and CBS 227.33) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of –3083.503236 is presented. The matrix had 224 distinct alignment patterns,
with 14.27% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.210590, C = 0.307570, G = 0.251651, T = 0.230189; substitution rates AC = 0.677624, AG = 3.002265, AT = 1.150518, CG = 0.720959, CT = 5.263152, GT = 1.000000; gamma distribution shape parameter \( \alpha = 1.009656 \). Bootstrap values for maximum likelihood greater than 50 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue bold.

**Fig. 103** – *Botryosphaeria dothidea* (MFLU 18-0177, new host record) a Material examined. b–c Appearance of conidia on host surface. d Conidiogenous cell attached to conidia. e Conidiogenous cell. f Conidia. g Upper view of colony on PDA. h Reverse view of colony on PDA. Scale bars: a–c = 500 µm, d–f = 100 µm.

Known distribution (based on molecular data) – Italy (Giambra et al. 2016, Linaldeddu et al. 2016), Italy, Forlì-Cesena Province, Strada San Zeno Province, Galeata Province (Dissanayake et al. 2016).

Known hosts (based on molecular data) – *Galium* sp. (Rubiaceae) (Ariyawansa et al. 2015), *Knautia* sp. (Caprifoliaceae) (this study).

Material examined – Italy, Province of Forlì-Cesena [FC], Civitella di Romagna, Buggiana, on dead aerial stem of *Knautia* sp. (Dipsacaceae), 12 February 2018, E. Camporesi, IT 3712 (JZB 3140014, new host record).

GenBank submissions – ITS: MN757871, tef1: MN854663.

Notes – Based on the phylogenetic analysis of combined ITS and *tef1* sequence data of *Diplodia* species (Fig. 104), our strain (JZB 3140014) was identified as *D. galiicola*. *Diplodia galiicola* is phylogenetically closely related with *D. seriata* (Fig. 104). The difference is observed in conidial length of *D. galiicola* being shorter as compared to *D. seriata*. The average conidial length of *D. seriata* is greater than or equal to 25 µm (Phillips et al. 2013), while the conidial length of *D. galiicola* is less than 25 µm. For our strain (JZB 3140014), the culture characteristics could not be obtained since the spores did not germinate in culture (PDA) and DNA was directly extracted carefully from fruiting bodies. Base pair comparison of ITS and *tef1* gene regions between our strain (JZB 3140014) and reference strain of *D. galiicola* (MFLU 15–1310) reveal less than 1% difference. The morphological characters such as conidiogenous cells and conidial dimensions also overlap confirming that the two strains are the same species (Ariyawansa et al. 2015). *Diplodia galiicola* previously has only been isolated as a saprobe on *Galium* sp. from Italy. This is the first time *D. galiicola* has been collected from *Knautia* sp.
Fig. 104 – Phylogram generated from maximum likelihood analysis based on combined ITS, and tef1 sequence data. Forty seven strains are included in the combined gene analyses comprising 866 characters after alignment (578 characters for ITS, 288 characters for tef1). Lasiodiplodia theobromae CBS 164.96 is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -3349.531698 is presented. The matrix had 261 distinct alignment patterns, with 12.08 % undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.203138, C = 0.302938, G = 0.263561, T = 0.230363; substitution rates AC = 0.989254, AG = 4.136603, AT = 0.730217, CG = 1.833881, CT = 4.680274, GT = 1.000000; gamma distribution shape parameter α = 0.775702. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.
**Fig. 105** – *Diplodia galiicola* (IZB 3140014, new host record). a, b Conidiomata on host substrate. c Cross section of conidioma. d Peridium. e, f Mature and immature conidia attached to conidiogenous cells. g Immature conidia attached to conidiogenous cells. h, i Mature conidia. Scale bars: c = 100 μm, d–g = 20 μm, h, i = 10 μm.

**Dothiorella** Sacc.

We follow the latest treatment and updated accounts of *Dothiorella* in Yang et al. (2017), Hyde et al. (2019) and Phookamsak et al. (2019). Yang et al. (2017) treated *Spencermartinsia* as a synonym of *Dothiorella* based on molecular phylogeny and the morphological character of apiculate ascospores which were not reliable to distinguish *Spencermartinsia* from *Dothiorella* (Yang et al. 2017, Hyde et al 2019). Phookamsak et al. (2019) mentioned that *Spencermartinsia alpina* and *S. yunnana* had to be transferred to *Dothiorella* in further studies. In this study, we collected *Spencermartinsia alpina* from a different host in Yunnan Province, China. Based on molecular data and phylogenetic analyses, we thus, transfer *S. alpina* to *Dothiorella*. A new combination species, *Dothiorella alpina* is introduced. Furthermore, *Dothiorella symphoricarpicola* is also reported from *Acer monspessulanum* in Italy for the first time.
Fig. 106 – Phylogram generated from maximum likelihood analysis based on a combined ITS and tefl sequence dataset. Forty-six strains are included in the combined gene analyses comprising 740 total characters including gaps (ITS: 1–509 bp, tefl: 510–740 bp). The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring of the ML tree is selected to represent the phylogenetic relationships of taxa in Dothiorella, with the final ML optimization likelihood: -3536.406809. The matrix had 293 distinct alignment patterns, with 18.47% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.211039, C = 0.292321, G = 0.252470, T = 0.244170; substitution rates AC = 1.328514, AG = 2.488703, AT = 1.264091, CG = 0.687931, CT = 4.488656, GT = 1.000000; Tree-Length = 1.050171; gamma distribution shape parameter α = 0.517888. Bayesian posterior
probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.009627. Bootstrap support for maximum likelihood (ML) greater than 50% and Bayesian posterior probabilities (PP) greater than 0.95 are defined above the nodes as ML/PP. The tree is rooted to *Neofusicoccum parvum* (CMW 9081). The type strains are indicated in bold and newly generated sequences are shown in blue bold.

**Dothiorella alpina** (Y. Zhang ter & Ming Zhang) Phookamsak & K.D. Hyde, comb. nov.

Index Fungorum number: IF55723; Facesoffungi number: FoF07384

= *Spencermartinsia alpina* Y. Zhang ter & Ming Zhang, Mycosphere 7(7): 1058 (2016)

*Saprobic* on dead stolon of *Ipomoea* sp. Sexual morph: Undetermined. Asexual morph: *Conidiomata* 135–245 µm high, 150–235 µm diameter, pycnidial, immersed, raised, becoming erumpent through host epidermis, gregarious, visible as black, gnarled with dark area on host, globose to subglobose, 1-loculate, occasionally developed hyaline, palisade-like invaginations, raising from basal cavity, ostiole central, with a small, narrow papilla. *Pycnidial walls* 15–30 µm wide, composed of 3–5 layers, of large, dark brown to black pseudoparenchymatous cells, arranged in *textura angularis* to *textura globulosa*. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* 7–10 × 2–4(–5) µm (x̄ = 8.9 × 3.6 µm, n = 30), raising from the inner cavity, enteroblastic, phialidic, integrated, determinate, ampulliform to cylindrical, unbranched, aseptate, hyaline, smooth, with minute collarette, with thin periclinal wall thickening. *Conidia* 22–25(–28) × 10–12(–13) µm (x̄ = 24.4 × 11.1 µm, n = 50), initially hyaline, becoming brown to dark brown when mature, oblong with rounded ends, occasionally irregular in shape, 1-septate, smooth-walled.

![Image](image_url)

**Fig. 107** – *Dothiorella alpina* (KUN-HKAS 102212, new combination). a Appearance of conidiomata on host substrate. b Section through conidiomata. c Section through pycnidial wall. d–f Conidiogenous cells with the development of conidia. g–i Conidia. j Germinated conidium. k, l Culture characteristics on PDA (k = from above, l = from below). Scale bars: b = 100 µm, c = 50 µm, d–f, j = 20 µm, g–i = 10 µm.
Culture characteristics – Colonies on PDA fast growing, reaching 50 mm diameter after 1 week at 20–25 °C, colonies initially medium sparse with aerial mycelia, white grey, becoming dense, with hyphal lumps, circular, flat, surface smooth with edge entire, margin well defined, cottony to fluffy; colony from above and below, black with white to black hyphal lumps; not producing pigments on PDA.

Known distribution (based on molecular data) – China, Yunnan (Zhang et al. 2016), China, Yunnan (this study).

Known hosts (based on molecular data) – Platycladus orientalis (Cupressaceae) (Dissanayake et al. 2016, Zhang et al. 2016), Ipomoea sp. (this study).

Material examined – China, Yunnan Province, Kunming City, Kunming Institute of Botany, on dead stolon of Ipomoea sp. (hanging on the tree), 18 October 2016, R. Phookamsak, KIB008 (KUN-HKAS 102212, new host record), living culture, KUMCC 18–0135, KUMCC 18–0136.

GenBank numbers – ITS: MT002267, LSU: MT002266, SSU: MT002268, RPB2: MT005773 (KUMCC 18–0135); ITS: MT000718, LSU: MT000720, SSU: MT000725, RPB2: MT005774 (KUMCC 18–0136).

Notes – The NCBI BLASTn search of ITS sequence data shows that our strains (KUMCC 18–0135, KUMCC 18–0136) are identical to Spencermartinsia sp. ‘alpina’ strain CGMCC3.18001 and Spencermartinsia sp. JC-2017 strain CFCC 51564 (as Dothiorella magnoliae; You et al. 2017) with 100% similarity. Phylogenetic analyses of a concatenated ITS and tef1 sequence dataset indicated that our strains have a close affinity with S. alpina (CGMCC3.18001) with strong support (97% ML and 1.00 PP, Fig. 106). Morphological characters and phylogenetic analyses indicated that our collection is conspecific with S. alpina (Zhang et al. 2016). We, therefore, transfer S. alpina to Dothiorella. A new combination, Dothiorella alpina is designated and is reported from Ipomoea sp. in Yunnan, China for the first time.

**Dothiorella sarmentorum** (Fr.) A.J.L. Phillips, J. Luque & A. Alves, Mycologia 97: 522 (2005)

Facesoffungi number: FoF 02148

*Saprobic on a wide range of hosts including Torilis arvensis. Sexual morph: Undetermined.*

Asexual morph: *Conidiomata* 162–243 μm high × 224–297 μm diameter (x̄ = 199 × 265 μm, n = 5), stromatic, solitary or scattered, semi-immersed, uniloculate, individual, black, globose to subglobose, ostiolate. *Peridium* 28–43 μm (x̄ = 34 μm, n = 15), outer layers composed of thick-walled, dark brown, cells of *textura angularis* and inner layers composed of thin-walled, hyaline *textura angularis* cells. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* 7–9.5 × 2.5–4 μm (x̄ = 8 × 3.5 μm, n = 15), phialidic, hyaline, cylindrical to subcylindrical. *Conidia* 19–25 × 8–10 μm (x̄ = 21.5 × 9 μm, n = 30), oval to ovoid, with a broadly rounded apex and truncate base, initially hyaline, 1-celled, becoming cinnamon to sepia and 1-septate, while still attached to conidiogenous cells; detached conidia, hyaline, sepia or dark brown, 1-celled or 1-septate, moderately thick-walled, wall externally smooth, slightly constricted at the septum.

Known distribution (based on molecular data) – England, The Netherlands, Norway, Sweden, and a worldwide distribution can be observed (Phillips et al. 2005), Italy (this study).

Known hosts (based on molecular data) – Plurivorous including *Acer, Malus, Menispermum, Prunus, Pyrus, Ulmus* – (Phillips et al. 2005, Farr & Rossman 2020), *Torilis arvensis* (this study).

Material examined – Italy, Province of Ravenna [RA], Faenza, Santa Lucia, on dead aerial stem of *Torilis arvensis* (Apiaceae), 15 January 2018, E. Camporesi, IT 3677 (MFLU 18–0253, new host record).

GenBank numbers – ITS: MN782001, tef1: MN854664.

Notes – Phillips et al. (2005) introduced *Dothiorella sarmentorum* based on the asexual morph of *Botryosphaeria sarmentorum*. This species has a worldwide distribution and has been recorded from 34 host species (Phillips et al. 2005, 2013). The sexual morph of *D. sarmentorum* is characterized by partially erumpent ascomata with papillate ostioles, 4–6(-8)-spored asci and
oblong to ovate (0–)1-septate, finely verruculose ascospores, widest in the middle part (Phillips et al. 2013).

![Image](image_url)

**Fig. 108** – *Dothiorella sarmentorum* (MFLU 18–0253, new host record). a, b Appearance of conidiomata on the host. c Vertical section through conidioma. d Ostiole. e Peridium. f Conidia developing on conidiogenous cells. g Mature dark brown conidia. Scale bars: c, d = 50 μm, e = 20 μm, f, g = 20 μm.

*Dothiorella symphoricarposicola* W.J. Li, Jian K. Liu & K.D. Hyde, Mycol. 35(3): 265 (2014)

Facesoffungi Number: FoF 06222

*Saprobic* on dead branch of *Acer opalus*. Sexual morph: Undetermined. Asexual morph: *Coelomycetous*. Conidiomata 220–265 μm high × 260–380 μm diameter (x̄ = 245 × 340 μm, n = 10), pycnidial, stromatic, solitary or clustered, immersed in the host, erumpent at maturity, dark brown to black, ostiolate, apapillate. **Peridium** 25–35 μm wide, outer and inner layers composed of dark brown and thin-walled hyaline **textura angularis**. Conidiogenous cells 10–22 μm high × 4–6 μm wide, phialidic, hyaline, thin-walled, smooth, cylindrical, and swollen at the base, integrated. **Conidia** 21–27 × 11–16 μm (x̄ = 24×13 μm, n = 50), globose to subglobose, with rounded apex, initially hyaline, becoming dark brown and 1-septate at maturity, with moderately thick wall.

Culture characteristics – Colonies growing on PDA, covering the entire plate in 5 days at 28 °C, mycelium grey to olivaceous black at the surface and olivaceous black from below.

Known distribution – Italy (Dissanayake et al. 2016, Farr & Rossman 2020 and this study).

Known hosts (based on molecular data) – On dead aerial branch of *Acer opalus* (Rosaceae), *Dothiorella symphoricarposicola* strains have also been reported from *Corylus avellana*, *Symphoricarpus* sp., *Sambucus nigra*, *Laurus nobilis* and *Laburnum alpinum* in Italy (Dissanayake et al. 2016, Farr & Rossman 2020, this study).
Material examined – Italy, Province of Forlí-Cesena [FC], Pieve di Rivoschio – Bangno di Romagna, on dead aerial branch of *Acer opalus* (Rosaceae), 3 May 2017, E. Camporesi, IT 310 (MFLU 3150027, new host record), living culture: JZB: 3150027.

GenBank number – ITS: MN989423.

Notes – Our collection obtained from dead aerial branches of *Acer opalus* was identified as *Dothiorella symphoricarposicola* based on morphology and phylogeny. Our collection clustered with the reference strain of the *Dothiorella symphoricarposicola* (MFLUCC 13–0196), in the combined ITS and *tef1* sequence phylogeny with 71% statistical support. Furthermore, our isolate showed 97% ITS similarity with *Dothiorella symphoricarposicola*. This is the first record of *Dothiorella symphoricarposicola* reported from *Acer opalus* from Italy.

**Fig. 109** – *Dothiorella symphoricarposicola* (MFLU 17–0729, new host record). a Appearance of conidiomata on the host. b Section through the conidioma. c Conidioma cell wall d, e Immature conidia attached to conidiogenous cell. f Mature conidia. Scale bars: a, b = 100 µm, c, d = 50 µm, e, f = 10 µm.
Lasiodiplodia Ellis & Everh.

Lasiodiplodia is a common genus of Botryosphaeriaceae. They are saprobes, endophytes and pathogens occur on numerous hosts with a wide geographic range (Dissanayake et al. 2016, Doilom et al. 2017b, Tibpromma et al. 2018a, Hyde et al. 2019). Lasiodiplodia is characterised by initially hyaline, aseptate and thick-walled conidia, becoming dark brown and septate with irregular longitudinal striations, and pycnidial paraphyses (Phillips et al. 2008, Liu et al. 2012b, Dou et al. 2017b). There are 67 epithets reported for Lasiodiplodia in Index Fungorum (2020). Recently many new species were introduced (Dou 2017b, Tibpromma et al. 2018a, de Silva et al. 2019).

Lasiodiplodia theobromae (Pat.) Griffon & Maubl., Bull. Soc. Mycol. Fr. 25: 57 (1909)  
Fig. 111

Facesoffungi number: FoF 00167

Saprobic on dead leaf sheath of Phoenix roebelenii. Sexual morph: not established. Asexual morph: Conidiomata pycnidial, dark brown to black, solitary or clustered, convex on host tissue, uni to multi-loculate, immersed in the host becoming erumpent when mature. Paraphyses 1.5–3.5 μm wide, hyaline, septate, branched. Conidiogenous cells hyaline, cylindrical, phialidic. Conidia 19–25 × 10–16 (x = 25 × 13 μm, n = 20), initially hyaline, aseptate, subovoid to ellipsoid-ovoid, with granular content, finally becoming dark brown, 1–septate at centre, thick-walled, upper cells wider than lower cells, truncate or rounded at the base, broadly rounded at the apex, with longitudinal striations from apex to base, guttulate.

Culture characteristics – Conidia germinating on PDA within 4–5 h. Colonies on PDA initially whitened in a few days, after 5–7 days becoming grey to black, filamentous, raised, fluffy, dense, reaching the edge of the Petri-dish after 2 days.

Known distribution (based on molecular data) – Widely distributed in Asia, Africa, America, Europe (Farr and Rossman 2020); e.g. Papua New Guinea (Phillips et al. 2005), South Africa (Gauteng) (Jami et al. 2015), (KwaZulu-Natal) (Osorio et al. 2017, Begoude et al. 2010), (Limpopo) (Mehl et al. 2017), (Western Cape) (Van Niekerk et al. 2004), Uganda (Slippers et al. 2004), USA (Arizona, California) (Inderbitzin et al. 2010), (Florida) (Mehl et al. 2017), (Hawaii) (Marincowitz et al. 2008), Thailand (Tibpromma et al. 2018a), (Phayao Province) (Doilom et al. 2015, 2017b), China (Tennakoon et al. 2016).

Known hosts (based on molecular data) – Barringtonia racemosa (Osorio et al. 2017), Eucalyptus amplifolia (Mehl et al. 2017), Leucospermum sp. (Marincowitz et al. 2008), Mangifera indica (Mehl et al. 2017), Pandanus sp. (Tibpromma et al. 2018a), Pistacia vera, Prunus domesticus (Inderbitzin et al. 2010), Tectona grandis (Doilom et al. 2015, 2017b), Terminalia catappa (Begoude et al. 2010), Vachellia karroo (Jami et al. 2015), Vitex donniana (Slippers et al. 2004), Vitis vinifera (Van Niekerk et al. 2004), Phoenix roebelenii (this study), For more information see Farr & Rossman (2020).

Material examined – Thailand, Chiang Mai Province, Sansai District, on dead leaf sheath of Phoenix roebelenii (Arecales), 2 February 2012, M. Doilom & R. Phookamsak (MFLU 19-1559, new host record), living culture (MFLUCC 12-0173).

GenBank numbers – ITS: MN582743, tef1: MN629743.

Notes – Lasiodiplodia theobromae has been recorded on numerous plant species in Thailand such as Hevea brasiliensis, Licuala longicalycata, Pandanus sp. and Tectona grandis (Pinruan et al. 2007, Scepheueak et al. 2011, Doilom et al. 2015, 2016, 2017b, Tibpromma et al. 2018a, Farr & Rossman 2020). The species is also known on several palm species such as Phoenix canariensis in Florida, P. dactylifera in Egypt, India, Oman, Venezuela, and P. hanceana in China and Hong Kong (Anonymous 1960, Mathur 1979, Lu et al. 2000, Zhuang 2001, El-Deeb et al. 2007, Al-Sadi et al. 2013, Li et al. 2018, Farr & Rossman 2020). However, it has not yet been reported on Phoenix roebelenii (palm) in Thailand (Farr & Rossman 2020). We identify specimen (MFLU 19–1559) as L. theobromae based on morphology and phylogeny. Phylogenetic analysis of the combined sequence data of ITS and tef1 showed that our strain (MFLUCC 12–0173) clustered with the ex-neotype strain of L. theobromae (CBS 164.96) and other strains of L. theobromae with good
support (Fig. 110, 69% MP/ 95% ML). This is a new host record of *Lasiodiplodia theobromae* on *Phoenix roebelenii* in Thailand.

**Neofusicoccum** Crous, Slippers & A.J.L. Phillips

*Neofusicoccum* species have a wide geographical and host range (Lopes et al. 2016). They are endophytes, saprobes and pathogens which caused dieback, cankers and trunk diseases (Golzar & Burgess 2011, Massonnet et al. 2017, Jami et al. 2018). Recently, many species have been introduced based on morphology and multiple-gene phylogenies by Marin-Felix et al. (2017a), Zhang et al. (2017) and Tibpromma et al. (2018a).

Fig. 110 – Phylogram generated from maximum parsimony analysis based on combined ITS and *tef1* sequence data. Thirty-five strains are included in the combined gene analyses comprising 739 characters after alignment (482 characters for ITS, 257 characters for *tef1*). *Diplodia mutila* (CBS 112553) and *D. seriata* (CBS 112555) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -2080.598320 is presented. The matrix had 125 distinct alignment patterns, with 6.41% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.205263, C = 0.292383, G = 0.256816, T = 0.245539; substitution rates AC = 0.821151, AG = 3.299106, AT = 1.173326, CG = 0.763958, CT = 3.692977, GT = 1.000000; gamma distribution shape parameter α = 0.067254. Bootstrap values for maximum parsimony and maximum likelihood equal to or greater than 50 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.

**Neofusicoccum parvum** (Pennycook & Samuels) Crous, Slippers & A.J.L. Phillips, Stud. Mycol. 55: 248 (2006)  
Facesoffungi number: FoF02411
Saprobic on dead branch. Ascostromata 155–250 μm diameter, 115–190 μm high (including the papilla), black dots and convex on host tissue, appearing through cracks in bark, solitary or clustered, when cut horizontally locules visible as white contents and dark ascospore dots, semi-immersed to erumpent under epidermis, individually globose to subglobose, with papilla. Ostiole central, circular, papillate. Peridium 25–65 μm, comprising two layers, outer layer composed of dark brown to black, thick-walled cells of textura angularis, inner layer composed of hyaline, thin-walled cells of textura angularis. Hamathecium comprising 2–4 μm wide, hyaline, hyphae-like, numerous, septate, pseudoparaphyses, slightly constricted at the septa, embedded in gelatinous matrix. Asci 55–135 × 16–23 μm (x̄ = 84 × 19 μm, n = 20), 8-spored, bitunicate, fissitunicate, clavate to cylindro-clavate, short to long pedicellate, with ocular chamber, apically rounded. Ascospores 17–25.5 × 7–10 μm (x̄ = 20 × 8 μm, n = 20), uniseriate at the base, 2–3-seriate at the centre and end, hyaline, ellipsoidal to fusiform, aseptate, wider in the centre, rounded or acute at the ends, thick-walled.

Fig. 111 – Lasiodiplodia theobromae )MFLU 19–1559, new host record. a Conidiomata on host substrate. b Vertical section through conidioma. c Conidiogenous cells with paraphyses. d Conidia attached to conidiogenous cells. e Immature conidium. f–i Mature conidia. j Germinated conidium. k, l Colony on PDA (k from above view, l from below view). Notes: c, e stained in lactophenol cotton blue. Scale bars: b = 50 μm, c, d = 10 μm, e–i = 5 μm, j = 20 μm.

Culture characteristics – Colonies on PDA fluffy, fairly dense, aerial, raised, filamentous, pigments not produced, white at first, becoming gray to grey-black after one to two weeks.

Known distribution (based on molecular data) – worldwide e.g. Algeria (Berraf-Tebbal et al. 2014), Australia, Indonesia (Sumatra), Brazil, China (Sichuan), France (Van Niekerk et al. 2004), Italy (Milan) (Moricca et al. 2012), New Zealand (TePuke) (Alves et al. 2005), Portugal (Palmela) (Phillips et al. 2008), (Hogsback, KwaZulu-Natal, Mpumalanga) (Slippers et al. 2004), (e.g. Gauteng, Limpopo) (Mehl et al. 2017), USA (California) (Úrbez-Torres et al. 2007), (Hawaii) (Slippers et al. 2004), Thailand (Chiang Mai Province) (Liu et al. 2012b, Trakunyingcharoen et al. 2015), (Phayao Province) (this study), For more information see Farr & Rossman (2020).
Known hosts (based on molecular data) – *Actinidia deliciosa*, *Heteropyxis natalensis*, *Malus sylvestris*, *Populus nigra*, *Eucalyptus* sp., *E. grandis*, *E. smithii*, *Ribes* sp., *Sequoia gigantea* (Slippers et al. 2004), *Acer pseudoplatanus*, *Quercus robur* (Moricca et al. 2012), *Actinidia deliciosa*, *A. chinensis*, *Populus nigra*, *Eucalyptus obliqua*, *Prunus cerasoides* (Trakunyingcharoen et al. 2015), *Pinus nigra*, *Vitis vinifera* (Crous et al. 2006a, Phillips et al. 2008), *Aesculus hippocastanum*, *Eucalyptus globulus*, *Ferula communis* (Lopes et al. 2016), *Juglans regia*, *Salix sp.*, *Sclerocarya birrea* subsp. *caffra* (Mehl et al. 2017), *Avicennia marina*, *Bruguiera gymnorrhiza*, *Lumnitzera racemosa*, *Rhizophora mucronata* (Osorio et al. 2017), *Linum usitatissimum* (Liu et al. 2012b), *Vitis vinifera* (Alves et al. 2005), *Mangifera indica* (Mehl et al. 2017, this study), For more information see Farr & Rossman (2020).

Material examined – Thailand, Phayao Province, Muang District, on dead branch of *Mangifera indica* (Anacardiaceae), 12 March 2012, M. Doilom (MFLU 19–1562, new host record), living culture (MFLUCC 12–0380).

GenBank numbers – ITS: MN582744, *tef1*: MN629744, TUB: MN643160.

![Phylogram](image)

**Fig. 112** – Phylogram generated from maximum likelihood analysis based on combined ITS, *tef1* and TUB sequence data. Fifteen strains are included in the combined gene analyses comprising 1247 characters after alignment (499 characters for ITS, 314 characters for *tef1*, 434 characters for TUB). *Botryosphaeria corticis* (CBS 119047) and *B. dothidea* (CMW 8000) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -3032.363425 is presented. The matrix had 192 distinct alignment patterns, with 6.38% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.201816, C = 0.314088, G = 0.262406, T = 0.221689; substitution rates AC = 1.071994, AG = 4.639898, AT = 1.573574, CG = 1.216826, CT = 6.907567, GT = 1.000000; gamma distribution shape parameter α = 0.171757. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.
Notes – *Neofusicoccum parvum* has been previously reported on *Eucalyptus obliqua*, *Linum usitatissimum* and *Prunus cerasoides* in Thailand (Liu et al. 2012b, Trakunyingcharoen et al. 2015). The holotype is from dead branches of *Populus nigra* in New Zealand (Pennycook & Samuels 1985). Although this species is widely distributed on mango in many countries such as Australia, Brazil, Italy, México, Peru, Puerto Rico, Spain and Taiwan (Javier-Alva et al. 2009, Ni et al. 2012, Ismail 2013, Sandoval-Sanchez et al. 2013, Serrato-Diaz et al. 2013, Arjona-Girona & Lopez-Herrera 2016), it has not yet been reported on *Mangifera indica* (mango) in Thailand (Farr & Rossman 2020). Conidial size of our specimen is similar to the type *N. parvum* (20 × 8 μm vs 20.8 × 9.2 μm) (Pennycook & Samuels 1985) as well as the *N. parvum* isolate identified by Liu et al. (2012b) (20 × 8 μm). The isolate MFLUCC 12-0380 clustered with the ex-type strain of *N. parvum* (CMW 9081) and *N. parvum* (MFLUCC 11-0184) (Fig. 112). Thus we identify our collection as *N. parvum* and this is the first report of *N. parvum* on *Mangifera indica* in Thailand.

**Fig. 113 – Neofusicoccum parvum** (MFLU 19–1562, new host record). a Ascomata on host substrate. b Close up of ascomata on host. c Vertical section through ascoma. d Peridium. e, f Immature and mature asci with pseudoparaphyses. g, h Mature asci. i–k Ascospores. Notes: e–i stained in lactophenol cotton blue. Scale bars: c = 50 μm, d = 20 μm, e = 30 μm, f–h = 10 μm, i–k = 5 μm.
**Venturiales** Y. Zhang ter, C.L. Schoch & K.D. Hyde  
**Sympoventuriaceae** Y. Zhang ter, C.L. Schoch & K.D. Hyde

Sympoventuriaceae belongs to Venturiales (Dothideomycetes). This family is related to *Verruconis* and nested within the Sympoventuriaceae. In their natural habitat, fungi from this family commonly occur in soils and decaying plant material (Barron & Busch 1962, Dwivedi 1959). Species are also known as opportunistic vertebrate pathogens (de Hoog et al. 2000, Horré et al. 1999) and can cause infections in humans and animals (Ge et al. 2012, Padhye et al. 1994, Salkin et al. 1990, Singh et al. 2006, see review in Yarita et al. 2007).

**Ochroconis** de Hoog & Arx

de Hoog & von Arx (1973) introduced *Ochroconis constricta* as the type species and they are characterised with sympodial conidiogenesis and septate, ellipsoidal conidia which were liberated rhexolytically. Previously, species of *Ochroconis* have been placed in different genera: *Diplorhinotrichum* (*D. gallopavum*; Georg et al. 1964), *Dactylaria* (*D. humicola* and *D. gallopava*; Bhatt & Kendrick 1968), and *Scolecobasidium* (*S. terreum* and *S. constrictum*; Abbott 1927).

**Ochroconis musae** (G.Y. Sun & Lu Hao) Samerp. & de Hoog, Mycological Progress 14 (2/6): 8 (2015)  
Facesoffungi number: FoF06957  
Saprobic on *Dracaena*. Sexual morph: Undetermined. Asexual morph: Full descriptions of this species were given by Hao et al. 2013( and by Samerpitak et al. )2015( under its synonymous name *O. mirabilis*. Conidiophores mostly arising laterally from vegetative hyphae, erect or flexuous, cylindrical with 1–2 septa, at 25 °C. *Conidia* were T or Y shaped or bilobed, cylindrical, ellipsoid, clavate or fusiform.

Culture characteristics – Conidia germinated on PDA within 24 hours with germ tubes produced from one or both end cells, mostly from basal cell of conidia. Colonies on PDA reaching 25–30 mm in diameter after 2 weeks at 25–30 °C, colonies circular, flat, moderately expanding, smooth, dry, greyish brown to dark brown, submerged colony margin, reverse as dark brown in the central portion; not producing pigmentation in agar. Mycelium hyphae subhyaline to pale brown, smooth- and thick-walled.

Material examined – Thailand, Chiangmai Province, on dead leave of *Dracaena*, 27 September 2017, Napalai Chaiwan, NCCM003 living culture, MFLUCC 17–2598 (new host record).

GenBank numbers – ITS: MN788641, LSU: MN788642

Notes – Our strain shares similar characters with *O. musae* which was collected from banana, *Musa basjoo*, in Ledong county, Haikou City, Hainan Province, China (Samerpitak et al. 2015). Phylogenetic analysis (Fig. 114) also supports the close relatedness to *O. musae*. The mycelium of *Ochroconis* consists of smooth, pale brown to medium brown. However, previously *O. musae* has only been reported from banana. This study provides the first report of new host record of this species from *Dracaena* as well as its first report from Thailand.

**Class Eurotiomycetes** Tehler ex O.E. Eriksson & K. Winka  
**Chaeotothyriales** M.E. Barr  
**Trichomeriaceae** Chomnunti & K.D. Hyde.

Species are epiphytic on the surface of leaves associated with honey dew insect excretions and colonies are often mixed with capnodiaceous taxa. Phylogenetic data (LSU and ITS) clearly shows that Trichomeriaceae belongs in Chaetothyriales (Chomnunti et al. 2012a). The type genus is *Trichomerium*.

**Trichomerium** Speg.

*Trichomerium* was introduced by Spegazzini (1918) with *Trichomerium coffeicolum* as the type species. Species in this genus are foliar epiphytes with superficial, setiferous, uniloculate
ascostromata surrounded by loosely interwoven mycelium, with bitunicate asci and hyaline, septate ascospores (Spegazzini 1918, Chomnunti et al. 2012a, Hongsanan et al. 2016). The asexual morph of *Trichomerium gloeosporum* was recorded as *Tripospermum*-like (Hongsanan et al. 2016). Some species of *Trichomerium* can be found on rocks and are slow-growing. Molecular analyses indicated that species of *Trichomerium* cluster separately from Capnodiaeae and Chaetothyriaceae (Chomnunti et al. 2012a, b, 2014, Yang et al. 2014, Hongsanan et al. 2016).

**Fig. 114** – Phylogram generated from maximum likelihood analysis based on combined LSU and ITS sequence data. Thirty-five strains are included in the combined gene analyses comprising 2019 characters after alignment (972 characters for LSU, 1047 characters for ITS). *Fusicladium africanum* CPC12828 is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -16950.850707 is presented. The matrix had 1069 distinct alignment patterns, with 37.28% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.239425, C = 0.234508, G = 0.322263, T = 0.203804; substitution rates AC = 0.815774, AG = 1.380551, AT = 0.895824, CG = 0.990791, CT = 3.616055, GT = 1.000000; gamma distribution shape parameter α = 0.259219. Bootstrap values for maximum likelihood equal
to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. The new host record generated sequence is indicated in blue.

Fig. 115 – *Ochroconis musae* )MFLUCC 17–2598, new host record(, a Culture. b–f Conidiophore g Conidiogenous cell h–k Conidia. Scale bars: b–g = 20 μm, h–k = 10 μm.

*Trichomerium camporesii* Marasinghe & K.D. Hyde sp. nov.

Index Fungorum Number: IF556916; Facesoffungi number: FoF06236

Etymology – Named after Erio Camporesi, a great fungi collector

Holotype – MFLU 19–2251

_Epiphytic_ on the surface of _Psidium guajava_ leaves. Black sheets of mycelia cover the leaves of the host with dark brown superficial, scattered, ascostromata. Hyphae septate, cylindrical, pale brown to brown with constrictions at the septa, 4.5–6 μm wide (\(\bar{x} = 5.5\) μm, \(n = 20\)). _Ascostroma_ 55–65 × 45–55 μm (\(\bar{x} = 60 \times 48\) μm, \(n = 10\)) globose to subglobose, brown, covered with attached aseptate to septate, dark brown to brown setae. _Ascostroma wall_ 70-80 μm wide (\(\bar{x} = 65\) μm, \(n = 10\)), thick-walled, inwardly hyaline, pale brown and brown towards the outside, comprised 2–3 layers of _textura angularis_. _Asci_ 55–60 × 20–25 μm (\(\bar{x} = 57 \times 22\) μm, \(n = 10\)), 8-spored, bitunicate, ellipsoid to clavate, some subglobose, with or without short pedicle. _Paraphyses_ not observed. _Ascospores_ 25–30 × 5–10 μm (\(\bar{x} = 28 \times 8\) μm, \(n = 10\)), fasciculate, hyaline, fusoid, ends rounded, 3 septate, with 1-2 distinct guttules, slightly curved at the middle, smooth-walled. Asexual morph: Undetermined.
Material examined – Thailand, Chiang Rai, on living leaf of *Psidium guajava* (Myrtaceae), 21 June 2018, M.W.D Sandamali (MFLU 19-2251, holotype).

GenBank Numbers – LSU: MN644511, ITS: MN644590.

**Fig. 116** – Phylogram generated from maximum likelihood analysis based on combined LSU and ITS sequence data representing Trichomeriaceae, Cyphellophoraceae, Chaetothyriaceae and the Capnodiaceae (outgroup). Related sequences are taken from Maharachchikumbura et al. (2018). Twenty five strains are included in the combined analyses which comprise 1629 characters (791 characters for LSU and 837 characters for ITS) including gaps. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -10206.223000 is presented. The matrix had 701 distinct alignment patterns, with 20.98% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.244909, C = 0.243885, G = 0.276707, T = 0.234500; substitution rates AC = 1.442198, AG = 1.950067, AT = 1.512274, CG = 1.034625, CT = 4.856831, GT = 1.000000; gamma distribution shape parameter $\alpha = 0.293070$. Bootstrap values for maximum likelihood (ML) equal to or greater than 60% and clade credibility values greater than
0.80 (the rounding of values to 2 decimal proportions) from Bayesian-inference analysis labeled on the nodes. Ex-type strains are in bold and black, the new isolate is indicated in bold and blue.

**Fig. 117** – *Trichomerium camporesii* (MFLU 19–2251 holotype). a Dead leaf specimen. b, c Ascostromata on the surface of leaf. d Ascostroma with setae. e Section through ascoma. f Ascoma setae. g–j Asci. k, l Ascospores. Scale bars: d–f = 100 µm, g = 50 µm, h–j = 20 µm, k, l = 10 µm.

Notes – *Trichomerium camporesii* is similar to *T. foliicola*, but differs in having ellipsoid to clavate asci and slightly curved ascospores with relatively large 1–4 guttules, while *T. foliicola* has cylindrical to clavate asci with relatively small 1–2-guttulate ascospores. Phylogenetic analysis based on LSU and ITS sequence data shows that *T. camporesii* constitutes an independent lineage basal to *T. eucalypti* (Fig. 116, 92 % ML, 1.00 BYPP).
Class Lecanoromycetes O.E. Erikss. & Winka
Lecanoromycetidae families incertae sedis
Ostropales Nannf
Stictidaceae Fr.

The family contains both lichenized and non-lichenized fungi (Wedin et al. 2005).

**Stictis** Pers.

*Stictis* species are widely spread in tropical and temperate evergreen forests (Joshi et al. 2012). The genus is characterized by orbicular ascoma opening by a pore, periphysoids in apothecial margins that extend down the whole length, a hymenium that splits from the margin when dry, a thick crystalline layer in the ascoma margin. *Stictis* was monographed by Sherwood (Sherwood 1977) and Lücking et al. (2017).

**Stictis urceolata** (Ach.) Gilenstam (Ach.) Gilenstam, Lichenologist 37(1): 74 (2005)  
Fig. 119

Facesoffungi number: FoF07378

Non-lichenized fungus, corticolous, forming a hyaline epiphloeodal hyphal felt or thallus, 30–60 μm thick. Sexual morph: *Ascomata*, urceolate, solitary, sometimes aggregated in two, at first immersed, opening by a pore, becoming erumpent and finally nearly superficial, 350–420 μm in diameter, round, chroodiscoid; disc brownish to flesh coloured, densely pruinose, splitting away from the margin, up to 0.4 mm in diameter, deeply immersed; margin radiate, effigurate, lacerate, 5–6 lobed, white-pruinose, eroded in older apothecia, 60–120(–145) μm thick in cross section, hyaline to darken in older *apothecia*, sometimes layered, entirely encrusted in crystals. Outer exciple layer thicker; inner *exciple* layer brown, branched periphysoids present, forming the innermost layer of the margin, separated from the outer wall by crystals; crystals forming a dense layer along the inner margin of apothecia, 50–60 μm thick. *Epihymenium* indistinct, granular, hyaline to slightly brownish, usually covered by 40–55 μm high crystalline layer, *hymenium* hyaline, interspersed, separated from the margin in dry condition, *Paraphyses* filiform, branched, with thickened apical cell, dense, conglutinate. *Asci* 50–70 (–80) × 3.5–5 μm, 8-spored, cylindrical, bitunicate. *Ascospores* 30–55 × 3.5–4.5 μm, hyaline, cylindrical to fusiform, transversely septe, sheathed, locules broader than longer. Asexual morph: Undetermined.

Known distribution (based on molecular data) – North America, England (Tuckerman, et al. 1847), Taiwan, Chia Yi, Dahu (this study)

Known hosts (based on molecular data) – *Miscanthus* sp. (this study)

Material examined – Taiwan, Chia Yi, Dahu, on decaying stems of *Miscanthus* sp. (Poaceae), 27 April 2018, A. Karunarathna, AKTW 52 (MFLU 19–2695, new host and geographical record); ex-type living culture,NCYUCC 19–0365.

GenBank number – LSU: MN989186.

Notes – Herein, we provide the new host and geographical records for *Stictis urceolata* based on morphology and phylogeny (Fig. 118).

**Vibrisseaceae** Korf

Vibrisseaceae was introduced by Korf (1990). Five genera and about 80 species are currently accepted in this family (Zheng & Zhuang 2017).

**Phialocephala** W.B. Kendr.

Kendrick (1961) introduced *Phialocephala*, which was previously known as *Leptographium*. Thirty-four species were accepted in *Phialocephala* (Tanney et al. 2016, Crous et al. 2017) and they occur as endophytes (Rashmi et al. 2019). However, several phylogenetic studies revealed that members of this genus are polyphyletic (Grüning et al. 2009, Wong et al. 2015, Ekanayaka et al. 2019).

**Phialocephala humicola** S.C. Jong & E.E. Davis, Mycologia 64 (6): 1352 (1972)  
Fig. 121
Facesoffungi number: FoF06717

*Saprobic* on decaying wood. Sexual morph: Undetermined. Asexual morph: *Colonies* on natural substate effuse, white, velvety. *Mycelium* mostly immersed, composed of hyaline, branched, septate and constricted at the septa, guttulate hyphae, 1.5–3 μm wide. *Conidiophores* up to 300 μm long, 5–7 μm wide at base, 3.5–4.5 μm wide at tip, macronematous, mononematous, erect, straight or broadly curved, subcylindrical, wider at base, septate, median brown, thick-walled, branched at the apex. *Conidiogenous cells* phialidic, discrete, determinate, terminal, clustered at the apex of conidiophores, hyaline, thin-walled, cylindrical, with inconspicuous openings, with minute collarettes. *Conidia* 2.5–4 μm long, 2–3.5 μm at widest (\(\bar{x} = 3.5 \times 2.5 \mu m, n = 30\)), acrogenous, aggregated in slimy and white masses, acrogenous, solitary to catenate, hyaline, cuneate, rhomboid or shield-shaped, aseptate, smooth.

Culture characteristics – Conidia germinated within 48 h. Germ tubes produced from one angle. Mycelia superficial, decumbent, irregular, with entire edge, yellowish brown at center, pale brown at circumference from above.

Known distribution (based on molecular data) – Japan, Tokyo, USA, New Jersey (Kiyuna et al. 2012), China, Guizhou (Jie et al. 2013), Thailand (this study).

Known hosts (based on molecular data) – Decayed tree, soil (Kiyuna et al. 2012), Decaying wood (this study)

Material examined – Thailand, Phrae Province, Rong Kwang, on decaying wood, 10 January 2018, N.G. Liu, N016 (MFLU 19–0005)

GenBank numbers – ITS: MN999924, LSU: MN901120, SSU: MN901150.

Notes – Jong and Davis (1972) introduced *Phialocephala humicola* from a soil sample collected from Cape May, New Jersey, USA. Subsequently, this species was isolated from soil in Japan (Matsushima 1975), USA (Ellis 1976), China (Matsushima 1980) and Australia (Matsushima 1989). Matsushima (1975) reported *P. humicola* on *Quercus* sp. in Nara City, Japan, and Matsushima (1980) found this species on leaves of *Areca catechu* in Taiwan, China. Our collection differs from *P. humicola* (ATCC 22801) in having cuneate, rhomboid or shield-shaped conidia, while the latter has ellipsoidal conidia, but the phylogenetic analysis of combined LSU and ITS sequence data confirmed that our collection is *P. humicola*. This is a new geographical record of *P. humicola* in Thailand (Fig. 120).

**Rhytismatales genera incertae sedis**

*Apiculospora* Wijayaw., Camporesi, A.J.L. Phillips & K.D. Hyde

*Apiculospora* is monotypic. This genus was placed in Helotiales genera incertae sedis by Wijayawardene et al. (2016). However, our phylogenetic analysis based on LSU, SSU and ITS sequence data show that *Apiculospora* is related to Porodiplodiaceae (Rhytismatales). The sexual morph of *Apiculospora* is undetermined. Future collections and DNA sequence analyses are needed to connect *Apiculospora* to its sexual morph.

*Apiculospora spartii* Wijayaw., W.J. Li, Camporesi, A.J.L. Phillips & K.D. Hyde, Fungal Diversity: (2016) Fig. 123

Facesoffungi number: FoF01426

*Saprobic* on dead twigs of *Clematis vitalba*, forming conspicuous rounded, black, conidiomata. Sexual morph: undetermined. Asexual morph: *Conidiomata* 150–300 μm diameter, 250–300 μm high, dark brown to black, acervular, solitary to gregarious, initially immersed, ultimately erumpent, globose to subglobose, unilocular. *Ostiole* absent, dehiscence by apical wall in the middle part. *Conidioma* wall composed of thin-walled, pale brown cells of *textura angularis* in the basal part. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* 7–18 × 2–3 μm, hyaline, holoblastic, cylindrical, thick-walled, smooth-walled, arising from the uppermost cells of the basal wall. *Conidia* 13–20 × 7–11 μm (\(\bar{x} = 15 \times 8.5 \mu m; n = 20\)), hyaline when young, becoming dark brown with age, fusiform, obtuse at apex, narrowed and slightly truncated at base, straight or slightly curved, 1-septate, constricted at septum, thick and smooth-walled, guttulate.
Culture characteristics – Colonies on PDA, reaching 10 mm diameter, after 7d, circular, whitened, spreading, flattened, felt-like, sparse, aerial, surface, smooth with crenate edge, filamentous; reverse yellowish in the central zone, whitened in edge.

Known distribution (based on molecular data) – Italy (Wijayawardene et al. 2016, this study)

Known hosts (based on molecular data) – Spartium junceum (Wijayawardene et al. 2016), Clematis vitalba (this study)

Material examined – Italy, Province of Forlì-Cesena, Civitella di Romagna, Voltre, on dead twigs of Clematis vitalba (Ranunculaceae), 2 December 2013, E. Camporesi, IT1553-2 (MFLU 19–2559, new host record), living culture MFLUCC 15–0584 = CMP 21841.

Fig. 118 – Phylogenetic tree generated from maximum likelihood (ML) based on LSU and mtSSU sequences. Bootstrap support (BS) values above 50% and Bayesian posterior probabilities equal or greater than 0.90 are shown above the branches at nodes. The tree is rooted with Cryptodiscus foveolaris. The best RaxML tree with a final likelihood value of -4629.887606 is presented. The matrix had 356 distinct alignment patterns, with 7.44 % undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.310275, C = 0.181487, G = 0.251212, T = 0.257026; substitution rates AC = 0.679339, AG = 2.161324, AT = 1.633529, CG = 0.387952, CT = 5.104996, GT = 1.000000; gamma distribution shape parameter α = 0.637625. Newly generate sequence is indicated in blue bold.
**Fig. 119** – *Stictis urceolata* (MFLU 19–2695, new host and geographical record). a Appearance of apothecia on host. b Apothecia. c Longitudinal section of apothecia. d Excipulum. e Paraphyses. f Asci. g, h Ascospores. i Germinating ascospore. Scale bars: c = 20 µm, d–e = 10 µm, f–h = 10 µm, i = 5 µm.

GenBank Numbers – (MFLUCC: 13–0400); LSU: MN660233, ITS: MN688212, SSU: MN688207, tef1: MN683863, rpb2: MN683866; (MFLUCC: 15–0584); LSU: MN688206, SSU: MN688208, tef1: MN683864.

Notes – This genus is characterized by acervular conidiomata with brown to dark brown, fusiform, 1-septate conidia (Wijayawardene et al. 2016). Our collection shares similar conidiomata, conidiogenous cells and conidia with *A. spartii*, but the latter has slightly larger conidia (17–25 × 8–11 µm (x̄ = 21.3 × 9.5 µm). *Apiculospora spartii* was collected from *Spartium junceum* (*Leguminosae*), thus our collection on *Clematis vitalba* is regarded as a new host record (Fig. 122).

Additional notes – The sequence of *Chalara* sp. (voucher: MFLU 18–1812, MFLU 18–1813) submitted by Ekanayaka et al. (2019) is in fact *Apiculospora spartii*. The sequence similarities between MFLU 18–1812 and MFLUCC 13–0400 (type strain) are 99% (875/877, 2 gaps) in LSU, 100% (474/474) in ITS, and 100% (910/910) in SSU. The morphology of these two collections is almost similar to *Apiculospora spartii* (not shown here).

**Class Sordariomycetes** O.E. Erikss. & Winka
**Subclass Diaporthomycetidae** Senan. et al.
**Atractosporales** H. Zhang et al.
**Conlariaceae** H. Zhang, K.D. Hyde & Maharachch.

We follow the latest treatments and updated accounts in Zhang et al. (2017), Phookamsak et al. (2019), Xie et al. (2019) and Hyde et al. (2020). *Conlarium aquaticum* is reported from *Thysanolaena maxima* Kuntze from Yunnan, China for the first time.
Fig. 120 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, nad1, nat and RPB1 sequence data. Twenty-eight strains are included in the combined gene analyses comprising 4587 characters after alignment (922 characters for LSU, 609 characters for ITS, 1384 characters for nad1, 859 characters for nat, 813 characters for RPB1). Holwaya mucida (B-70-0009352) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -14269.181523 is presented. The matrix had 795 distinct alignment patterns, with 70.07% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.277515, C = 0.201458, G = 0.229770, T = 0.291258; substitution rates AC = 1.632078, AG = 3.050144, AT = 1.027293, CG = 1.759203, CT = 5.265005, GT = 1.000000; gamma distribution shape parameter α = 0.676190. Bootstrap values for maximum likelihood equal or greater than 75% and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches, respectively. Ex-type strains are in black bold. The newly generated sequence is indicated in blue bold.
**Fig. 121** – *Phialocephala humicola* (MFLU 19–2852, new geographical record). a, b Conidiophores and conidia. c, d Conidiogenous cells and conidia. e, f Conidia. g Colony on PDA media. Scale bar: a = 20 μm, b = 30 μm, c = 5 μm, d, e, f = 3 μm.

**Conlarium** F. Liu & L. Cai.

*Conlarium* is the only genus of Conlariaceae introduced by Liu et al. (2012a) and is typified by *C. dupliciasmcosporum* collected from submerged wood in Guangdong Province, China. The genus is characterized by semi-immersed to superficial, dark brown to black, gregarious ascomata, with elongate necks, 8-spored, unitunicate, cylindrical asci, with a bipartite apical ring and hyaline, fusiform, aseptate to multi-septate ascospores, with or without appendages at one or each end (Liu et al. 2012a). The asexual morph of *Conlarium* was described as hyphomycetous, producing micronematous or semi-macronematous, mononematous, aseptate or septate, hyaline to brown conidiophores, determinate, doliiform, cylindrical, conidiogenous cells and brown, muriform, irregularly globose or subglobose conidia (Liu et al. 2012a, Zhang et al. 2017, Phookamsak et al. 2019, Xie et al. 2019). Currently, *Conlarium* comprises six species viz. *C. aquaticum*, *C. baiseense*, *C. dupliciasmcosporum*, *C. nanningense*, *C. sacchari* and *C. thailandense* X (Zhang et al. 2017, Phookamsak et al. 2019, Xie et al. 2019).
Fig. 122 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU and ITS sequence data. Related sequences are taken from Ekanayaka et al. (2019). Thirty-four strains are included in the combined analyses which comprise 2669 characters (897 characters for LSU, 1031 characters for SSU and 714 characters for ITS) after alignment. *Cyttaria hariotii* (isolate 55) and *C. darwinii* (isolate 14) in Cyttariales are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood values of -12211.970160 is presented. The matrix had 758 distinct alignment patterns, with 44.15% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.247343$, $C = 0.226463$, $G = 0.279975$, $T = 0.246219$; substitution rates $AC = 1.333617$, $AG = 2.238128$, $AT = 1.196767$, $CG = 1.158902$, $CT = 6.126029$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.194594$. Bootstrap values
for maximum likelihood (ML) equal to or greater than 60% and clade credibility values greater than 0.95 (the rounding of values to 2 decimal proportions) from Bayesian-inference analysis labeled on the nodes. The new isolate is indicated in bold and blue.

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**Fig. 123 –** *Apiculospora spartii* (MFLU 19–2559, new host record). a, b Appearance of black conidiomata on the host. c Section of peridium. d, e Vertical sections of conidiomata. f–j Conidiogenous cells and developing conidia. k Germinating conidium, l–o Conidia. p–q Culture on PDA. Scale bars a = 500 µm, b = 200 µm, c = 50 µm, d–e = 100 µm, f–j, l–o = 5 µm, k = 10 µm, p–q = 5 mm.
**Conlarium aquaticum** W. Dong., H. Zhang & K.D. Hyde, Fungal Diversity 85: 75–110 (2017)

Fig. 125

Facesoffungi number: FoF 03337

Saprobic on dead culm of *Thysanolaena maxima*. Sexual morph: Undetermined. Asexual morph: Colonies sporodochial, broadly punctiform, scattered to gregarious, visible as dark brown to black, velvety, shiny colony on the host surface. Mycelium mostly immersed in host substrate, hyaline to light brown, branched, septate, thin- and smooth-walled hyphae. Conidiophores 142–192 × 4.4–5.8 μm (x̄ = 167.1 × 5.1 μm, n = 20), semi-macronematous or macronematous, mononematous, septate, unbranched, straight or flexuous, brown to dark brown. Conidiogenous Cells 15–20 × 3.5–4.9 μm (x̄ = 15.6 × 4.2 μm, n = 20), monoblastic, holoblastic, integrated, determinate, cylindrical, hyaline to pale brown, smooth. Conidia 37–50 × 28–28.5 μm (x̄ = 43.4 × 33.4 μm, n = 20), acrogenous, dictyosporous, muriform, globose to subglobose or irregular in shape, initially hyaline, becoming brown to dark brown when mature, multi-septate, sectored, with small air bubble-like in each cell, slightly constricted at the septa. *Conidia* secession schizolytic.

Known distribution (based on molecular data) – Thailand (Zhang et al. 2017), China (this study).

Known hosts (based on molecular data) – Submerged wood in freshwater (Zhang et al. 2017), *Thysanolaena maxima* (this study).

Material examined – China, Yunnan Province, Xishuangbanna, Mengla County, Xishuangbanna Tropical Botanical Garden (XTBG), on dead stems of *Thysanolaena maxima* (Poaceae), 22 April 2017, R. Phookamsak, IS005 (MFLU 20–0139, new host record), living culture, KUMCC 18–0189.

GenBank numbers – ITS: MN994320, LSU: MN994322, SSU: MN994323, tef1: MT005780 (KUMCC 18–0189A); ITS: MN994328, LSU:MN994330, SSU: MN994331, tef1: MT005779 (KUMCC 18–0189B).

Notes – Our new collection has a similar morphology with the type species of *Conlarium aquaticum* in having brown to dark brown, muriform, multi-septate, sectored conidia (Zhang et al. 2017). However, the conidia of the new collection are slightly smaller than the type (MFLU 20–0139: 37–50 × 28–28.5 μm versus 45–70 × 20–57 μm: MFLU 15–2703; Zhang et al. 2017). Phylogenetic analyses of a combined LSU, SSU and ITS sequence dataset indicated that our strains form a strongly supported subclade with *C. aquaticum* (Fig. 124, MFLUCC 15–0992; 98% ML and 1.00 PP). A comparison of the ITS sequence between these strains reveals less than 1.5% nucleotide base differences, which demonstrate that our new collection is conspecific with *C. aquaticum* (MFLU 15–2703) (Jeewon & Hyde 2016). Therefore, we report our new collection as *C. aquaticum* which is collected from *Thysanolaena maxima* in Yunnan, China for the first time in this study.

**Diaporthales** Nannf.

**Cryphonectriaceae** Gryzenh. & M.J. Wingf.

Cryphonectriaceae was established by Gryzenhout et al. (2006), which comprises mostly phytopathogens, saprobes and endophytes. Recent classification of the family accept 23 genera as *Amphilogia*, *Aurantioporth*, *Aurantiosacculus*, *Aurapex*, *Aurifilum*, *Celoporthe*, *Chromendothia*, *Chrysofalia*, *Chrysoporthe*, *Chrysospherella*, *Cryphonectria*, *Cryptometria*, *Diversimorbus*, *Endothia*, *Foliocryphia*, *Holocryphia*, *Immersiporthe*, *Latruncellus*, *Luteocirrhus*, *Mastigosporella*, *Microthia*, *Rostraureum* and *Ursicollum* (Senanayake et al. 2017, 2018, Wijayawardene et al. 2018).

**Aurifilum** Begoude, Gryzenh. & Jol. Roux

*Aurifilum* is a monotypic genus introduced by Begoude et al. (2010) with *A. marmelostoma* associated with *Terminalia ivorensis* as a pathogen from Cameroon. The genus is characterized by large, pulvinate to pyriform, semi-immersed ascostroma and hyaline, ellipsoidal to fusoid, 1-septate ascospores and broadly convex conidiomata, lack of conidiomatal sack and paraphyses as
compared with other similar genera in Cryphonectriaceae with uniformly orange ascostromata (Begoude et al. 2010). Morphology and phylogenetic analyses using concatenated genes and pathogenicity tests have been confirmed the occurrence of cankers on *Terminalia catappa*, *T. ivorensis* and *T. mantali* in Cameroon and Taiwan (Begoude et al. 2010, Vermeulen et al. 2011, Shen et al. 2016).

Fig. 124 – Phylogram generated from maximum likelihood analysis based on a combined LSU, SSU and ITS sequence dataset. Twenty-one strains are included in the combined gene analyses comprising 2354 total characters including gaps (LSU: 1–872 bp, SSU: 873–1824 bp, ITS: 1825–2354 bp). The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring of the ML tree is selected to represent the phylogenetic relationships of taxa in Conlariaceae and other related families, with the final ML optimization likelihood: -9128.236780. The matrix had 631 distinct alignment patterns, with 21.35% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.258526, C = 0.222031, G = 0.276323, T = 0.243120; substitution rates AC = 1.896853, AG = 2.980974, AT = 1.889217, CG = 0.816766, CT = 7.838961, GT = 1.000000; Tree-Length = 1.051267; gamma distribution shape parameter α = 0.518124. Bayesian posterior probabilities (BYPP) from MCMC were evaluated with a final average standard deviation of split frequencies of 0.007152. Bootstrap support for ML greater than 70% and Bayesian posterior probabilities greater than 0.95 are defined above the nodes as ML/PP. The tree is rooted to *Lentomitella cirrhosa* (ICMP 15131). The type strains are indicated in bold and newly generated sequences are shown in blue.
**Fig. 125** – *Conlarium aquaticum* (MFLU 20-0139, new host record). a Colonies on host. b Squash of mycelium and conidiophores. c, e–m Conidia. d Conidiogenous cell. Scale bars: b = 50 μm, c = 30 μm, d, f–h, j, k = 20 μm, e, i, l = 10 μm.

*Aurifilum marmelostoma* Begoude, Gryzenh. & Jol. Roux, Antonie van Leeuwenhoek 98 (3): 273 (2010)

Facesoffungi number: FoF 04130

Saprobic on dead branches and twigs of *Terminalia ivorensis* (Combretaceae). Sexual morph: *Ascostromata* 520–600 μm high, 540–640 μm diameter, gregarious or single, scattered, medium to
large, usually beneath or erumpent through the bark, semi-immersed, pulvinate to pyriform, yellow to orange, black ostiole, upper region eustromatic, lower region pseudozystromatic, pseudoparenchymatous to prosenchymatous tissue. *Perithecia* 190–330 μm diameter, valsooid, 1–6 per stroma, globose to subglobose, embedded in the stroma at irregular levels, loosely connect with the host. *Peridium* 7.9–13.2 μm (\( \bar{x} = 9.9 \mu m, n = 15 \)) wide, composed of dark brown to reddish brown cells, 4–5 layers of textura angularis to textura epidermoidea. Perithecial necks 140–310 μm (\( \bar{x} = 210 \mu m, n = 10 \)) long, 33.5–57.6 μm (\( \bar{x} = 44.3 \mu m, n = 10 \)) wide, periphysate, black, emerging at stromatal surface as black ostioles, surrounded with orange stromatal tissue to form papillae, composed of with textura porrecta. *Asci* 47.9–51.7 × 8.1–12.3 μm (\( \bar{x} = 50.4 \times 9.8 \mu m, n = 20 \)), 8-spored, fusoid to ellipsoidal, unitunicate with J-, refractive apical rings, short pedicel in immature and sessile at maturity. *Ascospores* 8.3–10.8 × 2.5–4 μm (\( \bar{x} = 9.7 \times 3.3 \mu m, n = 40 \)), L/W 2.9, hyaline, fusoid to ellipsoidal, 1-median septum with tapered apex. Asexual morph: Undetermined.

Culture characteristics – Colonies on PDA fast-growing, reaching 55 mm diameter, after one week at 25–30 °C, medium dense, circular in shape, flattened, surface smooth, with entire edge, velvety to floccose, colony from above greyish white, from below yellowish brown.

Known distribution (based on molecular data) – Cameroon, Kribi, Mbamalyo and Yaounde, Taiwan (Begoude et al. 2010, Vermeulen et al. 2011, Shen et al. 2016), Thailand (this study).

Known hosts (based on molecular data) – *Terminalia catappa*, *T. ivorensis* and *T. mantaly* (Begoude et al. 2010, Vermeulen et al. 2011, Shen et al. 2016), *Terminalia ivorensis* (this study).

Material examined – Thailand, Chiang Rai, Mae Fah Luang University, on dead branches and twigs of *Terminalia ivorensis* (Combretaceae), 9 July 2017, MC. Samarakoon, SAMC082 (MFLU 18–0831, HKAS 102322, new host and geographical record), living culture MFLUCC 18–0536.

GenBank numbers – ITS: MN473057, LSU: MN473051, TUB: MN987000.

Notes – Our strain is similar to the type species, *Aurifilum marmelostoma* in having gregarious or single, yellow ascostromata, the diameter of perithecia overlapping (190–330 μm diameter vs 190–310 μm) and hyaline, fusoid to ellipsoidal, 1-septate ascospores with a tapered base. Our ITS-LSU based phylogenies reveal that our strain clusters with other *A. marmelostoma* strains with high support (Fig. 126, 100%/100%/0.99 PP). In addition, the ITS, LSU and BT sequences are highly similar to those from the type strain (100%, 100% and 99.5%). However, there is no known record of *A. marmelostoma* in Thailand. This is the first record of *Aurifilum marmelostoma* from *Terminalia ivorensis* from Thailand and probably the first host record from Asia. Shen et al. (2016) discussed the pathogenicity of *Aurifilum marmelostoma* on *Terminalia mantaly* and *T. catappa* seedlings.

**Cytosporaceae** Fr.

Senanayake et al. (2017) excluded several genera and accepted only *Cytospora, Pachytrype, Paravalsa, Xenotypa* and *Waydora* within this family. Most members in this family are pathogens that cause various diseases on commercial crops and forest trees (Adams et al. 2005, 2006). Fan et al. (2015a), Jiang et al. (2020) and Shang et al. (2020) have recently introduced several new taxa to this family.

**Cytospora** Ehrenb.

*Cytospora* was introduced by Ehrenberg (1818). Currently, there are more than 100 estimated species in *Cytospora* (Norphanphoun et al. 2018, Species Fungorum 2020, Shang et al. 2020). Rossman et al. (2015) conserved *Cytospora* over *Valsa* giving priority to the older and commonly used name. The asexual morph of *Cytospora* is commonly produced in nature and has fruiting bodies that contain a single or labyrinthine of locules, filamentous conidiophores and allantoid hyaline conidia (Fan et al. 2015a). Some *Cytospora* species are phytopathogenic and form cankers and diebacks (Dar & Rai 2014, Fan et al. 2015a). Some are saprobes and are involved in litter decomposition (Adams et al. 2006).
Fig. 126 – Phylogram generated from maximum likelihood analysis based on combined ITS and LSU sequence data. Eight strains are included in the combined gene analyses comprising 1165 characters after alignment (547 characters for ITS, 618 characters for LSU). *Endothia gyrosa* (CMW 2091) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -2120.791979 is presented. The matrix had 94 distinct alignment patterns, with 8.42% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.240, C = 0.249, G = 0.281, T = 0.230; substitution rates AC = 1.051005, AG = 1.236751, AT = 0.728926, CG = 0.729107, CT = 5.437760, GT = 1.000000; gamma distribution shape parameter $\alpha = 0.759635$. Bootstrap values for maximum parsimony and maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in black bold and newly generated sequence is indicated in blue.

Cytospora cedri Syd., P. Syd. & E.J. Butler, Annls mycol. 14(3/4): 193 (1916)

Facesoffungi number: FoF06871

Saprobic on aerial branches of *Lonicera* sp. Sexual morph: Stromata superficial, ascomata embedded in stromatic tissues, black, irregular to oval, internally brown, pseudoparenchymatous tissues and outer, black, shiny, tightly-packed cells. Ascostromata 350–500 × 150–250 µm ($\bar{x} = 440 \times 200$ µm, n = 20), immersed in stromatic tissue, solitary or aggregated, black to dark brown, coriaceous, uniloculate, with ostiolar neck. Papilla 160–200 × 60–100 µm ($\bar{x} = 180 \times 80$ µm, n = 20), dark brown, internal canal covered with hyaline filamentous hyphae. Peridium comprising several layers of dark brown, thick-walled, cells of textura angularis. Hamathecium comprising cellular, hyaline paraphyses. Asci 25–35 × 5.8–7.1 µm ($\bar{x} = 30 \times 6.7$ µm, n = 20), 8-spored, unitunicate, clavate to elongate obovoid, with a J-, refractive, apical ring. Ascospores 5.5–7.5 × 2–2.5 µm ($\bar{x} = 6.4 \times 2.3$ µm, n = 20), biseriate, hyaline, elongate fusiform to allantoid, 1-celled, smooth-walled. Asexual morph: Undetermined.

Known distribution (based on molecular data) – India (Index Fungorum 2020), Italy (this study).

Known hosts (based on molecular data) – *Cedrus libanus*, *Lonicera* (Sydow et al. 1916), *Lonicera* sp. (this study).

Material examined – Italy, Province of Forlì-Cesena, Modigliana, Trebbio, dead aerial branch of *Lonicera* sp. (Caprifoliaceae), E. Camporesi, 5 April 2015, IT2434 (MFLU 15–1120 new host record).

GenBank numbers – ITS: MN764316, LSU: MN764359.

Notes – According to the ITS blast results, this collection shows a high similarity (99.12%) with *Cytospora fraxinigena*, *C. rosae* (98.77%) and *C. cedri* (99.63%). However, combined gene analysis of ITS, LSU, Actin, and RPB2 in this study, shows that this *Cytospora* collection groups with *Cytospora cedri* (CBS 196.50), which was also collected from Italy. The holotype of *Cytospora cedri* was collected from Himachal Pradesh in India, on a branch of *Cedrus libanus*.
However, our specimen was obtained from a branch of *Lonicera* species in Italy. *Cytospora cedri* is reported on *Lonicera* for the first time, but we could not obtain a culture for this specimen, therefore DNA was extracted directly from fruit-bodies. Our collection is similar to the type described by Sydow et al. (1916). Herein, we report this collection as *Cytospora cedri* based on morphology and phylogeny.

**Fig. 127** – *Aurifilum marmelostoma* (MFLU 18–0831, new host and geographical record). a Host. b–e Ascostromata on the substrate. f Vertical section of ascostroma. g, h Ostiole section. i Peridium. j–n Asci (j–n in Congo Red). o Apical ring (in Congo Red). p–s Ascospores. t Upper view of the colony. u Reverse view of the colony. Scale bars: f = 200 µm, g, h = 100 µm, i–n = 20 µm, p–s = 10 µm, o = 5 µm.
**Cytospora fraxiicola** Chaiwan, Bulgakov, T.C., Jayaward & K.D. Hyde, sp. nov.

Facesoffungi number: FoF06958

Etymology: The specific epithet reflects the host genus *Fraxinus excelsior.*

Holotype: MFLU 17–2392

*Saprobic on branches of Fraxinus excelsior.* Sexual morph: Undetermined. Asexual morph: *Pycnidia* immersed on *Fraxinus excelsior.* *Conidiomata* 600–700 μm high × 1000–1200 μm diameter (x̄ = 650 × 1100 μm, n = 5), scattered, erumpent, multi-loculate. *Locules* composed of numerous interconnecting, chambers arranged radially or irregularly within a continuous, mass of ectostromatic tissue, pycnidial, stromatic, solitary or clustered, immersed in the host when young, semi-erumpent at maturity, dark brown to black, globose, ostiolate. *Ostioles* 360–400 μm high × 380–480 μm diameter (x̄ = 380 × 430 μm). *Peridium* comprising a few to several layers of cells of textura angularis, with inner most layer thin, brown, outer layer dark brown to black. *Conidiophores* branched, reduced to conidiogenous cells. *Conidiogenous cells* enteroblastic, phialidic, formed from the inner most layer of pycnidial wall, hyaline, smooth-walled. *Conidia* 5.8–8.7 × 1.2–2.3 μm (x̄ = 7.25 × 1.75 μm, n = 30), hyaline, elongate-allantoid, 1-celled, slightly curved, smooth-walled.

Material examined – Russia, on dead and dying branches of *Fraxinus excelsior. 21 March 2017, T-2051 MFLU 17–2392 (holotype).

GenBank numbers – LSU: MN764356, ACT: MN995562.

Notes – In the multi-gene analyses, *C. fraxiicola* is distinct and forms a moderately supported lineage clade (Fig. 128). Based on multi-gene analyses, *Cytospora fraxiicola* is distinct from the first group of *C. populina, C. fugax, C. hippophaes, C. gigalocus* and the second group of *C. cotini, C. ampluliformis, C. longispora, C. ribis, C. tanaatica, C. prunicola and C. ulmi.* However, *C. fraxiicola* is similar to *C. gigalocus* in having multi-loculate conidiomata in their asexual morph and has larger conidia (7.25 × 1.75 μm) than *C. gigalocus* (4.8 × 1.1 μm) (Fan et al. 2015a. *Cytospora fraxiicola* differs from *C. populina* and *C. hippophaes* which was reported as a sexual morph.

**Cytospora malicola** Z. Urb., Česká Mykol. 10(4): 209 (1956)

Facesoffungi number: FoF06902

*Saprobic on branches of Salix sp.* Sexual morph: *Stromata* immersed in bark. *Ascostromata* 500–900 × 150–500 μm, immersed in host tissue, scattered, uni or multi-loculate, with ostiolar neck. *Papilla* 350–400 μm, wide at the top than bottom, dark brown to black, internal canal covered with hyaline filamentous hyphae. *Peridium* comprising several layers of cells of textura angularis, with innermost layer thick, pale brown, outer layer dark brown to black. *Hamathecium* comprising long cylindrical, cellular, anastomosed paraphyses. *Asci* 50–65 × 9–11 μm (x̄ = 58 × 10 μm, n = 15), 8-spored, unitunicate, clavate to elongate obovoid, with a J-, refractive, apical ring. *Ascospores* 11–17 × 2.5–3 μm (x̄ = 15.7 × 2.8 μm, n = 20), biseriate to crowded, hyaline, elongate fusiform to allantoid, 1-celled, smooth-walled. Asexual morph: Undetermined.

Known distribution (based on molecular data) – Asia, Europe and North America. (Adams et al. 2006, Mehrabi et al. 2011), Italy (this study).

Known hosts (based on molecular data) – *Malus sylvestris, Prunus domestica, Prunus spinosa, Crataegus oxyacantha* (Hayova and Minter 1998), *Salix sp.* (this study).

Material examined – Italy, Province of Forlì-Cesena, Santa Sofia, Corniolo, dead aerial branch of *Salix* sp. (Salicaceae), E. Camporesi, 16 July 2014, IT2003, MFLU 14–0831, new host record), living culture MFLUCC 14-0831.

GenBank numbers – ITS: MN764317, LSU: MN764358.

Notes – Blast results of ITS of this species show a high similarity (99.31%) with *Cytospora malicola,* and *C. salicina* (98.81%) (Fig. 128). However combined gene analysis of ITS, LSU, Actine, and RPB2 shows that our *Cytospora* species groups with *Cytospora malicola* (Ch92) with a moderate support. Additionally, this subclade is basal to *C. germanica* (CXY1322), *C. kantschavelii* (CXY1383), *C. parakantschavelii* (MFLUCC 15–0857), *C. salicicola* (MFLUCC 14-
Fig. 128 – Phylogram generated from maximum likelihood analysis based on combined ITS, LSU, Actine, and RPB2 sequence data. Bootstrap support for ML greater than 50% and Bayesian posterior probabilities greater than 0.90 are given near nodes. The tree is rooted with Diaporthe vaccinii (CBS 160.32). Ex-type strains are in black bold and the newly generated sequences are indicated in blue bold.
1052) and *C. viticola* (Cyt6). This species was recorded from *Malus sylvestris*, *Prunus domestica*, *Prunus spinosa*, and *Crataegus oxyacantha* with both sexual and asexual morph
Cytospora malicola causes dieback of Malus twigs, especially on twigs frozen, burnt, wounded, injured by insects or attacked by other pathogens (Hayova & Minter 1998). This is the first record of *C. malicola* on *Salix* sp. Morphological characters and measurements of all the characters of our strain are very similar to the protologue described by Hayova & Minter (1998). Therefore, we report this strain as *Cytospora malicola* based on morphology and phylogeny.

**Fig. 129** – *Cytospora cedri* (MFLU 15–1120, new host record). a Ascostromata on substrate. b Cross section of ascoma. c Peridium. d–g Asci. h–j Ascospores. Scale bars: b = 100 μm, c = 50 μm, d–g = 15μm, h–j = 10μm.

*Cytospora melnikii* Norphanphoun, Doilom, Daranagama, Phookamsak, Wen, Bulgakov & Hyde, Mycosphere 8(1): 68 (2017)  
Facesoffungi number: FoF552606

_Saprobic_ on dead branches of *Salix alba*. Sexual morph: Undetermined. Asexual morph: _Pycnidia_ immersed on *Salix alba*. _Conidiomata* 500–600 μm high × 600–700 μm diameter (\(\bar{x} = 550 \times 650\) μm, n = 5), semi-immersed in host tissues, solitary, scattered, erumpent, discoid, circular to ovoid, unilocular, with long ostiolar necks, ostiolate. _Ostioles* 250–300 μm high × 350–400 μm diameter (\(\bar{x} = 275 \times 375\) μm), _Peridium_ comprising a few to several layers of cells of _textura angularis_, with inner most layer thin, brown, outer layer dark brown to black. _Conidiophores_ branched, reduced to conidiogenous cells. _Conidiogenous cells_ enteroblastic, phialidic, formed from the inner most layer of pycnidial wall, hyaline, smooth-walled. _Conidia* 5.5–7.5 ×1.5–2.5 μm (\(\bar{x} = 6.5 \times 2\) μm, n = 30), hyaline, 1-celled, elongate-allantoid, slightly curved, smooth-walled.

Known distribution (based on molecular data) – Russia (Norphanphoun et al. 2017, this study)

Known hosts (based on molecular data) – *Malus domestica* Borkh, _Populus nigra_ var. _italica_ Münchh (Norphanphoun et al. 2017), _Salix alba_ (this study)

Material examined – Russia, on dead and dying branches of *Salix alba* (Salicaceae), 25 July 2017, T-2061 (MFLU 17–2402, new host record).

GenBank numbers – ITS: MN764318, LSU: MN764355, ACT: MN995563, RPB2: MN995564.
Notes – Norphanphoun et al. (2017) introduced *Cytospora melnikii* collected from *Malus domestica* and *Populus nigra* in Russia. In this study, we introduced a new record based on phylogenetic analyses. The morphological characters of our isolate is similar to *C. melnikii* as described by Norphanphoun et al. (2017). It has semi-immersed, larger conidiomata (500–600 vs 420–520 μm) diameter, with larger ostioles (250–300 vs 200–230 μm); and has larger conidia (6.5 × 2 vs 4.6 × 1.2 μm).

Based on multi-gene analyses, our isolate clusters with *C. melnikii* with a high bootstrap support (Fig. 128). Our BLAST search on NCBI GenBank ITS, LSU, ACT and RPB2 sequences show 100% similarity to *C. melnikii*.

**Fig. 130** – *Cytospora fraxiicola* (MFLU 17–2392, holotype). a–c Conidiomata on host substrate. d, e Vertical sections of conidiomata. f–i Conidia. j–n Conidiogenous cells. Scale bars: a = 1000 μm, b, c = 500 μm, d, e = 100 μm, h, i = 50 μm, f, g, j–n = 10 μm.
Diaporthales Nannf.

Diaporthaceae Höhn. ex Wehm.

The most recent taxonomic treatments of Diaporthaceae is by Senanayake et al. (2018).

Diaporthe Nitschke

Diaporthe encompasses species which are endophytic, pathogenic and saprobic on a wide range of hosts. At the same time, single species of Diaporthe can be found on diverse hosts and co-occur on the same host or lesion in different life modes (Dissanayake et al. 2017, Santos et al. 2017). Most recent treatments of this genus are from Wanasinghe et al. (2018b), Marin-Felix et al. (2019) and Manawasinghe et al. (2019).

Diaporthe ambigua Nitschke, Pyrenomyc. Germ. 2: 311 (1870)

Faces of fungi number: FoF 07324

Saprobic on dead aerial stem of Iris sp. Sexual morph Undetermined. Asexual morph: Conidiomata globose, 150–200 μm diameter, mostly embedded in host tissue and erumpent at maturity, up to 450 μm diameter, 65–100 μm high, walls parenchymatous consisting of 3–4 layers of medium brown textura angularis. Conidiophores 7–9 × 1–2 μm, hyaline, smooth, occurring in dense clusters. Conidiogenous cells phialidic, cylindrical, terminal, with slight tapering towards apex, 0.5–1 μm diameter. Conidia 8–9 × 2–3 μm, aseptate, and hyaline, smooth, ovate to ellipsoidal, biguttulate or multi-guttulate.
Culture characteristics: colonies on PDA relatively slow growing reaching 80 mm diameter, after 21 days, On PDA white, fluffy aerial mycelium, and reverse with ash colour.

Known distribution – Chile, South Africa, Portugal, United Kingdom, Netherlands, Armenia, California, Uzbekistan, Canada, Cuba, Germany, China, and Spain (Farr & Rossman 2020), Italy (this study).

Fig. 132 – *Cytospora melnikii* (MFLU 17–2402, new host record). a, b Conidiomata on host substrate. c, d Vertical sections of conidiomata. e–g Conidiogenous cell. h–m Conidia. Scale bars: a = 2000 µm, b = 1000 µm, c–g = 100 µm, h–m = 10 µm.
Fig. 133 – Phylogram generated from maximum likelihood analysis based on combined ITS, TUB, tef1 and CAL sequence data representing the species of Diaporthe. Related sequences are obtained from Manawasinghe et al. (2019). In the combined analyses, 118 strains are included and they comprise 899 characters (558 characters for ITS, 399 characters for TUB, 353 characters for tef1, and 370 characters for and CAL). The tree is rooted by Diaporthella corylina (CBS 121124) The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with final likelihood values of -30526.060484 is presented. The matrix had 1055 distinct alignment patterns, with 15.59% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.221027, C = 0.312667, G = 0.239910, T = 0.226397; substitution rates AC = 1.384376, AG = 3.984787, AT = 1.276868, CG = 0.999850,
CT = 5.491949, GT = 1.000000; gamma distribution shape parameter $\alpha = 0.956001$. The dataset consisted of 1694 characters with 713 constant characters and 758 parsimony-informative and 223 parsimony-uninformative characters. The maximum number of trees generated was 1000, and the most parsimonious trees had a tree length of 5911 (CI = 0.315, RI = 0.710, RC = 0.224, HI = 0.685). Maximum likelihood (ML) and maximum parsimony (MP) equal to or greater than 50% are given as MP/ML on the respective node. Ex-type strains are in bold and black, the new host records are indicated in blue.

Fig. 133 – Continued.
Known hosts (based on molecular data) – *Actinidia deliciosa*, *Aspalathus linearis*, *Diospyros kaki*, *Foeniculum vulgare*, *Helianthus annuus*, *Malus* spp., *Pistacia vera*, *Prunus* spp., *Pyrus* spp., *Vitis vinifera* (Farr & Rossman 2020), *Iris* sp. (this study)

Material examined – Italy, Province of Forlì-Cesena [FC], Pievequinta, on dead aerial stem of *Iris* sp. (Iridaceae), 5 February 2018, E. Camporesii, IT 3695, living culture, JZB320172

GenBank number – ITS: MN989426.

Notes – The collection obtained from the dead aerial stem of *Iris* sp. was identified as *Diaporthe ambigua* with support from morphology and phylogeny. Our isolate clustered with the reference strain of the *Diaporthe ambigua* (JZB320172) in the combined ITS, *tef1*, CAL, HIS and TUB sequence phylogeny with 100% statistical support (Fig. 133). This is the first record of *Diaporthe ambigua* species reported from *Iris* sp. from Italy.

*Diaporthe foeniculina* Udayanga & Castl. Persoonia 32: 95 (2014)  
Faces of fungi number: FoF02183

*Saprobic* on dead aerial stem of *Artemisia vulgaris*. Sexual morph Undetermined. Asexual morph: *Conidiomata* visible as small round to oval dark brown to black dots on the host surface. Sexual morph: not observed. Asexual morph: Coelomycetous. *Conidiomata* 100–200 μm high, 80–200 μm diameter, pycnidial, solitary, scattered or gregarious, globose to subglobose, semi-immersed to immersed, sometimes erumpent, unilocular, thin walled. *Conidioma wall* composed of 3–4-layers, similarly dense at the apex and base, outer 1–2 layers dark brown to black, inner 1–2 layers hyaline, with thin walled cells of *textura angularis*. *Conidiogenous cells* 4–7 μm long × 2–4
µm wide, hyaline, phialidic, globose. **Conidia** 5–10 × 1–4 μm wide (̄x = 8 × 3 μm, n = 20), ellipsoidal to cylindrical, rounded at both ends, hyaline, straight, one septate to aseptate, thin smooth-walled, guttulate.

![Image](image_url)

**Fig. 134** – *Diaporthe ambigua* (JZB320172) a Appearance of conidiomata on the host. b Section through the conidioma. c Conidioma cell wall d Mature conidia.

Culture characters – Colonies on PDA reaching 80 mm diameter, after 14 days at 25 °C, circular colony, white, without any diffusible pigments. Pycnidia globose to subglobose. Conidia subcylindrical, aseptate.

Known distribution (based on molecular data) – New Zealand, Greece, Malta, California, Portugal, Spain, United States, South Africa, Serbia, Uruguay (Farr & Rossman 2020), Italy (this study).

Known hosts (based on molecular data) – Many economically important crops such as *Citrus* spp., *Foeniculum vulgare*, *Glycine max*, *Malus domestica*, *Prunus*, *Vitis* (Farr & Rossman 2020), *Artemisia vulgaris* (this study).

Material examined – Italy, Province of Forli-Cesena [FC], Marsignano – Predappio –Forli – Via Pietro Nenni, on dead aerial stem of *Artemisia vulgaris* (Asteraceae), 21 November 2017, E. Camporesii, IT 3578 (JZBH 320171) living culture, JZB 320171.

GenBank number – ITS: MN989424.

Notes – The collection obtained from the dead aerial stem of *Artemisia vulgaris* was identified as *Diaporthe foeniculina* with support from morphology and phylogeny. Our isolate clustered with the reference strain of the *Diaporthe foeniculina* (CBS 111553) in the combined ITS, tef1, TUB and CAL sequence phylogeny with 99% (ML)/ 100% (MP) statistical support. Furthermore, our isolate showed 99.45% base pair similarity with *Diaporthe foeniculina* in the ITS gene region. *Diaporthe foeniculina* has been reported from many economically important host
plants like *Citrus, Foeniculum vulgare, Glycine max, Malus domestica, Prunus* and *Vitis* worldwide (Farr & Rossman 2020). This is the first record of *Diaporthe foeniculina* on *Artemisia vulgaris* from Italy.

**Fig. 135** – *Diaporthe foeniculina* (JZB 320171, new host record). a Appearance of conidiomata on the host. b Section through the conidioma. c Conidioma wall d Immature conidia attached to conidiogenous cell. e Mature conidia. Scale bars: a = 200 μm, e = 10 μm.

*Diaporthe rumicicola* Manawasinghe, Camporesi & K.D. Hyde, Fungal Diversity [138] (2019)

Facesoffungi number: FoF 04940
Sexual morph: Undetermined. Asexual morph: Conidiomata 181–202 μm high × 203–215 μm diameter (∅ = 189.2 × 207.6 μm, n = 10), eustromatic, convoluted, mostly solitary, semi-immersed to immersed in the host, black, ostiolate. Ostiole 25–33 μm diameter, single, central, with a well-developed neck, thick-walled. Peridium multi-layered 15–20 μm wide at the base, 13–21 μm wide in sides, comprising 6–8 layers, heavily pigmented, thick-walled, textura angularis cells. Conidiogenous cells 8.5–11.7 μm high × 1.3–2.1 μm diameter (∅ = 9.6 × 1.7 μm, n = 10), phialidic, cylindrical, terminal, with slight taper towards apex. Alpha conidia 7.5–10.7× 2.4–3.5 μm (∅ = 9.2 × 2.7 μm, n = 50), aseptate, hyaline, smooth, ellipsoidal or fusiform, with one or two guttules, rarely with subtruncate base. Beta conidia not observed.

Culture characteristics – Colonies on PDA, circular, flattened, fimbriate, crenate edged, both surfaces white, slow growing, and reach 90 mm diameter in 9 days at 28 °C.

Known distribution (based on molecular data) – Italy (Hyde et al. 2019, this study)

Known hosts (based on molecular data) – Rumex sp. (Polygonaceae) (Hyde et al. 2019), Scrophularia canina (this study).

Material examined – Italy, Province of Forlì-Cesena [FC], Cusercoli – Civitella di Romagna, on dead aerial stem of Scrophularia canina (Scrophulariaceae), 7 February 2018, E. Camporesi (MFLU 18–0122, new host record), living culture (MFLUCC 19–0002).

GenBank numbers – ITS: MK066126; tef1: MK078545; TUB: MK078546.

Notes – Based on multi-gene phylogenetic analysis of combined ITS, tef1, β-tubulin (TUB) and calmodulin (CAL) sequence data of Diaporthaceae species (Fig. 133), our strain (MFLUCC 19–0002) clusters with the ex-type strain of Diaporthaceae rumicicola (MFLUCC 18–0739), but differs in alpha conidial length (∅ = 9.2 μm) as compared to the ex-type strain (∅ = 3.5 μm), Diaporthaceae rumicicola (MFLUCC 18–0739).

**Diaporthaceae** Lehman, Ann. Mo. bot. Gdn 10: 128 (1923)  
Facesoffungi number: FoF06582

*Saprobic* or pathogenic on diseased leaves of Rosa multiflora. Sexual morph: Undetermined. Asexual morph: Pycnidia on PDA, superficial, scattered, dark brown to black, globose, solitary in most. Conidiophores were not observed. Conidiogenous cells were not observed. Alpha conidia biguttulate, hyaline, fusiform or oval, both ends obtuse 5–7 × 2–4 (n = 40) μm (∅ = 5.5 × 3 μm). Beta conidia not observed.

Culture characteristics – Colonies on PDA reach 70 mm diameter after 7 days at 25 °C, producing abundant white aerial mycelia and reverse fuscous black.

Known distribution (based on molecular data) – China, Italy, Japan, Mexico, Thailand, and United States (Farr & Rossman 2020), Italy (this study).

Known hosts (based on molecular data – Arachis hypogaea, Arctium lappa, Asparagus officinalis, Aster exilis, Caperonia palustris, Citrus limon, Citrus reticulata, Citrus unshiu, Cucumis melo, Euphorbia nutans, Glycine max, Glycine soja, Phaseolus limensis, Phaseolus vulgaris, Vitis vinifera, Stokesia laevis (Farr & Rossman 2020), Ligustrum quihoui (this study).

Material examined – China, Shandong, Yellow River Park, on leaf spots of Ligustrum quihoui (Oleaceae), 7 October 2017, Y.Y. Hao (living culture JZB320142, new host record).

GenBank numbers – ITS: MN535308, TUB: MN561315.

Notes – In the phylogenetic analysis of combined ITS, TUB, tef1 and CAL gene regions, the taxon obtained in the present study clustered together with the type species, Diaporthaceae sojae (FAU 635) with 99% ML and 88% MP bootstrap support. The taxon is also identical to the type specimen (Udayanga et al. 2015). Diaporthaceae sojae has a wide range of hosts. This species is a well-known pathogen associated with Glycine max in many countries including China. Previously this species was reported from *Vitis* and *Citrus* Spp. in China (Huang et al. 2015, Udayanga et al. 2015). The Ligustrum quihoui is a native plant to Korea and China and grown mainly as an ornamental shrub. This is the first report of Diaporthaceae sojae on *L. quihoui* (Farr & Rossman 2020).
Fig. 136 – *Diaporthe rumicicola* (MFLU 18–0122, new host record). a, b Appearance of conidiomata on host substrate. c Vertical section of a conidioma. d Peridium of conidioma. e Immature and mature conidia attached to conidiogenous cells. f, g Mature conidia. h Culture characters on PDA. Scale bars: c = 50 μm, d = 20 μm, e = 10 μm, f = 20 μm, g = 10 μm.

**Melanconidaceae** G. Winter

Melanconidaceae was introduced by Winter (1886) and is typified by *Melanconis*. The family is characterized by circularly arranged perithecia immersed in stromata with a central column and ostioles erumpent through an ectostromatic disc with hyaline, 1-septate ascospores; pycnidia developing before the formation of the ascomata and produce 1-celled, dark-brown conidia (Barr
1978, Castlebury et al. 2002). Senanayake et al. (2017) restricted Melanconidaceae for *Melanconis*. Only the type genus, *Melanconis* is accommodated in the family. Species of Melanconidaceae are usually pathogens or endophytes on hardwood trees in Betulaceae. They are distributed worldwide especially in China.

**Fig. 137** – *Diaporthe sojae* (JZBH320142, new host record). a Material examined. b Appearance of conidia on PDA. c conidiogenus cell. d–e Conidia f Upper view of colony on PDA. g reverse view of colony on PDA. Scale bars: c–e = 20 μm.
Fig. 138 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS, RPB2 and TUB sequence data. Twenty-two strains are included in the combined gene analyses comprising 2747 characters after alignment (891 characters for LSU, 572 characters for ITS, 1126 characters for RPB2 and 425 characters for TUB). *Gnomonia gnomon* (CBS 199.53) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of \(-6277.206694\) is presented. The matrix had 263 distinct alignment patterns, with 17.80\% undetermined characters or gaps. Estimated base frequencies were as follows: \(A = 0.244490, C = 0.261036, G = 0.265173, T = 0.229301\); substitution rates \(AC = 1.367272, AG = 2.659677, AT = 1.874405, CG = 1.219603, CT = 7.861355, GT = 1.000000\); gamma distribution shape parameter \(\alpha = 0.738642\). Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.

*Melanconis* Tul. & C. Tul.

*Melanconis* was introduced by Tulasne & Tulasne (1863) with *Sphaeria stilbostoma* Fr. as the type species and an asexual state placed in *Melanconium* Link. *Melanconis* is characterized by circularly arranged perithecia, immersed in stromata with a central column and ostioles which arise through an ectostromatic disc with hyaline, 1-septate ascospores. The asexual morph is coelomycetous often with brown aseptate conidia. There are around 100 species epithets in *Melanconis*. Some of them been transferred to other families such as Amphiphasphaeriaceae, Cucurbitariaceae, Gnomoniaceae, Pseudovalsaceae and Sydowiellaceae (Index Fungorum 2020).
*Melanconis stilbostoma* (Fr.) Tul. & C. Tul., Select. fung. carpol. (Paris) 2: 115 (1863)

Pathogenic on stems of *Betula pendula* Roth. Sexual morph: Undetermined. Asexual morph: **Conidiomatal stromata** 134–172 × 100–159 μm (x̄ = 148.2 × 128.5 μm, n = 10) immersed or superficial in host bark, conical, black, circular to ellipsoid with an ectostromatic disc white to pale yellow, surrounded by bark. **Central column** beneath the disc more or less conical. **Conidiophores** 13–17 × 1.2–1.9 μm (x̄ = 17.2 × 1.6 μm, n = 20) branched at the base, elongate, hyaline to subhyaline, smooth. **Conidiogenous cells** monophialidic, integrated. **Conidia** 10–12 × 6–7 μm (x̄ = 11.1 × 7.2 μm, n = 20), ovoid, acute at one end and obtuse at the other, brown to olive brown, aseptate.

![Fig 139](image-url)
Culture characteristics – Colony on MEA white at first, producing creamy white to pale yellowish brown pigment after 15 days, felty, with an irregular edge; conidiomata sparse.

Material examined – Ukraine, Donetsk region, Donetsk City, Donetsk Botanical Garden, arboretum, on Betula pendula Roth (Betulaceae), 16 May 2017, Timur Bulgakov (MFLU 17–2444, new geographical record); living cultures; MFLUCC 18–0783, DSM 109785.

Known distribution (based on molecular data) – Sweden, China (Fan et al. 2016), Russia (this study).

Known hosts (based on molecular data) – Betula spp. (B. pendula, B. rotundifolia and B. tianschanica) (Fan et al. 2016), Betula pendula (this study).

GenBank Numbers – LSU: MN244205, SSU: MN244182, ITS: MN244222.

Notes – Melanconis stilbostoma is the type species of Melanconis and mainly distributed on Betula spp. Our isolate MFLUCC 18–0783 clustered with the strains of Melanconis stilbostoma (Fig. 138), which has been reported only in Sweden and China. We, therefore report our collection as a new geographical record from Russia based on morphology and phylogenetic analyses.

Schizoparmaceae Rossman

Schizoparmaceae was introduced by Rossman et al. (2007) to accommodate Schizoparmella with its asexual state Pilidella and Coniella in Diaporthales. Members of this family are pathogens that cause diseases in grapes, strawberry and other plants (Rossman et al. 2007) and saprobes in plants and soil (Alvarez et al. 2016).

Coniella Höhn.

The asexual genus Coniella was established by Von Höhnel (1918) and is typified by C. pulchella (= C. fragariae; Crous et al. 2014). Species of Coniella are saprobes, plant pathogens and endophytes (Alvarez et al. 2016). They have a wide host range i.e., Eucalyptus, Fragaria, Hibiscus, Psidium, Punica, Terminalia and Vitis occurring on leaf litter, rotting bark, and soil (Alvarez et al. 2016).

Coniella eucalyptorum (Crous & M.J. Wingf.) L.V. Alvarez & Crous in Alvarez, Groenewald & Crous, Stud. Mycol. 85: 15 (2016) Fig. 141

Facesoffungi number: FoF07370

Saprobic on dead leaves of undetermined host. Sexual morph: Undetermined. Asexual morph: Conidiomata 90–130 μm high × 160–200 μm in diameter (n = 10) solitary, globose, brown and dark brown from the top. Conidioma wall consisting of 2–3 layers of hyaline textura prismatica and 4-5 layers of brown textura angularis cells. Conidiophores densely aggregated, branched. Conidiogenous cells 12–15 × 1–2 μm, annellidic, narrowing at the tip, smooth and hyaline. Conidia 10–12 × 3–6 μm (x̄ = 11 × 5 μm, n = 30), hyaline to pale brown, becoming dark brown at maturity, smooth, broadly ellipsoidal, both sides gradually tapering, smooth-walled, and multi-guttulate with one or two prominent guttules.

Culture characteristics – Colony on PDA reaching 30 mm diameter after 7 days at 25 °C, colony circular, margin wavy, flat, velvety appearance, colony from above: light brown; reverse: dark brown.

Known distribution (based on molecular data) – Australia, Brazil, Chile, China, Indonesia, Malaysia, Mexico, Venezuela, Vietnam (Alvarez et al. 2016), Thailand (this study).

Known hosts (based on molecular data) – Eucalyptus sp., Corymbia nesophila, Corymbia torelliana (Alvarez et al. 2016), unidentified host (this study).

Material examined – Thailand, Chiang Mai Province, Mae Taeng, Ban Pa Deng, Mushroom Research Centre, on twigs of unidentified host, 27 May 2016, N.I de Silva, NI106 (MFLU 17–0675), living culture, MFLUCC 17–0870.

GenBank Numbers – ITS: MN836684, LSU: MN836664

Notes – Phylogenetic analysis of combined ITS, LSU, histone and tefl sequence data confirmed that our collection is Coniella eucalyptorum with high support (Fig. 140). Coniella
was previously recorded from Australia, Brazil, Chile, Indonesia, Malaysia, Mexico, Venezuela and Vietnam (Alvarez et al. 2016). Coniella eucalyptorum was commonly recorded from Eucalyptus sp. (Alvarez et al. 2016). This is the first record of C. eucalyptorum from twigs of unidentified host in Thailand.

**Fig. 140** – Phylogram generated from the maximum likelihood analysis based on combined ITS, LSU, histone and tef1 sequence data representing genus Coniella. Related sequences are taken from Chethana et al. (2017). Forty three strains are included in the combined analyses which comprise 2866 characters (586 characters for ITS, 1180 characters for LSU, 485 characters for histone, 615 characters for tef1) after alignment. Melanconiella sp. (CBS110385) is used as the outgroup taxon. The best RaxML tree with a final likelihood values of -15179.467302 is presented. The matrix had
796 distinct alignment patterns, with 23.91% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.249525$, $C = 0.245214$, $G = 0.256307$, $T = 0.248954$; substitution rates $AC = 1.240352$, $AG = 2.275249$, $AT = 1.362790$, $CG = 0.918763$, $CT = 5.013442$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.398515$. Bootstrap values for ML equal to or greater than 50% (first set) and MP equal to or greater than 50% (second set) are given above the nodes. Ex-type strains are in black bold and new strain is indicated in blue bold.

**Fig. 141** – *Coniella eucalyptorum* (MFLU 17–0675, new geographical record). a–c Appearance of conidiomata on dead branch. d Vertical section of conidioma. e Peridium. f, g Conidiophores. h Conidia. i Germinating conidia. j Upper view of culture. k Lower view of culture. Scale bars: d = 50 $\mu$m, e–h = 10 $\mu$m, i = 20 $\mu$m.

**Subclass Hypocreomycetidae** O.E. Erikss. & Winka

**Glomerellales** Chadef.

**Glomerellaceae** Locq ex Seifert & W Gams.

Glomerellaceae is a monotypic family mainly comprised of pathogens. This family is characterized by *Colletotrichum* (asexual morph) and *Glomerella* (sexual morph), which was synonymized under *Colletotrichum* (Hyde et al. 2014, Maharachchikumbura et al. 2016a).

**Colletotrichum** Corda

This genus was introduced by Corda (1831), for *C. lineola* Corda (Hyde et al. 2009). *Colletotrichum* comprises mainly pathogens, as well as endophytes and saprotrophs (Hyde et al. 2014, Jayawardena et al. 2016, Samarakoon et al. 2018). Kirk et al. (2001, 2008b) and Réblová et
al. (2011) placed *Colletotrichum* in Glomerellaceae and the study of Maharachchikumbura et al. (2016a) further confirmed the placement of this genus.

**Fig. 142** – Phylogenetic tree generated by maximum parsimony analysis of combined ITS, GAPDH, CHS, ACT and TUB sequence data of *Colletotrichum* species. Forty-five strains are included in the analyses, which comprise 1876 characters including gaps. The tree is rooted with *Colletotrichum boninense* (CBS 123755). The maximum parsimonious dataset consisted of 1293 constant, 268 parsimony-informative and 315 parsimony-uninformative characters. The parsimony analysis of the data matrix resulted in the maximum of ten equally most parsimonious trees with a length of 1945 steps (CI = 0.708, RI = 0.764, RC = 0.541, HI = 0.292) in the first tree. MP, ML bootstrap values ≥50% and Bayesian posterior ≥0.90 probability are shown near the nodes. The scale bar indicates 10 changes per site. The ex-type strains are in bold. Newly generated sequences are indicated in blue bold.

*Colletotrichum fructicola* Prihast., L. Cai & K.D. Hyde, Fungal Diversity 39: 96 (2009)

Facesoffungi number: FoF0767

*Saprotrophic* on dead leaves of *Rosa hybrida*. Sexual morph: Undetermined. Asexual morph: *Conidiomata* absent on culture. *Setae* numerous, base pale to dark grey, with scattered, dark brown
setae about 50–80 µm long. Conidiophores branched at the base, elongate, hyaline, smooth. Conidiogenous cells enteroblastic, phialidic, integrated, smooth, with minute collarette at the tip. Conidia $11–14 \times 3–5$ µm $\bar{x} = 12 \times 4$ µm, $n = 40$ hyaline, smooth or verruculose, aseptate, guttulate, fusiform, cylindrical with obtuse to slightly rounded ends. Appressoria not observed.

Culture characteristics – Colonies on PDA reaching a maximum of 80 mm diameter in 7 days at 25 ºC, at first white and becoming pale brownish to pinkish, reverse pale yellowish to pinkish, aerial mycelium greyish white, dense, cottony. Vegetative hyphae 2.1 µm hyaline, smooth-walled, septate, branched.

Known distribution (based on molecular data) – Brazil, China, Japan, Korea, Puerto Rico, Thailand (Bhunjun et al. 2019, Cao et al. 2019, Costa et al. 2019, Hirayama et al. 2018, Veloso et al. 2018, Jayawardena et al. 2016, this study)

Fig 143 – Colletotrichum fructicola (MFLU 18–1837, new host record). a Setae. b–d Conidiogenous cells. e–h Conidia. i Germinating conidium. j Upper view of 7 day old culture. k Reverse view of 7 day old culture. Scale bars: a = 20µm, b–i = 5µm.
Known hosts: based on molecular data (– Amaranthus blitum, Anacardium sp., Annona sp., Artocarpus sp., Camellia sp., Capsicum sp., Citrus sp., Coffea sp., Malus sp., Mangifera sp., Nephelium lappaceum, Pyrus sp., Vitis sp.) Bhunjun et al. 2019, Cao et al. 2019, Costa et al. 2019, Hirayama et al. 2018, Jayawardena et al. 2016, Veloso et al. 2018, Li et al. 2016b (Rosa hybrida) this study.

Material examined: Thailand, Chiang Rai, on dead leaves of Rosa hybrida (Rosaceae), 15 May 2018, Ruvishika S. Jayawardena RKB5 (MFLU) 18–1837, new host record (living culture MFLUCC 18–1160).

GenBank Numbers: – ITS: MN788675, GAPDH: MN995327, CHS: MN995335, ACT: MN995333

Notes: Colletotrichum fructicola has a wide host and geographical range as well as a wide distribution and is a well-known pathogen on many crops (Jayawardena et al. 2016). A strain in this study clustered with the type species of C. fructicola (Fig. 142). This is the first record of C. fructicola as a saprotroph on Rosa hybrida in Thailand.

Colletotrichum siamense Prihast., L. Cai & K.D. Hyde, Fungal Diversity 39: 98 (2009)

Facesoffungi number: FoF 03599

Saprotrophic on dead leaves of Rosa sp. Sexual morph: Undetermined. Asexual morph: Conidiomata 70–143 μm diameter, brown to dark brown, acervulate, oval, solitary to aggregated. Setae absent. Conidiophores hyaline, cylindrical or clavate, smooth-walled, simple, to 80 μm long. Conidiogenous cells 10–14 μm long, enteroblastic, phialidic, hyaline, smooth-walled, cylindrical to slightly inflated, Collarette 0.5–1 μm long, periclinal thickening visible. Conidia 7–19 × 3–4 μm. x̄ = 11 × 4 μm, n = 40 (hyaline, smooth or verruculose, aseptate, guttulate, fusiform, ovoid to cylindrical or clavate with rounded apices. Appressoria not observed.

Culture characteristics: – Colonies on PDA reaching a maximum of 80 mm diameter in 7 days at 25 °C, at first white and becoming pale brownish to pinkish, reverse pale yellowish to pinkish, aerial mycelium greyish white, dense, cottony. Vegetative hyphae 2.1 μm hyaline, smooth-walled, septate, branched.

Known distribution: based on molecular data (– Australia, Brazil, China, Japan, Korea, Nigeria, Puerto Rico, South Africa, Thailand, USA, Vietnam) Yang et al. 2009, Weir et al. 2012, Jayawardena et al. 2016, Shivas et al. 2016, this study.

Known hosts: based on molecular data (– Artocarpus sp., Camellia sp., Capsicum sp., Carica papaya, Citrus sp., Coffea sp., Dioscorea rotundata, Fragaria × ananassa, Hymenocallis sp., Jasminum sambac, Malus sp., Mangifera sp., Manihot sp., Persea americana, Pistacia vera, Pyrus sp., Vitis sp.) Bhunjun et al. 2019, Cao et al. 2019, Costa et al. 2019, Hirayama et al. 2018, Veloso et al. 2018, Li et al. 2016b (Rosa sp). This study.

Material examined: Thailand, Chiang Rai, on dead leaves of Rosa sp. (Rosaceae), 19 May 2018, Ruvishika S. Jayawardena RV1 (MFLU) 18–1839, new host record (living culture MFLUCC 18–1162).

GenBank Numbers: – ITS: MN788676, GAPDH: MN995328, CHS: MN995336, ACT: MN995334, TUB: MN995329

Notes: Colletotrichum siamense has a wide host range and a distribution (Jayawardena et al. 2016). A strain in this study clustered with the type species of C. siamense (Fig. 142). This is the first record of C. siamense as a saprotroph of Rosa sp. in Thailand.

Colletotrichum viniferum Li J. Peng, L. Cai, K.D. Hyde & Zi Y. Ying, Mycoscience 54(1): 36 (2013)

Facesoffungi number: FoF 03600

Endophytic and saprotrophic on leaves of Hemerocallis and Mangifera indica. Sexual morph: Undetermined. Asexual morph: Conidiomata 48–143 μm diameter, black, acervulus, oval, solitary to aggregated. Setae absent. Conidiophores hyaline to light brown, cylindrical or clavate,
smooth-walled, simple, wide at the base, to 46 µm long, occurring in densely arranged clusters. *Conidiogenous cells* enteroblastic, phialidic, hyaline, smooth-walled, cylindrical to slightly inflated, collarette 0.5–1 µm long, periclinal thickening visible, *Conidia* 7–12 × 2–6 µm $\bar{x} = 9 \times 4$ µm, n = 40, hyaline, smooth or verruculose, aseptate, ovoid to cylindrical or clavate with rounded apices. *Appressoria* formed in culture 9–16 × 5–6 µm $\bar{x} = 13 \times 5$ µm, n = 10, solitary to aggregated, in small groups or short chains, medium to dark brown, smooth-walled, round, oval or irregular.

Fig 144 – *Colletotrichum siamense* (MFLU 18–1839, new host record). a Conidiomata on host. b Conidiophores and conidiogenous cells. c–d Conidia. e Upper view of 7 day old culture. f Reverse view of 7 day old culture. Scale bars: b–d = 10µm.
Culture characteristics – Colonies on PDA reaching 75 mm diameter in 7 day at 25 °C, at first white becoming dark grey, reverse pale yellowish, aerial mycelium greyish white, dense, cottony, vegetative hyphae 2.1 µm hyaline, smooth-walled, septate, branched.

Known distribution (based on molecular data) – China, South Korea) Peng et al. 2013, Jayawardena et al. 2016; (Russia, Thailand) this study.

Known hosts (based on molecular data) – Capsicum sp., Vitis sp. Peng et al. 2013, Jayawardena et al. 2016, Oo and Oh 2017, Hemerocallis sp., Mangifera indica this study.

Material examined – Russia, Rostov region, Shakhty City District, street flowerbed, on dead floriferous shoot of Hemerocallis sp. (Asphodelaceae), 20 November 2017, T. Bulgakov T2281-1) MFLU 18–0441, new host and geographical record, culture, MFLUCC 18–0748; Thailand, Chiang Rai, on leaves of Mangifera indica, 19 May 2018, Ruvishika S. Jayawardena MRV3) MFLU 18–1850, new host and geographical record, culture, MFLUCC 18–1179.

GenBank Numbers – (MFLUCC 18–0748); ITS:MN788677, TUB:MN995330 (MFLUCC 18–1179); ITS:MN788678, TUB:MN995331

Fig. 145 – Colletotrichum viniferum (MFLU 18–1850, new host and geographical record). a Conidiogenous cells. b Mature conidia. c Germinating spore. d–e Appressoria. Scale bars: a–e = 5 µm.

Notes – Colletotrichum viniferum was introduced by Peng et al. 2013. This species is a known pathogen causing grape ripe rot in China) Yan et al. 2015. Our strains clustered with the
type species of *C. viniferum* with high bootstrap support) Fig. 142. *Colletotrichum asianum* and *C. gloeosporioides* have been previously recorded from *Mangifera* spp. in Thailand (Weir et al. 2012, Jayawardena et al. 2016). This study provides both the first report of *C. viniferum* as a saprotroph from *Hemerocallis* sp. from Russia and as an endophyte from *Mangifera indica* in Thailand.

**Hypocreales** Lindau
**Cordycipitaceae** Kreisel ex G.H. Sung et al.

*Cordycipitaceae* was introduced by Sung et al. (2007) based on multi-gene analyses to accommodate 11 genera. Species in this family are mostly pathogenic on insects, spiders, nematodes and rust fungi, saprobic on decaying wood or occur on soil (Sung et al. 2007, Evans 2013, Nonaka et al. 2013, Mijeon et al. 2015, Zhang et al. 2017, Huang et al. 2018, Wei et al. 2018). Maharachchikumbura et al. (2016a) accepted 18 genera and provided family and generic descriptions for *Cordycipitaceae* and *Cordyceps*. Kepler et al. (2017) accepted 11 genera and rejected eight generic names in this family based on phylogenetic analyses. Wijayawardene et al. (2018) listed 17 genera while rejecting some generic names in Kepler et al. (2017), such as *Granulomanus*, *Isaria* and *Microhilum*. Previous research failed to account for *Leptobacillium* (Zare & Gams 2016) and *Samsoniella* (Mongkolsamrit et al. 2018). Up to now, there are many old species in *Cordycipitaceae* with several genera without verification with molecular data, such as *Beejasamuha*, *Coremiopsis* and *Granulomanus*.

**Cordyceps** Fr.

*Cordyceps* was established by Link (1833) based on the type species *Cordyceps militaris*. The representative characters of this genus are fleshy, pallid or brightly pigmented stromata, superficial to completely immersed ascomata, cylindrical asci with a thickened cap and hyaline, multi-septate, disarticulating into part-spores or non-disarticulating ascospores (Sung et al. 2007, Maharachchikumbura et al. 2016a).

**Cordyceps pruinosa** Petch, Trans. Br. mycol. Soc. 10(1-2): 38 (1924) Fig. 147

Facesoffungi number: FoF07468

*Parasitic* on limacodid pupa which resemble a plant seed. Sexual morph: Undetermined. Asexual morph: Hyphomycetous. *Stroma* emerging from host, single, erect, cylindrical, apically branched, red, pale yellow at the base, fleshy, with white straight hyphae and interlaced vegetative hyphae mass occurring on the surface of upper stroma. *In vitro. Synnemata* orange, erect, compacted below, apically branched, becoming finger-like. *Hyphae* 0.6–1.20 (x̄ = 1, n = 20) μm, aseptate, hyaline, smooth-walled. *Conidiophores* macronematous, prostrate, branched, bearing solitary or 2–3 whorls of phialides. *Conidiogenous cells* phialidic. *Phialides* 11–30 × 1.2–1.6 (x̄ = 16 × 1.3, n = 20) μm, aseptate, monophialidic, gradually tapering towards the apex, hyaline, straight. *Conidia* 1.8–4.8 × 0.9–2.7 (x̄ = 3 × 1.5, n = 30) μm, globose, oval, cylindrical, aseptate, smooth-walled, hyaline, aggregating in imbricated slimy chains at the tip of phialides.

Culture characters – slow-growing on PDA, reaching 4 cm in diameter after incubating at room temperature for 26 days, yellow from upper and reverse view, floccose, dense, circular, feathery margin, with several erect, orange synnemata developing at central regions, without diffused pigment. The sporulation occurs on upper part of synnemata.

Material examined – China, Yunnan Province, Kunming City, Western hill Park. On Limacodid pupa, 27 July 2018, Deping Wei (HKAS 102537, new sequence data), living culture KUMCC 18–0340.

Known distribution (based on molecular data) – Korea (Bae et al. 2002, Moon et al. 2018); Guangdong and Guizhou in China (Wang et al. 2008, Meng et al. 2014, Zha et al. 2019); Japan (Nikoh & Fukatsu 2001, Kepler et al. 2012).

Known hosts (based on molecular data) – Cocoon of a Limacodidae insect (Meng et al. 2014, Zha et al. 2019); *Iragoides fasciata* (Lepidoptera) (Spatafora et al. 2007, Sung et al. 2007).

GenBank numbers – ITS: MT012347, LSU: MT012354, SSU: MT012361, tef1:MT025053.
Notes – In the phylogenetic tree, our collection XS2703 is closest to *Cordyceps pruinosa* (ARSEF 5413) and *C. ninchuckispora* with moderate support (81% MP/ 0.99 PP). The strain ARSEF 5413 is not linked to any description (Kepler et al. 2017). *Cordyceps pruinosa* was initially described by Petch (1924) with its asexual morph linked to *Mariannaeas pruinosa* by Liang et al. (1991) but not confirmed with molecular data. The asexual morph of *Cordyceps pruinosa* was characterized by verticillate conidiophores and ovoid, ellipsoid, cylindrical, rarely septate, hyaline conidia, which adhere in imbricate chains (Liang et al. 1991). Our collection is consistent with the concept of *Cordyceps pruinosa* in the shape and size, as well as the arrangement of conidia. A comparison of *tef1* sequences between our isolates and two strains of *Cordyceps ninchuckispora* (EFCC 5197 and EFCC 5693) shows 17/862 bp (2%) differences, but identical to *Cordyceps pruinosa* (ARSEF 5413) within 862bp. In ITS, our collection is identical to *Cordyceps pruinosa* (ARSEF 5413) within 355bp. We report a new collection of *Cordyceps pruinosa* according to the guidelines of Jeewon & Hyde (2016).

*Cordyceps rosea* Kobayasi & Shimizu, Bull. natn. Sci. Mus., Tokyo, B 8(4): 112 (1982)

Fig. 148

Facesoffungi number: FoF07467

*Parasitic on Lepidoptera* larva or pupa. Sexual morph: Stroma arising from insect body, stipitate, multiple or single, flexuous or straight, orange, fleshy, unbranched. *Perithecia* 329–846 × 343–667 (\(\bar{x} = 555 \times 486, n = 20\)) µm, superficial, born on upper region of stroma, ovoid. *Peridium* 22–49 (\(\bar{x} = 34, n = 20\)) µm wide, membranous, comprised of yellow, thick-walled cells of *textura angularis*. *Asci* 212–595 × 2.5–5.5 (\(\bar{x} = 395 \times 4, n = 30\)) µm, 8-spored, unistomate, narrowly cylindrical, with round, attenuate base and thickened, hemispherical cap. *Ascospores* 2–5.5 × 0.7–1.5 (\(\bar{x} = 3 \times 1, n = 100\)) µm, filiform, hyaline, disarticulating into secondary spores and spraying from perithecia when mature. *Secondary ascospores* cylindrical, hyaline, smooth-walled, aseptate, truncated at both ends, straight, occasionally swollen spores present. Asexual morph: Undetermined.

Culture characteristics – *Colonies* growing on PDA, circular, slightly raised, edge entire, velvety, reaching 18 mm in 16 days at 25 °C, white to pale yellow in PDA medium.

Material examined – China, Yunnan Province, Honghe County, Jiayin Village, Amushan protected area, on *Lepidoptera* pupa, 23 October 2018, De-Ping Wei, AMS10 (HKAS 102492, reference specimen designated here), ibid; AMS06 (HKAS 102495), TSQ09B (HKAS 102493), TSQ09C (HKAS 102494), TSQ09E (HKAS 102496); KUMCC 20-0002, living culture.

Known distribution (based on molecular data) – Taiwan (Kepler et al. 2017).

Known hosts (based on molecular data) – Lepidopteran larva (Kepler et al. 2017).

GenBank numbers – ITS: TSQ09B = MT012342, TSQ09C = MT012343, TSQ09E = MT012344, AMS06 = MT012345, AMS10 = MT012346; LSU: TSQ09B = MT012349, TSQ09C = MT012350, TSQ09E = MT012351, AMS06 = MT012352, AMS10 = MT012353; SSU: TSQ09B = MT012356, TSQ09C = MT012357, TSQ09E = MT012358, AMS06 = MT012359, AMS10 = MT012360; *tef1*: TSQ09B = MT025048, TSQ09C = MT025049, TSQ09E = MT025050, AMS06 = MT025051, AMS10 = MT025052; *RPB2*: TSQ09B = MT025044, TSQ09C = MT025045, TSQ09E = MT025046.

Notes – *Cordyceps rosea* was initially discovered on larva of *Lepidoptera* in Honshu Province, Japan and was introduced by Kobayasi & Shimizu (1982). Its macro-morphology and molecular data were made available in Kepler et al. (2017), but the details of micro-morphology are still lacking. Our isolates resemble the reference specimen in having fleshy, orange stroma and superficial, conical perithecia. In addition, the multi-locus based phylogenetic tree shows that our isolates form a clade sister to *Cordyceps rosea* (spat 09-053) with high support (Fig. 146, 99% ML/0.99 PP). The nucleotide differences between our isolates and *Cordyceps rosea* (spat 09-053) are lower than 1.5% for LSU and SSU sequences, and higher for *tef1* sequences (Table 1). Unfortunately, ITS sequences of *Cordyceps rosea* (spat 09-053) are not available for comparison. The ITS nucleotide sequences of our isolates TSQ09B, TSQ09C, TSQ09E, AMS10 are identical,
but differ from AMS06 in two gaps and one base pair. To conclude, there are many differences in the tef1 sequences, but the subtle differences in the ITS sequence and phylogenetic analysis as well as the macro-morphologic features support our isolates to be a new geographical record of *Cordyceps rosea* in mainland China. Given that the molecular data and morphological descriptions of *Cordyceps rosea* were untraceable, we propose a reference specimen for this species with support from phylogeny and morphological data.

Fig 146 – Phylogram generated from maximum likelihood analysis based on combined LSU, SSU, ITS and tef1 sequence data. Sixty-one strains are included in the combined gene analyses comprising 3020 characters after alignment (802 characters for LSU, 980 characters for SSU, 375 characters for ITS, 862 characters for tef1). *Beauveria bassiana* (ARSEF 1564) and *Beauveria bronniartii* (BCC 16585) are used as the outgroup taxa. The tree topology derived from the
Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of \(-12920.722667\) is presented. The matrix had 870 distinct alignment patterns, with 20.99% undetermined characters or gaps. Estimated base frequencies were as follows: \(A = 0.242336\), \(C = 0.262549\), \(G = 0.268942\), \(T = 0.226174\); substitution rates \(AC = 1.424390\), \(AG = 2.347648\), \(AT = 1.121613\), \(CG = 0.997130\), \(CT = 7.134373\), \(GT = 1.000000\); gamma distribution shape parameter \(\alpha = 0.699986\). Bootstrap values for maximum likelihood equal to or greater than 70 and Bayesian posterior probabilities equal or greater than 0.90 are placed above or below the branches. The newly generated sequences are indicated in red bold.

![Image of Cordyceps pruinosa](image.jpg)

Fig 147 – *Cordyceps pruinosa* (HKAS 102537, new sequence data). a Stroma emerging from host. b, f White hypha produced on stroma. c, d Upper and reverse view of cultures on PDA after 34 days incubation. e Vertical section of stroma. g Interlaced hypha occurring on upper region of stroma. h, j Conidiophore. k, l Phialides. i, m–o Conidia adhering in imbricated chains. Scale Bars:
Microascales Luttr. ex Benny & Kimbr

Microascaceae Luttr. ex Malloch

Luttrell (1951) proposed Microascaceae and Microascales and they were validly published with Latin descriptions by Malloch (1970), Benny & Kimbrough (1980). Malloch (1970) reviewed the family with sexual and asexual characteristics, while Réblová et al. (2011) and Maharachchikumbura et al. (2015) investigated the phylogenetic placement of Microascaceae. Currently, this family includes 23 genera (Wijayawardene et al. 2020). Our marine fungal collection from the UK yielded *Parascedosporium putredinis* from submerged marine wood and this is the first record of *P. putredinis* from a marine habitat.

**Table 1** A comparison of nucleotide sequences between *Cordyceps rosea* (spat 09-053) and our isolates.

| Species  | LSU (bp) | SSU (bp) | tefl (bp) |
|----------|----------|----------|-----------|
| TSQ09B   | 3/796 (0.37%) | 1/907 (0.1%) | 24/834 (2.87%) |
| TSQ09C   | 4/796 (0.5%) | 1/907 (0.1%) | 26/834 (3.11%) |
| TSQ09E   | 3/796 (0.37%) | 1/907 (0.1%) | 24/834 (2.87%) |
| AMS06    | 3/796 (0.37%) | 1/907 (0.1%) | 26/834 (3.11%) |
| AMS10    | 3/796 (0.37%) | 1/907 (0.1%) | 26/834 (3.11%) |

*Parascedosporium* Gilgado, Gené, Cano & Guarro

*Parascedosporium* was introduced by Gilgado et al. (2007) based on *Graphium tectonae* (CBS 127.84) isolated from seeds of *Tectona grandis* in Jamaica (Lackner & de Hoog 2011). *Graphium tectonae* was synonymized as *Parascedosporium tectonae* by Gilgado et al. (2007). Subsequently, this species was considered as a synonym of *Parascedosporium putredinis* by Lackner & de Hoog (2011) based on the analysis of ITS sequence data and examination of the ex-type culture (CBS 127.84). This treatment was accepted by de Beer et al. (2013). Sympodial conidia produced from denticulate conidiogenous cells are the unique features of this genus (Gilgado et al. 2007). *Parascedosporium* synanamorphs has solitary conidiophores that emerge from aerial mycelium and comprises of graphium-like synnemata (Gilgado et al. 2007). The sexual morphs of this genus are undetermined.

*Parascedosporium putredinis* (Corda) Lackner & de Hoog

[Fig. 150](#)

Facesoffungi Number: FoF 04482

*Saprobic* on unidentified wood in a stream. Sexual morph Undetermined. Asexual morph Hyphomycetous. *Synnemata* 460–620 × 7.5–15 μm (\(\bar{x} = 587 \times 8.5 \mu m, n = 5\)), scattered but abundant, arising from the host surface. *Stipes* 175–223 × 28–42 μm, pale to dark brown, branched and flared above. *Conidiogenous cells* 16–24 × 0.8–2.1 μm (\(\bar{x} = 18.5 \times 1.7 \mu m; n = 20\)), annellidic, pale brown, verrucose. *Conidia* 5.8–7.3 × 2.6–3.7 μm (\(\bar{x} = 6.2 \times 3 \mu m, n = 30\)), aseptate, hyaline, cylindrical to obovoid, hyaline, smooth, produced in a mucilaginous mass on synnema.

Culture characteristics – Conidia germinating on PDA within 12 h. Colonies on MEA circular, entire edge with white to cream, raised on surface media reverse gray in the center, becoming white towards the margin.

Known distribution (based on molecular data) – Australia, China, Cuba, Czech Republic, France, Italy, Jamaica, Japan, Malaysia, Madagascar, New Zealand, Nicaragua, Poland, Thailand, The Netherlands, USA (Lackner & de Hoog 2011, Perera et al. 2018), UK (this study).

Known hosts (based on molecular data) – Human, foot mycetoma, on rotten stem of *Echium* sp., dried seed pod of *Delonix regia* (Lackner & de Hoog 2011, Perera et al. 2018), wood (this study).
Material examined – UK, Isle of Wight, Calbourne, on unidentified wood, stream, 7 September 2017, E.B.G. Jones, GJ429 (MFLU 19–1237, new geographical record), living culture MFLUCC 17–2483.

Fig. 148 – Cordyceps rosea (HKAS102492, lectotype). a–e Stroma raised from insect body. f Perithecia in stroma. g Vertical section through perithecia. h Perithecium. i Peridium. j Asci. k–m Disarticulating ascospores. n Part of asci. o Apical region of asci. p, q Upper and lower view of culture on PDA. a, f, g, h, j, k, m, p, q from HKAS102492 (neotype), b from HKAS102493, c from HKAS102494, d from HKAS102496, e, l, n, o from HKAS102495. Scale bar: f, g = 1000 μm, h = 500 μm, i = 30 μm, j = 50 μm, k, n, o = 20 μm, l, m = 5 μm, k = 10 μm, n–p = 3 μm. (m stained with cotton blue solution).
Maximum likelihood analysis with 1000 bootstrap replicates yielded a best tree with the likelihood value of -7124.535878. The combined ITS and LSU sequence datasets comprised 32 strains of Microascales with *Graphium fimbriasporum* (CMW5605), *G. penicilloioides* (CBS 102632) and *Trichoderma viride* (DAOM JBT1003) as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The matrix had 564 distinct alignment patterns, with 32.45% of undetermined characters.
or gaps. Estimated base frequencies were as follows; A = 0.239641, C = 0.266161, G = 0.273521, T = 0.220676; substitution rates AC = 2.021294, AG = 2.471263, AT = 1.983840, CG = 1.481755, CT = 5.565293, GT = 1.000000; gamma distribution shape parameter α = 0.245860. Maximum likelihood bootstrap (ML, black) values > 65% and Bayesian posterior probabilities (PP, green) > 0.80% are given above the nodes. The scale bar indicates 0.06 changes. The ex-type strains are in bold and new isolate is in blue bold.

GenBank numbers – LSU: MN017875, ITS: MN047111.
Additional GenBank number – SSU MN017934.
Notes – The morphological characters of our strain resemble *P. putredinis* (Lackner & de Hoog 2011). Our phylogenetic analyses with combined ITS and LSU sequence data indicate that our new strain, MFLUCC 17-2483 grouped together with the *Parascedosporium putredinis* strains (Fig. 149). Furthermore, based on a megablast search using the ITS sequence, the closest matches of *P. putredinis* (MFLUCC 17-2483) in NCBI's GenBank nucleotide database were *P. putredinis* (GenBank MF782705; similarities 558/559(99%), 1/559(0%)), *P. putredinis* (GenBank MH048678; similarities 544/549(99%), 0/549(0%)), *P. putredinis* (GenBank MF782706; similarities 555/559(99%), 1/559(0%)) and *P. putredinis* (GenBank KC894850; similarities 555/559(99%), 1/559(0%)).

Subclass Xylariomycetidae O.E. Erikss & Winka
*Amphisphaeriales* D. Hawksw. & O.E. Erikss.
*Apiosporaceae* K.D. Hyde et al.

Hyde et al. (1998) re-examined the type (*Apiospora montagnei* Sacc.) and introduced Apiosporaceae to accommodate *Apiospora* (Sexual morph of *Arthrinium*) and *Appendicospora*. Subsequently, this family has been recognized Xylariales by several authors (Huhndorf et al. 2004, Zhang et al. 2006, Jaklitsch & Voglmayr 2012, Sharma et al. 2014, Senanayake et al. 2015, Dai et al. 2016).

*Nigrospora* Zimm.

*Nigrospora* was introduced by Zimmerman (1902) for *N. panici* which was isolated as an endophyte from leaves of *Panicum amphibium* in Java, Indonesia. Mason (1927) transferred several black-spored hyphomycetes occurring on monocotyledonous hosts to *Nigrospora* and included several species in this genus. Based on its conidial characters, *Nigrospora* was placed in Dermateaceae (Moniliales) by Barnett & Hunter (1998) and Kirk et al. (2008b) assigned *Nigrospora* and its *Khuskia* sexual morph to the Trichosphaeriaceae (Trichosphaeriales). Wang et al. (2017) placed *Nigrospora* in Apiosporaceae based on phylogenetic analyses of combined ITS, tef1-α and TUB sequence data of 165 strains from China and Europe.

*Nigrospora oryzae* (Berk. & Broome) Petch, J. Indian bot. Soc. 4: 24 (1924) Fig. 152

Facesoffungi number: FoF06596

*Saprobi* on fallen dead leaves of *Phyllostachys heterocycla*. Sexual morph: Undetermined.
Asexual morph: Hyphomycetes. *Conidiophores* 24–32 × 2–3 μm (μ = 28 × 2.5 μm, n = 10) macronematous, mononematous, flexuous, smooth, hyaline to pale brown, branched. *Conidiogenous cells* 3.2–5.6 × 2–3 μm (μ = 4.9 × 2.6 μm, n = 20) holoblastic, monoblastic, solitary, hyaline to subhyaline, ampulliform to subsphaerical, discrete, smooth. *Conidia* 4–5.5 × 4.2–5 μm (μ = 5.2 × 4.8 μm, n = 100) dry, solitary, acrogenous, simple, spherical to broadly ellipsoidal, aseptate, black, smooth, velvety.

Culture characteristics: Conidia germinating on PDA within 12 hours and germ tubes produced from all cells. Colonies growing on PDA, hairy, black, reaching 5 mm in 7 days at 30 ºC, mycelium partly superficial, partly immersed, slightly effuse, radially striate, with irregular edge, black; Asexual spores were formed after 30 days and sexual spores not formed within 60 days.
Known distribution (based on molecular data) – Australia (Barkat et al. 2016), China (Sun et al. 2011, Chen et al. 2018, Zhang et al. 2019b), Iran (Hashemian Kalati et al. 2014), Italy (Lorenzini et al. 2016), Kazakstan (Eken et al. 2016), Pakistan (Alam et al. 2017b), Sri Lanka (Wang et al. 2017), Thailand (this study).

Fig. 150 – *Parascedosporium putredinis* (MFLU 19–1237, new geographical record). a–b Conidiophores on host substrate. c–e Erect synnematous conidiophores. f Conidia attached to the conidiogenous cells. g Conidia. h, i Colonies on seawater MEA. (h-upper, i-lower). Scale bars: c–e = 100 μm, f, g = 10 μm.
Known hosts (based on molecular data) – Acer truncatum (Sun et al. 2011), Aloe vera (Alam et al. 2017b), Castanopsis sp. (Wang et al. 2017), Cephalotaxus sinensis (Wang et al. 2017), Citrullus lanatus (Chen et al. 2018), Citrus reticulata (Wang et al. 2017), Clearya japonica (Wang et al. 2017), Daphniphyllum macropodum (Wang et al. 2017), Daphniphyllum oldhamii (Wang et al. 2017), Gossypium hirsutum (Zhang et al. 2019b), Hamamelis mollis (Wang et al. 2017), Nelumbo nucifera (Chen et al. 2018), Nelumbo sp. (Wang et al. 2017), Neolitsea sp. (Wang et al. 2017), Oryza sativa (Wang et al. 2017), Osmanthus fragrans (Wang et al. 2017), Osmanthus sp. (Wang et al. 2017), Pennisetum americanum (Hashemian Kalati et al. 2014), Pentactina rupicola (Wang et al. 2017), Photinia serrulata (He et al. 2019a), Rhododendron simiarum (Wang et al. 2017), Rhododendron sp. (Wang et al. 2017), Rubus reflexus (Wang et al. 2017), Rubus sp. (Wang et al. 2017), Symylcos zizyphoides (Wang et al. 2017), Ternstroemia sp. (Wang et al. 2017), Triticum aestivum (Eken et al. 2016, Barkat et al. 2016), Tutcheria microcarpa (Wang et al. 2017), Vaccinium corymbosum (Zhang et al. 2019b), Vitis vinifera (Lorenzini et al. 2016), Phyllostachys heterocycla (this study).

Material examined – Thailand, Chiang Rai Province, Doi Mae Salong, on fallen dead leaves of Phyllostachys heterocycla (Poaceae), 18 May 2016, J. F Li, DMS12-2 (MFLU 17–0199, new host and geographical record), living culture at KUMCC 16–0041.

GenBank numbers – ITS: MN648321, LSU: MN64832, SSU: MN64831.

Notes – Nigrospora oryzae has a cosmopolitan distribution and a wide host range (Wang et al. 2017, Raza et al. 2019). In this study, our collection (KUMCC 16-0041) collected from Doi Mae Salong, Thailand, clustered in a single clade with N. oryzae (Fig. 151) and their morphological characteristics were similar to those of N. oryzae. Therefore, we regard this isolate (KUMCC 16-0041) as N. oryzae as the first report.

Nigrospora sphaerica (Sacc.) E.W. Mason, Trans. Brit. Mycol. Soc. 12. 158. (1927) Fig. 153

Facesoffungi number: FoF06599

Hyphae smooth, hyaline, branched, septate, 3–8 μm diameter Sexual morph: Undetermined. Asexual morph: Conidiophores micronematous or semi-macronematous, septate, branched, flexuous or straight, hyaline to pale brown, smooth, 3–8 μm thick; hyaline vesicles usually surrounding the septum to delimit the conidia and their conidiogenous cells. Conidiogenous cells pale brown, holoblastic, monoblastic, determinate, subsphearical, 7–11 μm diameter \( \bar{x} = 6, n=10 \). Conidia formed abundantly, solitary, globose or subglobose, brown, black, shining, smooth, aspetate, 16–21 μm diameter \( \bar{x}=18, n=20 \).

Culture characteristics – On PDA, colonies floccose, margin circular. Colonies initially white, becoming black with age, reaching 8 cm diameter in 7 d at 25 °C.

Known distribution )based on molecular data( – China and Pakistan )Alam et al. 2017a, Wang et al. 2017, ( Russia )this study(.

Known hosts )based on molecular data( – Actinidia sp., Camellia sp., Citrus sp., Clearya japonica, Deutzia sp., Harpullia longipetala, Musa paradisiaca, Nelumbo sp., Rhododendron arboreum, Rosa sp., Saccharum sp., Sesamum indicum )Zhao et al. 2014, Wang et al. 2017, Alam et al. 2017a, Cui et al. 2018, Hemerocallis sp. )this study(.

Material examined – Russia, Rostov region, Rostov-on-Don, Zelezhodorozhny City District, street flowerbed, on dead floriferous shoot of Hemerocallis fulva (Asphodelaceae), 11 October 2017, T. Bulgakov T2279-2 )MFLU 18–0439, new host and geographical record(, culture, MFLUCC 18–0895.

GenBank Numbers – ITS: MN788673, tef1: MN995332.

Notes – We were unable to find any report of Nigrospora sphaerica from Hemerocallis sp. )Farr & Rossman 2020(, Therefore, herein we provide the first report of N. sphaerica from Hemerocallis from Russia.
Fig. 151 – Phylogram generated from the best scoring of the RAxML tree based on combined ITS, *tefl* and TUB sequenced data of taxa in *Nigrospora*. *Arthrinium vietnamense* (IMI 99670) was selected as the outgroup taxon. The best RAxML tree with a final likelihood value of -3487.815676 is presented. The matrix had 227 distinct alignment patterns, with 46.27% undetermined characters or gaps. Estimated base frequencies were as follows: $A = 0.242574$, $C = 0.243318$, $G = 0.252046$, $T = 0.262062$; substitution rates $AC = 1.131584$, $AG = 1.129186$, $AT = 0.645273$, $CG = 0.825588$, $CT = 4.542097$, $GT = 1.000000$; gamma distribution shape parameter $\alpha = 0.020014$. Bootstrap support for maximum likelihood (ML, black) equal to or greater than 60% and BYPP (red) equal to or greater than 0.95 are given above or below the nodes. Newly generated sequences are indicated in blue.
Fig. 152 – Nigrospora oryzae (MFLU 17-0199, new host and geographical record). a. Colonies on bamboo leaves. b, c. Conidia and conidiophores. d–j Conidiophores with conidiogenous cells. k–u Conidia. Scale bars: a = 200 µm, b = 50 µm, c = 20 µm, d–f = 10 µm, g–u = 5 µm.

Beltraniaceae Nann.

Beltraniaceae is based on Beltrania and similar genera, which are characterized by biconic, lageniform to navicular conidia, with or without a hyaline band, and with or without swollen separating cells (Seifert et al. 2011, Lin et al. 2017a). Conidiophores separate, or arise from, basal cells of setae, with or without radially lobed basal cells (Seifert et al. 2011, Lin et al. 2017a). In this study, we introduce two novel species, Beltrania dushanensis and Beltraniella brevis.

Beltrania Penz., Nuovo G. Bot. Ital. 14(2): 72 (1882)

Fourteen species are accepted in Beltrania (Lin et al. 2017a, Tibpromma et al. 2018b). Some
Beltrania species are plant pathogens, e.g., *B. pseudorhombica* was reported as the causal agent of pistachio leaf and fruit spot in Arizona, United States (Lichtemberg et al. 2019).

**Fig. 153** – *Nigrospora sphaerica* (MFLU 18–0439, new host and geographical record). a–b Upper surface and reverse overview of culture on PDA. c Conidiogenous cell giving rise to conidia. d Conidia. Scale bars: c, d = 10 μm.

**Beltrania dushanensis** C.G. Lin, Jian K. Liu & K.D. Hyde, sp. nov.

Index Fungorum Number: IF556728; Facesoffungi number: FoF06220

Etymology – Referring to Dushan, the type locality of this species

Holotype – MFLU 19–2252

*Saprobic* on decaying seed. Sexual morph: Undetermined. Asexual morph: *Colonies* on plant substrate effuse, dark brown, velutinous. *Mycelium* mostly immersed in the substratum. *Setae* numerous, erect, straight or slightly flexuous, unbranched, thick-walled, smooth, pale brown to dark brown, 180–260 μm long, 4–8 μm wide at the base, tapering to a pointed apex, arising from a dark brown, swollen, radially lobed basal cell, 10–17 μm diameter *Conidiophores* macronematous, single or in small groups, straight or flexuous, septate, smooth, thick-walled, cylindrical or clavate, pale brown, 10–45 μm long, 4–7 μm wide, arising from basal cells of setae or from separate dark brown, swollen, radially lobed cells, 8–18.5 μm diameter, often reduced to conidiogenous cells. *Conidiogenous cells* polyblastic, integrated, terminal, sympodial, denticulate, cylindrical, clavate, pale brown, smooth, 6.5–24 μm long, 4.3–8.5 μm wide. *Separating cells* ovoid, ellipsoidal, smooth, thin-walled, hyaline, 7.9–12.7 μm (ȳ = 10.5 μm, n = 23) long, 4.4–5.8 μm (ȳ = 5.1 μm, n = 23) wide in the broadest part. *Conidia* arise directly from conidiogenous cells or from separating cells, acrogenous, simple, smooth, dry, straight, biconic, rostrate, appendiculate, pale brown with a
subhyaline equatorial transverse band, 21.4–27.9 µm (\(\bar{x} = 24.3 \mu m, n = 30\)) long, 8.9–12.6 µm (\(\bar{x} = 10.4 \mu m, n = 30\)) wide in the broadest part; appendage 6.3–12.1 µm long, 0.7–1.6 µm wide at the base, tapering to a pointed apex.

Material examined – China, Guizhou Province, Qiannan Buyi Miao Autonomous Prefecture, Dushan County, Guizhou Zilinshan National Forest Park (Shengou District), on decaying seeds, 6 July 2018, Chuan-Gen Lin, DS 1-5 (MFLU 19–2252, holotype); *ibid.* (HKAS 105107, isotype); ex-type living culture GZCC18–0020.

**Fig. 154** – Phylogram generated from maximum likelihood analysis based on combined LSU and ITS sequence data. Forty-seven strains are included in the combined gene analyses comprising 1456 characters after alignment (856 characters for LSU, 600 characters for ITS). *Castanediella couratarii* (CBS 579.71) is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -4888.682999 is presented. The matrix had 384 distinct
alignment patterns, with 11.82% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.251976, C = 0.212516, G = 0.269311, T = 0.266196; substitution rates AC = 1.482978, AG = 2.788030, AT = 1.703126, CG = 0.797716, CT = 6.027412, GT = 1.000000; gamma distribution shape parameter α = 0.610669. Bootstrap values for maximum likelihood equal to or greater than 50 and Bayesian posterior probabilities equal or greater than 0.95 are placed near the branches. Ex-type strains are in bold and black. The newly generated sequences are indicated in blue.

GenBank numbers – ITS: MN252875, LSU: MN252882.

Notes – *Beltrania dushanensis* is most similar to *B. rhombica* Penz. in having biconic, symmetrical, light brown conidia with a V-shaped proximal end, but differs by its short conidiophores which are often reduced to conidiogenous cells (Pirozynski 1963, Morelet 2001). In addition, *B. dushanensis* and *B. rhombica* can be recognized as phylogenetically distinct lineages (Fig. 154). Phylogenetically, *B. dushanensis* shows a close relationship with *B. krabiensis* Tibpromma & K.D. Hyde (Fig. 154) since they clustered and formed a monophyletic group, however, they can be recognized as distinct lineages. *Beltrania dushanensis* differs from *B. krabiensis* by its shorter conidiophores and larger conidia (Tibpromma et al. 2018b).

**Beltraniella** Subram., Proc. Indian Acad. Sci., Sect. B 36: 227 (1952)

Twenty-six species are accepted in *Beltraniella* (Crous et al. 2016b, Lin et al. 2017b, Tibpromma et al. 2018, Crous et al. 2019a). However, only 11 species of *Beltraniella* have molecular data. A new species of *Beltraniella* from China is described based on phylogenetic analyses and morphological characters.

**Beltraniella brevis** C.G. Lin, Jian K. Liu & K.D. Hyde, sp. nov.

    Index Fungorum Number: IF556729; Facesoffungi number: FoF06219
    Etymology – Referring to the short conidiophores
    Holotype – MFLU 19–2254

*Saprobic* on decaying leaves. Sexual morph: Undetermined. Asexual morph: *Colonies* on plant substrate effuse, thin, pale brown. *Mycelium* mostly immersed in the substratum. *Setae* numerous, erect, arising from radially lobed basal cells, straight or flexuous, unbranched, single or in small groups, thick-walled, verrucose, dark brown, 89–251 μm long, 4.5–10.5 μm wide at the base, tapering to a pointed apex, arising from a dark brown, swollen, radially lobed basal cell, 15.5–27.8 μm diameter, imperfect setae single, straight, septate, verrucose, thick-walled, dark brown at the base and paler at the apex, up to 310 μm long, swollen at the base, slightly tapering to a pointed apex. *Conidiophores* macronematous, short, simple or branched, septate, sometimes reduced to conidiogenous cells, smooth-walled, swollen at the base, subhyaline to pale brown, thin-walled, arising from basal cells of setae or from separate, 6–16 μm long, 3.7–7 μm wide. *Conidiogenous cells* polyblastic, integrated, determinate, terminal, ampulliform, cylindrical, oblong, hyaline to subhyaline, smooth, 4.9–13.7 μm (x̄ = 8.3 μm, n = 29) long, 3.5–6.8 μm (x̄ = 5.3 μm, n = 29) wide at the base. *Separating cells* fusiform, thin-walled, smooth, hyaline to subhyaline, 1-denticulate at each end, 11–18 μm long, 3.4–4.1 μm wide in the broadest part. *Conidia* arise directly from conidiogenous cells or from separating cells, aggregated, acrogenous, simple, dry, straight, smooth, thin-walled, turbinate to pyriform, rostrate to pointed at proximal end, truncate at distal end, hyaline with a hyaline supraequatorial transverse band, 20–26.5 μm (x̄ = 22.3 μm, n = 30) long, 4.5–7.2 μm (x̄ = 5.9 μm, n = 30) wide in the broadest part.

Material examined – China, Guizhou Province, Qiannan Buyi Miao Autonomous Prefecture, Dushan County, Guizhou Zilinshan National Forest Park (Shengou District), unnamed road, on decaying leaves, 6 July 2018, Chuan-Gen Lin, DS 2–21 (MFLU 19–2254, holotype); ibid., HKAS 105159, isotype); ex-type living culture GZCC 18–0081; ibid, DS 2–23 (MFLU 19–2253, paratype, HKAS 105160, isoparatype); ex-paratype living culture GZCC 18–0082.

GenBank numbers – ITS: MN252876, MN252877, LSU: MN252883, MN252883.
**Fig. 155** – *Beltrania dushanensis* (MFLU 19–2252, holotype). a Host material. b, c Conidiophores on leaf surface. d Setae, conidiophores and conidiogenous cell. e, f Conidiogenous cells. g–i Separating cells. j–l Conidia. Scale bars: d = 50 μm, e, f, j–l = 10 μm, g–i = 5 μm.

Notes – *Beltraniella brevis* is most similar to *Bel. portoricensis* in having straight, verrucose setae and short non-setiform conidiophores arising from radially lobed basal cells, polyblastic conidiogenous cells, swollen separating cells, and turbinate, appendiculate conidia (Pirozynski 1963, Shirouzu et al. 2010). However, *Bel. brevis* differs from *Bel. portoricensis* by its hyaline conidia, whereas conidia are subhyaline to dilute yellow-olive in *Bel. portoricensis* (Pirozynski 1963, Shirouzu et al. 2010). In the tree generated from maximum likelihood analysis based on a
combined LSU and ITS sequence data (Fig. 154), Bel. brevis grouped together and formed a monophyletic group with 96% ML bootstrap support and 100% Bayesian posterior probabilities within Beltraniiella.

**Sporocadaceae** Corda

Sporocadaceae species have acervular or pycnidial conidiomata, and hyaline, or pale brown to dark brown, cylindrical to subcylindrical or fusiform, multi-septate conidia, usually with appendages at one or both ends (Nag Raj 1993, Liu et al. 2019). The sexual morph of this family is characterized by perithecial ascomata with simple, rarely branched paraphyses, unitunicate, 8-spored, cylindrical conidiomata with J+, apical ring, and pale, yellow to dark brown, fusiform or ellipsoidal and septate ascospores (Jaklitsch et al. 2016a, Liu et al. 2019). The taxonomy and phylogeny of Sporocadaceae species were by Jaklitsch et al. (2016a) and Liu et al. (2019). Approximately 30 well-supported monophyletic genera were included in Sporocadaceae (Liu et al. 2019).

**Bartalinia** Tassi

*Bartalinia* is characterized by pycnidial, unilocular to multi-loculate, glabrous conidiomata, and cylindrical to subcylindrical or fusiform, 3–4-euseptate conidia with a hyaline apical cell, bearing attenuated, flexuous, divergent, apical appendages with 2–4 branches and a tubular, single, unbranched, exogenous basal appendage (Nag Raj 1993, Jeewon et al. 2002). This genus has been assigned to Amphisphaeriaceae (Jeewon et al. 2003, Crous et al. 2014a) and Bartaliniaceae (Senanayake et al. 2015, Wijayawardene et al. 2016). Jaklitsch et al. (2016a) and Liu et al (2019) revived Sporocadaceae and reduced Bartaliniaceae to a synonym of this family.

**Bartaliniarobillardoides** Tassi, Bulletin Labor. Orto Bot. de R. Univ. Siena 3: 145 (1900)

*Facesoffungi* number: FoF06861

*Saprobic* on dead bark of *Picea excelsa*. Sexual morph: Undetermined. Asexual morph: *Conidiomata* 100–200 μm diameter, 100–150 μm high, brown, stromatic, pycnidial, scattered to gregarious, initially semi-immersed, ultimately erumpent, globose to subglobose, unilocular or multi-loculate, lacking an ostiole, dehiscing by an irregular break in the apical wall, glabrous. *Conidiomata wall* 30–60 μm wide, composed of thick-walled, brown cells of *textura angularis* in the outer layers, gradually merging with hyaline cells in the inner layers. *Conidiophores* arising all around the cavity of the conidiomata, cylindrical, hyaline, branched, septate, smooth-walled. *Conidiogenous cells* 1–4 μm wide, hyaline, holoblastic, ampulliform to cylindrical, integrated, determinate, smooth-walled. *Conidia* 8–15 × 1–2 μm (x = 10 × 1.5 μm; n = 50), cylindrical to subcylindrical, 4-septate, slightly constricted at septa, bearing appendages at both ends; basal cell pale brown, obconic with obtuse base, median cells 3, subcylindrical, thick-walled, apical cell hyaline, conical; apical appendage 5–10 μm long, hyaline, 3-branched, tubular, filiform, flexuous; basal appendage 1.5–3 μm long, single, filiform, unbranched, centric to eccentric.

Culture characteristics – Colonies fast growing, reaching 50 mm diameter after one week at 25 °C, circular, whitened in first three days, after two weeks becoming pale grey, flattened, felt-like, dense, aerial, surface smooth with crenate edge, filamentous, reverse whitened to pale brown in a week, finally brown, sporulation after one month.

Known distribution (based on molecular data) – Australia, Italy, Netherlands, South Africa, Thailand (Crous et al. 2014a, Wijayawardene et al. 2016, Liu et al. 2019), Italy (this study)

Known hosts (based on molecular data) – *Cupressus lusitanica* (Cupressaceae), *Eucalyptus* sp. (Myrtaceae), *Leptoglossus occidentalis* (Coreidae), *Poa* sp. (Poaceae) *Scadoxus puniceus* (Amaryllidaceae) (Crous et al. 2014a, Wijayawardene et al. 2016, Liu et al. 2019) *Picea excels* (this study).

Material examined – Italy, Province of Forlì-Cesena, Santa Sofia, Spescia, on dead cone of *Picea excels* (Pinaceae), 3 May 2014, E. Camporesi, IT1854 (MFLU 19–2560, new host record), living culture MFLUCC 15–0133, KUN, HKAS 101685.
GenBank Numbers – LSU: MN660234, ITS: MN688213, SSU: MN688209, tef1: MN683865, RPB2: MN683867.

Notes – Our collection fits well with the description of *B. robillardoides*. The sequence similarity between our collection and *B. robillardoides* (CPC 25385) is 100% (881/881) in LSU, 99% (549/552) in ITS, 100% (832/832) in RPB2 (Fig. 158). Our collection on *Picea excels* is regarded as a new host record.

**Fig. 156** – *Beltraniella brevis* (MFLU 19–2254, holotype). a Host material. b Conidiophores on leaf surface. c Setae with short conidiophores. d Imperfect setae. e, f Conidiogenous cells. g–j Conidia with separating cells. Scale bars: c, d = 20 μm, e–j = 10 μm.

**Discosia** Lib.

*Discosia* was introduced by Libert (1837) with *Discosia artocreas* selected as the lectotype by Vanev (1991). Senanayake et al. (2015) introduced Discosiaceae to accommodate *Adisciso*,...
Discosia, Discostroma, Immersidiscosia, Sarcostroma, and Seimatosporium but Jaklitsch et al. (2016a) treated Discosiaceae as a synonym of Sporocadaceae as the family has good phylogenetic support. The generic description of Discosia, coupled with the updated morphology as well as the phylogenetic relationship based ITS sequences were reviewed by Liu et al. (2019).

Fig. 157 – Beltraniella brevis (MFLU 19–2253, paratype). a Host material. b, c Conidiophores on leaf surface. d Setae with short conidiophores. e Imperfect setae with short conidiophores. f, g Conidiogenous cells. h–k Conidia with separating cells. Scale bars: d, e = 20 μm, f–k = 10 μm.
**Fig. 158**– Phylogram of *Bartalinia* and allied genera generated from maximum likelihood analysis based on combined LSU, ITS, tef1, RPB2 sequence data. Related sequences are taken from Liu et al. (2019). Fifteen strains are included in the combined analyses which comprise 2745 characters (LSU: 1–830, ITS: 831–1349, tef1: 1350–1913, RPB2: 1914–2745) after alignment. *Truncatella angustata* CBS 144025 is used as the outgroup taxon. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -8567.262112 is presented. The matrix had 512 distinct alignment patterns, with 4.60% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.255888, C = 0.231712, G = 0.245946, T = 0.266454; substitution rates AC = 1.331043, AG = 3.021366, AT = 1.507439, CG = 0.983080, CT = 7.311485, GT = 1.000000; gamma distribution shape parameter α = 0.131325. Bootstrap values for maximum likelihood (ML) equal to or greater than 50% and clade credibility values greater than 0.95 from Bayesian-inference analysis labeled on the nodes. The new isolate is indicated in bold and blue.

**Discosia pini** Heald, Mycologia 1(5): 216 (1909)

Facesoffungi number: FoF07469

Saprobic on dead fern leaves. Sexual morph: Undetermined. Asexual morph: Coelomycetous. *Conidiomata* 140–258 × 45–59 (x̄ = 195 × 54, n = 5) μm, sparse, acervular, superficial, bilocular, which is separated by interlocular wall and there are 2–3 hyaline, vertical supporting partitions inside the bigger loculus. *Peridium* 14–18 μm thick, brown, comprising thick-walled cells of *textura angularis*. *Conidiogenous cells* 4–6 × 1–2 (x̄ = 4.5 × 1.5, n= 15) μm, subcylindrical,
developing from the inner stromatic tissue, monophialidic, integrated, hyaline, smooth. Conidia 14–20 × 1.9–3 (\(\bar{x} = 17 \times 2.7, n = 50\)) μm, hyaline, fusiform with rounded ends, straight or slightly curved, 3-celled and the second cell from base is longer, with 2-appendages. Appendages 4–10 (\(\bar{x} = 8, n= 50\)) μm long, hyaline, straight, filiform, with rounded ends, with a round base, producing from the attachment between the middle cell and the terminal cell of conidia.

Fig. 159 – Bartalinia robillardoides (MFLU 19-2560, new host record). a, b Appearance of brown conidiomata on the host. c, d Vertical sections of conidiomata. e Section of peridium. f Germinating conidium. g, h Conidiophores, conidiogenous cells and developing conidia. i–l Conidia. m Culture on PDA. Scale bars: a = 1000 μm, b–d = 100 μm, e = 50 μm, f–l = 5 μm, m = 25 mm.
Fig. 160 – Phylogram generated from maximum likelihood analysis based on combined LSU, ITS and RPB2 sequence data. Thirty-one strains are included in the combined gene analyses comprising 2067 characters after alignment (475 characters for ITS, 784 characters for LSU, 808 characters for RPB2). *Sporocadus biseptatus* (CBS 110324) and *Sporocadus cornicola* (CBS 143889) are used as the outgroup taxa. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best RaxML tree with a final likelihood value of -6084.123259 is presented. The matrix had 320 distinct alignment patterns, with 23.85% undetermined characters or gaps. Estimated base frequencies were as follows: \(A = 0.251281\), \(C = 0.217598\), \(G = 0.258884\), \(T = 0.272237\); substitution rates \(AC = 1.723842\), \(AG = 6.440596\), \(AT = 1.103776\), \(CG = 1.039404\), \(CT = 12.923260\), \(GT = 1.000000\); gamma distribution shape parameter \(\alpha = 0.840718\). Bootstrap values for maximum likelihood equal to or greater than 70 and Bayesian posterior probabilities equal or greater than 0.90 are placed above or below the branches. The newly generated sequence is indicated in blue bold and the type strains are indicated in black bold.
Fig. 161 – *Discosia pini* (HKAS 101478, new host record). a, b Conidiomata on fern leaf. c Column. d Vertical section through conidioma showing two loculi and three columns. e, f Conidiogenous cell bearing conidia. g–i Conidia. j Germinating conidium. k, l Conidiogeneous cells formed from a mycelial strand on culture. m, n Upper and reverse review of culture after 8 days incubation on PDA. Scale bars: d = 50 µm, e, j, k = 20 µm, c, f–i, l = 10 µm. (The columns are indicated with black arrows, a–i observed from herbarium specimen, j–n described from culture collection).
Culture characters – Cultures were made from germinating conidia which germinated after 1d on PDA. The colonies are rapid-growing, attaining a diameter of 5 cm in 12 days at 22 °C. Colonies white with one grey annulation at the centre, becoming fully grey at the aging stage, velvety, circular, olive-green to grey from reverse. In vitro, the conidia form from the mycelial strand.

Material examined – China, Yunnan Province, Honghe State, Yunti County, on a decay fern leaf, 20 September 2017, De-Ping Wei, YT07 (HKAS 101478); living culture KUMCC 18-0033.

GenBank Numbers – ITS: MT012348, LSU: MT012355, RPB2: MT025047

Notes – The multiple gene tree indicates that our collection (HKAS 101478) has a close affinity to Discosia pini (MAFF 410149) and D. artocreas (NBRC 8975) with strong statistical support (87% ML/0.99 PP, Fig. 160). The nucleotide comparison shows that our isolate is identical to the above two reference species within 545 bp ITS and 782 bp LSU sequences. Unfortunately, except for BT gene sequence data, other gene sequences of D. pini (MAFF 410149) and D. artocreas (NBRC 8975) are not available and we could not successfully obtain BT gene sequences for our isolate in this study. Even though the full morphology of D. pini (MAFF 410149) and D. artocreas (NBRC 8975) are not linked to any literature (Tanaka et al. 2011), our isolate is phylogenetically separated from the ex-type strain of D. artocreas (CBS 124848). Our collection is similar to D. artocreas (CBS 124848) in having hyaline conidia with a longer second cell from base, but differs in the position where the appendages are produced. The apical appendage of D. artocreas is polar and the basal appendage is located far above the conidium base (Matsushima 1975, Liu et al. 2019). However, the appendage of our collection is borne from the junction of the terminal cell and the middle cell. The type specimen of D. pini (No. 758) was originally reported on Pini ponderosae by Heald (1909), and he provided a description and hand-drawing. Our collection resembles D. pini in having black, glabrous, bilocular conidiomata with vertical columns inside the larger loculus and similar size of conidia (14–20 × 1.9–3 μm in our collection vs 12–20 × 2.6 μm in NEB 758). Our collection differs slightly from D. pini (NEB 758) however, by having round ends to the conidia and shorter appendages (4–10 μm long), while D. pini (NEB 758) has relatively pointed ends to the conidia and longer appendages (10–12 μm long). Considering our collection is most similar to the type specimen of D. pini (NEB 758) rather than D. artocreas, we report our collection as a new host record of D. pini from fern leaf bases based on morphological and phylogenetic evidence. However, we suggest treating the strains D. artocreas (NBRC 8975) as an undetermined species as its morphology is not known and phylogenetically it separates from its type strain. Fresh collections with further sequence data may prove this conclusion to be incorrect.

Pestalotiopsis Steyaert

Pestalotiopsis was introduced by Steyaert (1949), with the type species P. guepinii (Desm.) Steyaert, a plant pathogenic taxon, on Rhododendron. Based on molecular evidence, the genus has 66 accepted species (Norphanphoun et al. 2019, Jayawardena et al. 2019b) and 257 species named in Index Fungorum (2020).

Pestalotiopsis diploclisiae Maharachch., K.D. Hyde & Crous, Stud. Mycol. 79: 160 (2014) Fig. 163

Facesoffungi number: FoF06982

Isolated from asymptomatic leaf of Kandelia candel. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial, globose, brown, semi-immersed on PDA, releasing conidia in a black, slimy, globose, glistening mass. Conidiophores indistinct. Conidiogenous cells annellidic, ampulliform to lageniform, hyaline, smooth- and thin-walled, simple, proliferating 1–2 times percurrently 5–10 × 5–8 μm. Conidia (18–)18.5–25(–26) × (5–)5.5–6(–6.5) μm (μ ± SD = 21 ± 2.2 × 5.5 ± 0.6 μm), fusiform to clavate, straight to slightly curved, 4-septate; basal cell obconic with a truncate base, hyaline or sometimes pale brown, thin- and smooth-walled, (3–)3.5–4(–5) μm long (μ ± SD = 3.8 ± 0.7 μm); three median cells (10–)10.5–15(–16) μm long (μ ± SD = 12.7 ± 1.7 μm), brown, septa and periclinal walls darker than rest of the cell, versicolored, wall rugose; second cell
from base pale brown, (3–)4–5–(6) μm long (± SD = 4.4 ± 0.8 μm); third cell brown, (3.5–)4–5–(6) μm long (± SD = 4.2 ± 0.7 μm); fourth cell brown, (3–)4–5–(5.5) μm long (± SD = 4.1 ± 0.5 μm); apical cell (4–)4.5–5–(6) μm long (± SD = 4.5 ± 0.7 μm), hyaline, conic to acute; with 1–3 tubular appendages on apical cell, inserted at different loci in a crest at the apex of the apical cell, branched, flexuous, (8.5–)9.5–33–(34) μm long (± SD = 20 ± 7.6 μm); single basal appendage, tubular, unbranched, centric, (3.5–)4–9–(10) μm long (± SD = 6.5 ± 1.6 μm).

Culture characteristics – Colonies on PDA reaching 6–7 cm diameter after 14 d at room temperature (±25 °C), under light 12 hr/dark 12 hr, colonies filamentous to circular, medium dense, aerial mycelium on surface flat or raised, fluffy, white from above and reverse.

Known distribution (based on molecular data) – China, Hong Kong (Maharachchikumbura et al. 2014), Taiwan (this study).

Known hosts (based on molecular data) – Diploclisia glaucescens (Menispermaceae), Psychotria tutcheri (Rubiaceae) (Maharachchikumbura et al. 2014), Kandelia candel (this study).

Material examined – Taiwan, Hsinchu City, tissue isolation from asymptomatic leaf of Kandelia candel (Rhizophoraceae), 16 July 2018, Norphanphoun Chada HsE1L-1A, living cultures, NCYU19-0356 (new host record). GenBank Numbers – TUB: MN885525, tef1: MN885527, ITS: MN887600, LSU: MN887606.

Notes – Pestalotiopsis diploclisiae was introduced by Maharachchikumbura et al. (2014) from the fruits of Diploclisia glaucescens and Psychotria tutcheri in Hong Kong. Our strain is similar to P. diploclisiae in spore size (in this study; 23 ± 1.5 × 5.5 ± 0.5 μm vs. 24 ± 1.3 × 5.7 ± 0.4 μm; Maharachchikumbura et al. 2014). Based on phylogenetic analysis our strain grouped with P. diploclisiae with low bootstrap support (66% ML, Fig. 162). Hence, we introduce our strain as P. diploclisiae, the first record from Kandelia candel in Taiwan.

Pestalotiopsis kenyana Maharachch., K.D. Hyde & Crous, Stud. Mycol. 79: 166 (2014) Fig. 164

Facesoffungi number: FoF06981

Isolated from asymptomatic leaf of Kandelia candel. Sexual morph: Undetermined. Asexual morph: Conidiomata pycnidial, globose, brown, semi-immersed on PDA, releasing conidia in a black, slimy, globose, glistening mass. Conidiophores indistinct. Conidiogenous cells holoblastic, ampulliform to lageniform, hyaline, smooth- and thin-walled, simple, 5–10 × 5–8 μm. Conidia (18–)18.5–25(–26) × (5–)5.5–6(–6.5) μm (± SD = 21 ± 2.2 × 5.5 ± 0.6 μm), fusiform to clavate, straight to slightly curved, 4-septate; basal cell obconic with a truncate base, hyaline or sometimes pale brown, thin- and smooth-walled, (3–)3.5–4(–5) μm long (± SD = 3.8 ± 0.7 μm); three median cells (10–)10.5–15(–16) μm long (± SD = 12.7 ± 1.7 μm), brown, septa and periclinal walls darker than rest of the cell, versicolored, wall rugose; second cell from base pale brown, (3–)4–5–(6) μm long (± SD = 4.4 ± 0.8 μm); third cell brown, (3.5–)4–5–(6) μm long (± SD = 4.2 ± 0.7 μm); fourth cell brown, (3–)4–5–(5.5) μm long (± SD = 4.1 ± 0.5 μm); apical cell (4–)4.5–5–(6) μm long (± SD = 4.5 ± 0.7 μm), hyaline, conic to acute; with 1–3 tubular appendages on apical cell, inserted at different loci in a crest at the apex of the apical cell, branched, flexuous, (8.5–)9.5–33–(34) μm long (± SD = 20 ± 7.6 μm); single basal appendage, tubular, unbranched, centric, (3.5–)4–9–(10) μm long (± SD = 6.5 ± 1.6 μm).

Culture characteristics – Colonies on PDA reaching 6–7 cm diameter after 14 d at room temperature (±25 °C), under light 12 hr/dark 12 hr, colonies filamentous to circular, medium dense, aerial mycelium on surface flat or raised, fluffy, white from above and reverse.

Known distribution (based on molecular data) – China (Yunnan Province, Jiangxi Province, Chongyi County) (Liu et al. 2017a), Kenya (Maharachchikumbura et al. 2014) Kandelia candel (this study).

Known hosts (based on molecular data) – Coffea sp. (Rubiaceae) strain number CBS 442.67, Camellia sinensis (Theaceae) strain numbers LC3291, LC3633 and LC6633 (Liu et al. 2017a, Maharachchikumbura et al. 2014, Taiwan (this study)).
Material examined – Taiwan, Hsinchu City, tissue isolation from asymptomatic leaf of *Kandelia candel* (Rhizophoraceae), 15 July 2018, Norphanphoun Chada HsE1P, living culture NCYUCC 19–0389 (new host record).

GenBank Numbers – TUB: MN885526, ITS: MN887601, LSU: MN887607.

Notes – *Pestalotiopsis kenyana* was introduced by Maharachchikumbura et al. (2014) from Kenya, with the character of two basal appendages. Our strain is similar to *P. kenyana* in spore size (22 ± 2.2 × 2.5 ± 0.6 μm vs. 25.5 ± 1.2 × 8 ± 0.4 μm; Maharachchikumbura et al. (2014)), and conidiogenous cell size (7–22 × 2–5 μm vs. 10–25 × 2–5 μm; Maharachchikumbura et al. (2014)). Based on phylogenetic analysis our strain is related to *P. kenyana* (Fig. 162). The collection of *P. kenyana* is the first record on Kandelia candel from Taiwan.

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

Maharachchikumbura et al. (2014) established *Pseudopestalotiopsis*. It was differentiated from its two sister genera *Pestalotiopsis* and *Neopestalotiopsis* based on conidial pigmentation and molecular phylogeny. As in *Pestalotiopsis*, the median cells in the conidia of *Pseudopestalotiopsis* are concolourous, but are much darker (brown to dark brown) and bear knobbed apical appendages (Liu et al 2010, Maharachchikumbura et al. 2014). In *Neopestalotiopsis* the median cells are versicolorous. *Pseudopestalotiopsis* species have been reported from terrestrial and mangrove habitats, as saprobes or as pathogens associated with leaf spots (Maharachchikumbura et al. 2014, 2016b, Norphanphoun et al. 2019).

**Pseudopestalotiopsis kubahensis** Lateef, M. Sepiah & Bolhassan, Current Research in Environmental & Applied Mycology 5(4): 378 (2014)  
Facesoffungi number: FoF01310

Endophytic or pathogenic causing irregular shaped lesions on branches of *Lonicera* sp. appearing as black, flattened patches. Sexual morph: Undetermined. Asexual morph: *Conidiomata* semi-immersed to erumpent, acervulus, exuding brown to black conidial masses, subglobose to globose with a flattened base, 250–300 μm in diameter, solitary, dark brown to black. *Conidioma wall* composed of light brown, thin-walled, flattened cells of *textura prismatica*, indistinct at the base, mostly fused with host tissue. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* holoblastic, arising from the innermost layers of the conidiomata, discrete, hyaline, smooth-walled, cylindrical to lageniform. *Conidia* broad fusiform, straight to slightly curved, 4-septate, slightly constricted at septa, 22.5–26.5 × 6.5–8.5 μm (x̄ = 25 × 7.5 μm, n = 40) basal cell conic with a truncate base, thin and smooth-walled, 2.5–3.5 μm long; three median cells 16–19.5 μm long, doliiform, concolourous, golden brown to dark brown, septa darker than the rest of the cell (second cell from base 4–6 μm long; third cell 3.5–4.5 μm long; fourth cell 5–6.5 μm long); apical cell conic, hyaline, thin and smooth-walled, 2.5–4 μm long; with 3 or occasionally 2 tubular apical appendages arising from the apical tip, flexuous, rarely branched, 15–35(40) μm long, occasionally knobbed; basal appendage not observed.

Culture characteristics – Colonies on PDA reaching approximately 7 cm diameter after 7 days at 16–18 °C, greyish-white, irregular, with undulate margins and black, gregarious conidiomata sporulating; reverse pale yellow.

Known distribution (based on molecular data) – Malaysia (Lateef et al. 2015), Italy (this study).

Known hosts (based on molecular data) – *Macaranga* sp. (Lateef et al. 2015), *Lonicera* sp. (this study).

Material examined – Italy, Province of Forlì-Cesena [FC], near Converselle - Castrocucco Terme e Terra del Sole, on living branch of *Lonicera* sp. (L.) (Caprifoliaceae), 13 December 2014,
Fig. 162 – Phylogram generated from maximum likelihood analysis based on combined ITS, TUB and refl sequence data for Pestalotiopsis. 82 strains are included in the combined gene analyses and Neopestalotiopsis protearum (CBS 114178) and N. acrostichi (MFLUCC 17-1754) are used as the out-group taxa. The best RAxML tree with a final likelihood value of -10930.743597 is presented. The matrix had 690 distinct alignment patterns, with 12.40% undetermined characters or gaps. Estimated base frequencies of Partition 0 (ITS) were as follows: A = 0.235848, C = 0.243464, G = 0.220143, T = 0.300545; substitution rates AC = 0.277769, AG = 2.981361, AT = 0.839816, CG =
Estimated base frequencies of Partition 1 (TUB) were as follows: A = 0.218592, C = 0.325639, G = 0.225308, T = 0.230461; substitution rates AC = 1.214354, AG = 5.267474, AT = 1.343109, CG = 1.322200, CT = 5.809285, GT = 1.000000; gamma distribution shape parameter α = 0.179684. Estimated base frequencies of Partition 2 (tefl) were as follows: A = 0.261334, C = 0.316753, G = 0.194433, T = 0.227480; substitution rates AC = 0.938048, AG = 2.227476, AT = 1.023715, CG = 0.645264, CT = 3.467662, GT = 1.000000; gamma distribution shape parameter α = 0.291483. Bootstrap values for maximum likelihood equal to or greater than 60 near the branches. The newly generated sequences are indicated in blue bold.

Fig. 163 – Pestalotiopsis diploclisiae (HsEIL-1A, new host record). a Habitat. b, c Kandelia candel. d Culture on PDA (leaf-above, right-reverse). e–g Colony sporulating on PDA. h Conidiogenous cells giving rise to conidia. i–m Conidia. Scale bars: e–f = 500 μm, g = 250 μm, h = 10 μm, i, m = 20 μm (j–l use same scale bar as m).
Fig. 164 – *Pestalotiopsis kenyana* (HsE1P, new host record). a Habitat. b, c *Kandelia candel*. d Culture on PDA (leaf-above, right-reverse). e–g Colony sporulating on PDA. h Conidiogenous cells giving rise to conidia. i–m Conidia. Scale bars: e = 2 mm, f, g = 1 mm, h = 250 µm, h–n = 10 µm (j–l use same scale bar as m).

E. Camporesi IT 2294 (MFLU 15–0844, new geographical record and host record), living culture MFLUCC 15–0565.

GenBank number – ITS: MG818971

Notes – In the phylogenetic analysis, the *Pseudpestalotiopsis* isolate (MFLUCC 15-0565) collected in the present study grouped with the type strain of *Ps. kubahensis* (UMAS KUB-P20), a species introduced from *Macaranga* sp. (Fig. 165), in Malaysia. Both produce spores with golden brown to dark, concolourous, median cells (Lateef et al. 2015). The conidial dimensions of the species from Malaysia is 27–30 × 5.6–7.3 µm and 22.5–26.5 × 6.5–8.5 µm for our collection. The lengths of apical appendages for UMAS KUB-P20 and MFLUCC 15-0565 are 16–29.5 µm and 15–35 µm respectively. However, morphologically our collection differs from *Ps. kubahensis* by lacking a
basal appendage. The relationship between the two strains is poorly resolved, as for both isolates only ITS data is available for sequence comparison. In a similar result obtained by Maharachchikumbura et al. (2016b) it was emphasized that the variation within the ITS region of *Pseudopestalotiopsis* species is limited, however, a combination of ITS, TUB and *tef1* gene data gave the best resolution as compared to any single marker for resolving *Pestalotiopsis* and related taxa. In the present study, we prefer to maintain our collection (MFLUCC 15-0565) as *Ps. kubahensis* until we have obtained more sequence data. This is the first *Pseudopestalotiopsis* sp. to be recorded on *Lonicera* sp. and is the first record from a temperate region, as all previous species introduced were from tropical regions.

**Fig. 165** – Phylogram generated from maximum parsimony (MP) analysis based on combined ITS, TUB, and *tef1* sequence data. Twenty-three strains are included in the combined gene analyses comprising 2323 characters after alignment (572 characters for ITS, 793 characters for TUB, 948 characters for *tef1*). *Neopestalotiopsis protearum* (CPC 1765) is used as the outgroup taxon. Bootstrap 50% majority-rule consensus tree is presented here. The combined dataset contained 1953 constant, 270 parsimony uninformative and 100 parsimony informative characters. Gaps were
treated as missing data. The alignment was subjected to 1000 bootstrap replicates. Statistics generated from MP analysis are as follows: TL = 492, CI = 0.852, RI = 0.741, RC = 0.631, HI = 0.148. The tree topology of the maximum likelihood (ML) analysis did not differ significantly from the maximum parsimony analysis. The best RaxML tree selected had a final likelihood value of -5981.645344. Bootstrap values for MP and ML analyses equal to or greater than 50 are placed above or below the branches. Ex-type strains are in bold and black. The newly generated sequence is indicated in blue.

Xylariales Nannf.
Lopadostomataceae Daranag. & K.D. Hyde
Lopadostomataceae was introduced by Senanayake et al. (2015) in Xylariales, to accommodate Lopadostoma and Creosphaeria. Jumillera and Whalleya were also included in the family by Daranagama et al. (2018).

Fig. 166 – Pseudopestalotiopsis kubahensis (MFLU 15–0844, new host record and geographical record). a Appearance of conidiomata on host. b Close-up of a conidioma. c Cross section of a conidioma. d, e Conidia arising from conidiogenous cells. f–i Conidia. Scale bars: a = 1000 μm, b = 200 μm, c = 100 μm, d, e = 10 μm, f–i = 20 μm.

Whalleya J.D. Rogers, Y.M. Ju & F. San Martín
Whalleya was introduced to accommodate two species previously known in Biscogniauxia (Rogers et al. 1997). Species of Whalleya are xylariaceous in sexual morph characters (Rogers et al. 1997) and conidial characters are similar to lopadostomataceous species. Previous studies have
placed Whalleya in Diatrypaceae (Glawe & Rogers 1986), because of similar characters to Xylariaceae and Lopadostomaceae. Wendt et al. (2017) and Daranagama et al. (2018) have treated Whalleya as a member of family Lopadostomataceae using both phylogeny and morphology.

**Whalleya microplaca** (Berk. & M.A. Curtis) J.D. Rogers, Y.M. Ju & F. San Martín, Mycotaxon 64: 48 (1997)

Faces of fungi number: FoF03015

*Saprobic* on woody litter of *Phoebe* spp. Sexual morph: *Stromata* applanate, solitary or confluent, bipartite, outer dehiscing layer dark brown, thin, exposing mature black surface, carbonaceous. *Ascomata* globose, 100–120 μm high, 213–245 μm diameter, tissue between ascomata mainly composed of fungal tissue, coriaceous, becoming brownish, tissue below the ascomatal layer inconspicuous. *Ostioles* umbilicate. *Asci* 8-spored, 33–38 × 2–3 μm, unitunicate, cylindrical, short-pedicellate, with discoid ring. *Ascospores* 8–9 × 2–3 μm, uniseriate, light brown, ellipsoid-inequilatertal, with narrowly rounded ends, 1-celled, with central prominent guttule. Asexual morph: Undetermined.

Culture characteristics – colonies on PDA white, covering entire plate in 14 days at 28 °C fluffy mycelium, and reverse white.

Known distribution – China, Yunnan, Mauritius, Louisiana, Florida, New Jersey, Philippines, and Taiwan (Farr & Rossman 2020), China (this study).

Known hosts (based on molecular data) – *Phoebe* spp., *Magnolia glauca*, *Melia* sp., *Persea* sp., *Sassafras albidum*, *Sassafras officinale* (Farr & Rossman 2020), *Phoebe* spp. (this study).

Material examined – China, Yunnan, on woody litter of *Phoebe* sp. (Lauraceae), 5 December 2018, D.N Wanasinghe, DW0122 (JZBH 3400001) living culture, JZB 3400001.

GenBank number – ITS: MN989425.

Notes – *Whalleya* is typified by *W. microplaca* and has sexual morph characters similar to *Lopadostoma pouzarii* (Daranagama et al. 2018). A collection obtained from the woody litter of *Phoebe* spp. was identified as *Whalleya microplaca* with support from morphology and phylogeny. Our isolate clustered with the reference strain of *Whalleya microplaca*, in the combined ITS, LSU and RPB2 sequence analysis with 100% support (Fig. 167). This is the first record of *Whalleya microplaca* species reported from *Phoebe* sp. in China.

**Xylariaceae** Tul. & Tul.

Xylariaceae was introduced by Tulasne & Tulasne (1863) using the term “Xylariei” (Stadler et al. 2013). According to Maharachckumbura et al. (2016a), Xylariaceae is one of the largest and most diverse families of Ascomycota. The exact number of accepted taxa varies within the family (Hyde et al. 2020). Presently, Xylariaceae comprises 44 genera and more than 1350 species (U’Ren et al. 2016, Wijayawardene et al. 2018). Xylariaceae is characterized by perithecial ascomata, embedded in a more or less well-developed, dark stromata or sometimes reduced or lack stromata, 8-spored, unitunicate, cylindrical asci with or without J+, apical rings and pigmented ascospores with germ slits or pores. The asexual morphs produce conidia holoblastically from sympodially, or occasionally percurrently proliferating conidiogenous cells (Rogers 2000, Daranagama et al. 2018).

**Rosellinia** De Not.

*Rosellinia* was introduced by De Notaris (1844) to accommodate *Rosellinia aquila* (Fr.) Ces. & De Not. as the type species. *Rosellinia* is a species rich genus (Petrini 1992, 2003, 2013). There are 142 accepted *Rosellinia* species including many type specimens based on morphological studies (Petrini 2013). *Rosellinia* is characterized by superficial, subglobose, semiglobose, mammate to cupulate or conical, ostiolate, unipерitheciate, brown to black stromata seated in a subiculum; cylindrical, pedicellate asci with J+, ascal apical rings with a rounded, angular or
Fig. 167 – Phylogram generated from maximum likelihood analysis based on a combined ITS, LSU and RPB2 sequence dataset. Thirty-two strains are included in the combined gene analyses comprising 2319 total characters including gaps. The tree topology derived from the Bayesian analysis was similar to that derived from the maximum likelihood analysis. The best scoring of the ML tree is selected to represent the phylogenetic relationships of taxa in Lopadostomataceae, with the final ML optimization likelihood: -3536.406809. The matrix had 293 distinct alignment patterns, with 18.47% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.211039, C = 0.292321, G = 0.252470, T = 0.244170; substitution rates AC = 1.328514, AG = 2.488703, AT = 1.264091, CG = 0.687931, CT = 4.488656, GT = 1.000000; Tree-Length = 1.050171; gamma distribution shape parameter $\alpha = 0.517888$. The newly generated sequences are indicated in blue.
indistinct bulge at the upper rim, the rings are usually higher than the width. The ascospores are slightly larger, 1-celled, with a germ slit in most species (Petrini 2013). The asexual morph produces conidia in the subiculum, on immature perithecia or in culture (Petrini & Petrini 2012).

Fig. 168 – *Whalleya microplaca* (JZB 3400001, new host record). a, b Stromata in wood. c, d Cross section of stroma showing ascomata encased in stromal tissue. e–g Asci. h Ascus tip. i Ascospores. Scale bars: a, b = 200 µm, e, f, g = 10 µm h–i = 10 µm.
Fig. 169 – Phylogram generated from maximum likelihood analysis based on combined ITS, RPB2, ACT, and TUB sequence data. Forty strains are included in the combined gene analyses comprising 2513 characters after alignment. *Amphiloga gyrosa* (YMJ 91123101) is used as the outgroup taxon. The tree topology of the Bayesian analysis was similar to the maximum likelihood analysis. The best RaxML tree with a final optimization likelihood value is -34670.729423. The matrix had 323 distinct alignment patterns, with 12.37% undetermined characters or gaps. Estimated base frequencies were as follows: A = 0.240651, C = 0.275519, G = 0.236248, T = 0.247582; substitution rates AC = 1.504126, AG = 4.316567, AT = 1.308364, CG = 1.338210, CT = 4.785013, GT = 1.000000; gamma distribution shape parameter α = 0.705373. Bootstrap values for maximum likelihood equal to or greater than 65 and Bayesian posterior probabilities equal or greater than 0.95 are placed above or below the branches. The newly generated sequences are indicated in bold and blue.

*Rosellinia* species have a wide distribution as saprobes, endophytes or pathogens of various host species (Su et al. 2016a). Many *Rosellinia* species cause plant diseases (Whalley 1996). In our study, we introduce *Rosellinia convexa* as a new host record from *Prunus subhirtella* (Rosaceae) in China.
**Rosellinia convexa** Q.R. Li & J.C. Kang, Mycoscience 57(3): 166 (2016)  
Facesoffungi number: FoF 00890

*Saprobic* on dead host stem of *Prunus subhirtella*. Sexual morph: *Subiculum* feltly, brownish grey, persistent, appressed. *Ascomata* 940–1330 μm diameter × 760–835 μm high (x̄=1165 × 798 μm, n=10), perithecial, scattered to gregarious, superficial with the base slightly sunken, globose to subglobose, greyish brown to black, apex with a finely conical, thick eostroma, slightly carbonaceous, black, finely roughened, thickened at the base and spreading around the stroma base beneath the subiculum, entostroma restricted to a thin fibrous gray color tissue encasing the perithecium, perithecia not collapsed in the stroma cavity, ostiolar papilla. *Paraphyses* 1–2 μm wide, sparse, thin walled, tapering above asci. *Asci* 125–190 × 7–13 μm (x̄ =155 × 11 μm, n =20) cylindrical, 8-spored, pedicel 19–40 μm (x̄ = 32 μm, n = 20) long, with apical ring, 6–7 × 6–9 μm (x̄ =8 × 8 μm, n =20), cuboid-cuneate with an upper lateral rim, J+ in Melzer’s reagent. *Ascospores* 16–19 μm × 7–10 μm (x̄ =18 × 8 μm, n =30), obliquely overlapping uniseriate, dark brown, narrowly to broadly ellipsoidal-equilateral with broadly rounded, with a conspicuous straight germ slit almost along entire spore-length on the ventral side, entirely surrounded by an ill-defined slimy sheath, 1.8–4.5 μm wide, visible in India ink, without appendages. Asexual morph: Undetermined.

Culture characteristics – *Ascospores* germinating on PDA within 24 h, from single spore isolation. Colonies on PDA reaching 25–30 mm diameter after one week at 16 °C, circular, entire edge, white, with dense, convex papillate surface, brown in reverse.

Known distribution (based on molecular data) – Guizhou Province, China (Su et al, 2016a and this study).

Known hosts (based on molecular data) – *Prunus subhirtella* (this study).

Material examined – China, Guizhou Province, Guiyang. Lives on a dead stem of *Prunus subhirtella* (Rosaceae), 22 Dec. 2018, Lakmali S. Dissanayake, GHP03 (MFLU 19–0773, new host record), living culture MFLUCC 19–0469.

GenBank Numbers – ITS: MN707567, RPB2: MN987003, ACT: MN987004, TUB: MN987002.

Notes – There is a large overlap between our newly collected species (MFLU 19–0773) and *Rosellinia convexa* (holotype, GZUCC13005). Both species were collected in the same locality, but the holotype was collected from an unidentified host (Su et al. 2016a). Both strains have globose to subglobose, scattered ascomata, papillate ostioles, 8-spored, cylindrical, uniseptate asci with an apical ring bluing in Melzer’s reagent and brown ascospores with germ slits (Su et al. 2016a). However, the ascospores of our new collection are different from the holotype in having narrowly to broadly ellipsoidal-equilateral ascospores with broadly rounded ends, while the holotype has lemon-shaped to podiform ascospores with convex umbilical ends. Multi-gene (ITS, RPB, ACT, TUB) analysis herein, also shows that our collection clusters with other *Rosellinia convexa* strains (Fig. 169). According to the guidelines of Jeewon and Hyde (2016), we have analyzed the nucleotide differences within the rRNA gene region to further clarify the identification. In comparison of ITS regions (ITS1-5.8S-ITS2) from 738 nucleotides, there are 3 bp (0.40%) differences between MFLU 19–0773 and GZUCC13005. We confirmed that our new collection (MFLU 19–0773) is another record of *Rosellinia convexa* and a new host record from dead stems of *Prunus subhirtella* (Rosaceae) in China.

*Sordariomycetes orders incertae sedis*  
**Catabotryales** K.D. Hyde & Senan.  
**Catabotryaceae** Petr. ex M.E. Barr

This family was referred to different orders including Sordariales (Barr 1990b) then transferred to Xylariales by Hyde et al. (2000), while Kirk et al. (2001) placed it in Boliniiales based on its morphology. Miller and Huhndorf (2005) placed Catabotryaceae in Diaporthales based on a multi-gene analysis. Maharachchikumbura et al. (2015, 2016a) placed it in Amplitrnomatales as it clustered with Amplitromataceae in phylogenetic analyses, while Hyde et al. (2020) introduced a new order to accommodate it based on divergence time estimates.

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**Fig. 170** – *Rosellinia convexa* (MFLU 19–0773, new host record). a, b Habit of stromata on host surface. c Stromata in lateral close-up showing the smooth ectostroma with small papillate ostioles. d Stroma in vertical section showing the thick carbonaceous ectostroma. e Peridium. f Paraphyses. g–j Immature and mature asci. k Apical ring. l Ascal apical ring in Melzer’s reagent. m–o Immature and mature ascospore with cellular appendages (arrow). p Ascospore in dorso-lateral view showing a germ slit (arrows). q Ascospore with slimy sheath in Indian ink. Scale bars: b, c = 500 μm, d = 200 μm, e, f = 10 μm, g–j = 50 μm, k, l = 5 μm, m–q = 10 μm.

*Catabotrys* Theiss. & Syd.

The monotypic genus *Catabotrys* typified by *Catabotrys deciduum* was introduced by Theissen & Sydow (1915). *Catabotrys* is characterized by a sexual morph with superficial, reddish
brown to black, discoid to pulvinate, multi-loculate stromata, ascomata that are perithecial, globose enclosed in stromatic columns, asci that are 8-spored, unitunicate, broad cylindrical, short pedicellate, with a J-, discoid, apical ring and ascospores that are hyaline, 1-celled, ellipsoidal to cylindrical and the asexual morph is hyphomycetous with ellipsoidal to ovoid conidia. Currently, this genus includes three species as listed in Index Fungorum (2020).

**Catabotrys deciduum** (Berk. & Broome) Seaver & Waterston, Mycologia 38(2): 184 (1946)

**Facesoffungi number:** FoF06530

** Saprobic** on decaying wood of *Rhizophora mucronata*. Sexual morph: *Stromata* 1–3 mm high, 1–4 mm diameter (\(\bar{x} = 1.8 \times 2.3\) mm, \(n = 10\)), irregularly scattered, occasionally coalescing, conspicuous, multi-loculate, superficial, with base slightly penetrating the epidermis at regular intervals, discoid to pulvinate, reddish brown to black surface, scurfy, flat or slightly convex, composed of rather thin-walled, reddish yellow cells of *textura angularis*, *textura globosa* and *textura epidermoidea*. *Ascomata* perithecial, 390–500 µm high, 180–200 µm diameter (\(\bar{x} = 426 \times 197\) µm, \(n = 10\)), deeply imbedded in stromatic columns, globose, with a long, periphysate, ostiolar neck. *Ostioles* 120–200 × 35–45 µm (\(\bar{x} = 160 \times 40\) µm, \(n = 10\)). *Peridium* 40–55 µm (\(\bar{x} = 45\) µm, \(n = 5\)) thick, composed of several layers of compressed, reddish brown cells of *textura angularis*. *Paraphyses* 1–3 µm (\(\bar{x} = 1.7\) µm, \(n = 10\)), hypha-like, numerous, tapering towards the apex, not embedded in a gelatinous matrix. *Asci* 35–45 × 4–6 µm (\(\bar{x} = 41 \times 5\) µm, \(n = 10\)), 8-spored, unitunicate, broad cylindrical, short pedicellate, apically rounded or truncate, with a J- discoid, refractive, apical ring. *Ascospores* 7–9× 3–4 µm (\(\bar{x} = 8 \times 3.5\) µm, \(n = 10\)), uniseriate to bi-seriate, hyaline, 1-celled, ellipsoidal to cylindrical, smooth-walled, lacking a mucilaginous sheath. Asexual morph: Hyphomycetous, vegetative hyphae smooth-walled, septate, hyaline, up to 3 µm wide. *Conidiophores* micronematous, sympodial, weakly differentiated arising from vegetative hyphae. *Conidiogenous* cells reduced, holoblastic, polyblastic. *Conidia* ellipsoidal to ovoid, smooth surfaced, thick-walled, often with denticles or left over scars of detachment from conidiogenous cells, 3–5.5 × 2–5 µm (\(\bar{x} = 4.3 \times 3.4\) µm, \(n = 10\)), hyaline and 1-celled.

**Culture characteristics** – *Ascospores* germinating on seawater agar within 24 hours, germ tubes arising from both ends of the ascospores. Colonies on MEA, moderately growing, reaching 25–45 mm diameter after 25 days of incubation at room temperature, hyaline to cream and pale yellow at center margin, reverse creamy to pale yellow, velvety, circular, surface raised.

**Known distribution** (based on molecular data) – Chumphon, Thailand (Hyde et al. 2019), Panama (Miller and Huhndorf 2005), Puducherry, India (this study).

**Known hosts** (based on molecular data) – *Salacca* sp, oil palms (Hyde et al. 2019), *Rhizophora mucronata* (this study).

**Material examined** – India, Puducherry, Thengaithittu mangroves (11.5°N 79.5°E), on decaying wood of *Rhizophora mucronata* (Rhizophoraceae), 20 January 2016, B. Devadatha (AMH-10015), living culture, NFCCI-4395.

**GenBank numbers** – LSU: MN061355, SSU: MN173346, RP2B: MN546859, *tef1*: MN184789.

**Notes** – The monotypic genus *Catabotrys* is typified by *C. deciduum* and belongs to Catabotrydaceae (Petrak 1952) in Catabotryales (Hyde et al. 2020). Our collection of *Catabotrys deciduum* (AMH-10015) has similar morphological characters and measurements to *Catabotrys deciduum* (Seaver & Waterston 1946). The nucleotide sequence data of our collection of *Catabotrys deciduum* (NFCCI-4395) is 100% identical in the LSU region and less than 1% nucleotide differences in protein-coding gene regions with the sequence data of *C. deciduum* (SMH3436).
Fig. 171 – Phylogram generated from maximum likelihood analysis based on LSU, ITS, tef1 and RPB2 gene regions of Amplitromatales and related genera. Bootstrap support for maximum likelihood (ML, green), equal to or greater than 75 % and the values of Bayesian posterior probabilities (BIPP, purple) equal to or greater than 0.95 are given above each branch. The ex-type strains are in bold and new isolate is in blue. The tree is rooted with Xylaria hyphoxylon AFTOLID 51. The newly generated sequence is indicated in blue bold.
Fig. 172 – *Catabotrys deciduum* (AMH–10015, new geographical record). a Stromata on decaying wood of *Rhizophora mucronata*. b Horizontal section of stromata. c–e Longitudinal sections of ascomata. f Ostiole with periphyses. g Section of peridium comprising inner hyaline to pale brown cells of *textura angularis*. h Paraphyses and immature asci. h–j Immature and mature asci. k Ascus in Lugol’s iodine showing dextrinoid reaction. l–m Ascospores n Germinating ascospore Scale bars: b = 100 μm, c = 50 μm, d–n = 10 μm.
Further multi-gene phylogenetic analyses also showed that *Catabotrys deciduum* (NFCCI-4395) clustered together with *Catabotrys deciduum* strains in a monophyletic clade with high statistical support (ML100%, 1.00 BYPP) (Fig. 171). In the present study, we observed that *Catabotrys deciduum* (NFCCI-4395) produced an asexual morph with hyaline, ellipsoidal, 1-celled conidia in culture (MEA). *Catabotrys deciduum* is a saprobe, which commonly occurs on palms and larger monocots, with a pantropical distribution (Maharachchikumbura et al. 2016a). However, our collections of *Catabotrys deciduum* (AMH-10015) shows that it is frequent on intertidal decaying wood of *Rhizophora mucronata* thus extending its host range. Furthermore, its occurrence in Indian mangroves is a new country record and extends its geographical range.

**Agaricomycetes**

**Agaricomycetidae** Locq.

**Agaricales** Underw.

**Agaricaceae** Chevall.

Agaricaceae was established by Chevallier in 1826 (Chevallier 1826, Kirk 2008b). The classification of the family has changed considerably with the use of molecular dating results (Vellinga et al. 2004, 2011, Matheny et al. 2007, He et al. 2019b).
**Lepiota** (Pers.) Gray

Lepiota is a large genus of white-spored mushrooms of Agaricaceae which is commonly distributed in tropical and temperate countries (Dennis 1952). Based on its morphology, the genus was divided into six sections (Vellinga 2001). However, DNA based studies did not support the sections (Vellinga 2003, Liang et al. 2011). There were 37 species of *Lepiota* recorded in Thailand (Sysouphanthong et al. 2011b, 2012, 2013, 2016, Tibpromma et al. 2017). In this study, four species of *Lepiota* are new records to Thailand.

**Lepiota angusticystidiata** J.F. Liang & Z.L. Yang, Mycologia 110(3): 496 (2018)

Facesoffungi number: FoF06198

**Pileus** 15–40 mm diameter when young conical to parabolic, expanding to hemispherical to campanulate or umbonate, finally plano-convex, with inflexed margin; when young granulose or glabrous, greyish brown to light brown (6B4–5), soon breaking up into concolorous squamules around glabrous umbo towards the margin on white background; margin with white cortinate and connected with stipe when young, when mature squamulose, fringed. Lamellae free, crowded, ventricose, 3–6 mm wide, white, with colored floccose edge. Stipe 50–75 × 5–8 mm, cylindrical or slightly tapering to apex; covered with concolorous squamules as those on pileus, slightly crowded from base to annular zone, with white to yellowish white (4A2) background. Anulus an annular zone, with white fibrillose or cortinate. Context white in pileus, 2–3 mm wide; white to yellowish white (4A2) in stipe, hollow, with white fibrils in central cavity. Smell and Taste not observed. Spore print white.

Basidiospores 6.5–7 × 4–5 μm, avl x avw = 6.9 × 4.6 μm, Q = 1.4–1.7, avQ = 1.5, in side-view ellipsoid ovoid, in frontal view ellipsoid, hyaline, slightly thick-walled, dextrinoid, congophilous, cyanophilous, not metachromatic. Basidia 17–22 × 7–9 μm, clavate, 4-spored, hyaline, slightly thick walled. Cheilocystidia 16–26 × 7–9 μm, clavate, utriform to narrowly utriform, often fusiform, hyaline, slightly thick-walled, rarely with apical excrescence. Pileus covering a trichoderm made up of narrowly cylindrical elements with tapering to apex, 120–250 (–300) × 10–17 μm, sometimes septate at base, thick-walled, with parietal pale brown pigment, with concolorous short narrowly clavate elements under these long elements, 30–50 × 13–20 μm. Stipe covering of squamules a trichoderm similar to pileus covering. Clamp-connections present in all tissues.

Known distribution (based on molecular data) – China (Liang et al. 2018), northern Thailand (this study).

Known hosts (based on molecular data) – Terrestrial on soil or soil mixed with decayed leaves and wood (Liang et al. 2018, this study).

Material examined – Thailand, Chiang Mai Province, Mae Taeng District, Phadeng Village, N 19O 07' 13.7", E 98° 43' 52.9", alt. 905 m., 8 October 2008. P. Sysouphanthong, PS219 (MFLU 09–0202); Chiang Mai Province, Mae Taeng District, Pongduad Village, N 16° 06' 16.1", E 99° 43' 07.9", alt. 780-805 m., 18 July 2008. P. Sysouphanthong, PS86 (MFLU 090159); Chiangrai Province, Mae Jan District, Forest of Huay Kang Pla Village, N 20° 05' 28.1", E 99° 46' 52.7", alt. 512 m., 2 July 2008. P. Sysouphanthong, PS135 (MFLU 09–0208).

GenBank numbers – ITS (MFLU 09–0159 = MN240415, MFLU 09–02020 = MN240414, MFLU 09–0208 = MN240416).

Notes – From this study, all specimens of *Lepiota angusticystidiata* are recognized by greyish brown to light brown squamules covering the pileus and stipe, ellipsoid ovoid basidiospores, clavate to utriform cheilocystidia, a trichodermal pileus and stipe covering made up of narrowly cylindrical elements which taper at the apex with short narrowly clavate elements and clamp-connections (see Figs 175, 176).
Fig. 174 – Maximum likelihood phylogenetic tree based on nrITS sequences of Lepiota species. Dataset was analysed in RAXML7.2.6 (Stamatakis et al. 2008) with GTR+GAMMA as the model of evolution, and branch support was estimated over 1000 bootstrap partitions (BP) with the rapid bootstrap option. Bootstrap support ≥ 60% are indicated at the nodes. New sequences from this study are in blue bold. The GenBank accession numbers are indicated after species name. Abbreviation L= Lepiota, M = Macrolepiota. The tree is rooted with Macrolepiota orienrriex.
*Lepiota angusticystidiata* was described from southwest China and this species belongs to section *Ovisporae* (J.E. Lange) Kühner (Liang et al. 2018). All Thai specimens have larger basidiomata than the type specimen from China, but all other characteristics are similar. In nrITS analysis the Thai specimens clustered with the type and other specimens from China (see Fig. 174).

*Lepiota angusticystidiata* is a first record for Thailand and is distributed in deciduous forest of Chiang Mai and Chiang Rai of northern Thailand with an elevation around 500–900 m.

![Fresh basidiomata of *Lepiota angusticystidiata* on field.](image)

**Fig. 175** – Fresh basidiomata of *Lepiota angusticystidiata* on field. (a, b = MFLU 09–0202; c = MFLU 09–0159), (d = MFLU 09–0208). Scale bars: a–d= 10 mm.

*Lepiota attenuata* J.F. Liang & Zhu L. Yang, Mycologia 103(4): 821 (2011)

Facesoffungi number: FoF06197

*Pileus* 40–95 mm diameter campanulate or convex, expanding to plano-convex or planar, with straight to inflexed margin; with granulose or glabrous at center, sometime with patch-like at center, greyish brown to brownish orange (6C2-3), with centrically concolorous squamules or fibrilllose squamules around patch towards the margin, with radially reddish white fibrilllose between squamules towards the margin on white background; margin striate or sulcate at marginal zone, fringed, sometime attached with white remnants and concolorous squamules. *Lamellae* free, crowded, broadly ventricose, 5–12 mm wide, white, with colored eroded edge. *Stipe* 70–110 × 6–12 mm, cylindrical or slightly tapering to apex, covered with white to yellowish white (4A2) fibrilllose on white background. Annulus attached at upper site of stipe, membranous or white cortinate with greyish brown to brownish orange (6C2-3) squamules upper part. *Context* white in pileus, 4–7 mm wide; white to yellowish white (4A2) in stipe, hollow, with white fibrils in central cavity. *Smell and Taste* not observed. *Spore print* white.
Fig. 176 – Microcharacters of Lepiota angusticystidiata (MFLU 09–02020). a = Basidiospores, b = Cheilocystidia, c = A trichodermal pileus structure of pileus covering.

**Basidiospores** 16–19 × 4.8–5.2 μm, avl × avw = 17.32 × 5.02 μm, Q = 3.2–3.6, avQ = 3.45, in side-view ellipsoid penguin-shaped with straight adaxial side and a suprahilar depression, with attenuate to apex, in frontal view fusiform, hyaline, slightly thick-walled, dextrinoid, congophilous, cyanophilous, not metachromatic. **Basidia** 28–32 × 7–10 μm, clavate, 4-spored, rarely 2-spored, hyaline, thick walled. **Cheilocystidia** 25–28 × 6–10 μm, clavate to utriform or fusiform, rarely with apical excrescence, often with short clavate and septate, 6–10 × 4–6 μm. **Pileus covering** a trichoderm made up of narrowly cylindrical elements with tapering to apex, 120–300 × 7.5–11.5 μm, sometimes septate at base, thick-walled, with pale brown parietal pigment; with concolorous short clavate elements with septate under these long elements. Stipe covering not observed. **Clamp-connections** present in all tissues.

**Known distribution** (based on molecular data) – China (Liang et al. 2011), Northern Thailand (this study).

**Known hosts** (based on molecular data) – terrestrial on soil or soil mixed with decayed leaves and wood (Liang et al. 2011, this study).

**Material examined** – Thailand, Chiang Rai Province, Muang District, Mae Fah Luang University, N 18° 05' 59.1", E 102° 40' 22.9", alt. 488 m., 6 November 2008. P. Sysouphanthong, PS123 (MFLU 09–196); ibidem, 6 Nov. 2008. P. Sysouphanthong, PS126 (MFLU 09–0199).

**GenBank numbers** – ITS (MFLU 09–0196 = MN240413, MFLU 09–0199 = MN240412).

**Notes** – Thai specimens of Lepiota attenuata are characterized by greyish brown to brownish orange squamulose to fibrillose squamules on the pileus, a striate or sulcate pileus margin, with a white to yellowish white fibrillose stipe, with an annulus, penguin-shaped basidiospores, clavate to utriform cheilocystidia, a trichodermal pileus cover, and clamp-connections.
Lepiota attenuata was described from southern China by Liang et al. (2011). The species is placed in section Lepiota which comprises species with long basidiospores and a trichodermal pileus covering, and taxa similar to this species are Lepiota metalispora, Lepiota thrombophora, Lepiota cortinarius, Lepiota magnispora and Lepiota spheniscispora. Lepiota attenuata however, has basidiospores which are distinctly narrowed at the apex (Vellinga 2001, Wang 2004, Liang et al. 2011).

The nrITS sequences analysis showed that two Thai specimens clustered with type specimens from south China (see Fig. 174). The species is a new record for Chiang Rai of northern Thailand.

Fig. 177 – Fresh basidiomata of Lepiota attenuata on field. (a, b = MFLU 09–0196; c, d = MFLU 09–0199). Scale bars: a–d= 20 mm.

Lepiota magnispora Murrill, Mycologia 4: 237. 1912.
Facesoffungi number: FoF06195

Pileus 65 mm diameter, umbonate, with straight or slightly inflexed margin; covered with light brown to golden brown (5D5–7) granulose at center, with orange white to light orang (6A2–4) granulose, with crowded squamules towards the margin on white background; margin with white to light orange (6A4) floccules. Lamellae free, crowded, ventricose, up to 6 mm wide, white, with colored eroded edge. Stipe 85 × 5–10 mm, tapering to apex, covered with white to light orang (6A4) granulose, very crowded from base to annular zone, with light brown to golden brown (5D5–7) squamules at base zone on orange white (6A2) background. Annulus an annular zone with white
to light orang (6A4) granulose. Context white in pileus, 5 mm wide; white to orange white (6A2) in stipe, hollow, with white fibrils in central cavity. Smell and Taste not observed. Spore print white. Basidiospores 14–18 × 4.5–5.5 μm, avl × avw = 15.22 × 5.02 μm, Q = 2.8–3.4, avQ = 3.05, in side-view ellipsoid penguin-shaped with straight adaxial side and a suprahilar depression, with slightly attenuate to apex, in frontal view fusiform, hyaline, slightly thick-walled, dextrinoid, congophilous, cyanophilous, not metachromatic. Basidia 25–31 × 9–11 μm, clavate, 4-spored, rarely 2-spored, hyaline, thick walled. Cheilocystidia 23–38 × 6–12 μm, normally utriform or fusiform, rarely with clavate, often with 2–3 septate at base. Pileus covering a trichoderm made up of narrowly cylindrical elements with rounded apex, 175–310 × 11–15 μm, sometimes septate, with encrusted at base layer, thick-walled, with pale brown parietal pigment; with concolorous short clavate elements with septate under these long elements. Clamp-connections present in all tissues.

**Fig. 178** – Microcharacteristics of *Lepiota attenuata* (MFLU 09–0196). a Basidiospores. b Basidia and basidiomes. c Cheilocystidia. d A trichodermal pileus structure of pileus covering.

Known distribution (based on molecular data) – China, Germany, Netherlands, USA (Liang et al. 2011, Vellinga 2004), Thailand (this study).

Known hosts (based on molecular data) – Terrestrial on soil or soil mixed with decayed leaves and wood (this study).

Material examined – Thailand; Chiang Mai Province, Mae Taeng District, Phadeng Village, N 19° 07’ 13.7”, E 98° 43’ 52.9”, alt. 905 m., 9 July 2008. P. Sysouphanthong, PS51 (MFLU 09–0124).

GenBank numbers – ITS (MFLU 09–0124 = MN240411).
Notes – The description of *Lepiota magnispora* is based on a single collection. The species is characterized by a light brown to golden brown pileus with white to light orange floccose margins, white and free lamellae, a white to light orange floccose stipe, ellipsoid penguin-shaped, utriform or fusiform cheilocystidia, a trichodermal pileus cover made up of narrowly cylindrical elements with rounded apices and clamp-connections.

*Lepiota magnispora* is mostly found in temperate countries, and taxonomical history of the species was discussed in Vellinga (2000). With regards to distribution in Asia, *Lepiota magnispora* is distributed in China and Japan (Sysouphanthong et al. 2011a). In this study, *Lepiota magnispora* is also found in quite high elevation (905 m.) in north Thailand. Phylogenetic analysis based on nrITS sequences indicate that our specimen of *Lepiota magnispora* from Thailand is identical and clustered with those specimens from temperate countries such as Belgium, Canada, China and USA.

![Image of Lepiota magnispora](image.png)

**Fig. 179** – Fresh basidiomata of *Lepiota magnispora* on field. (a, b = MFLU 09–0124). Scale bars: a–b = 20 mm.

*Lepiota thrombophora* (Berk. & Broome) Sacc., Syll. fung. (Abellini) 5: 53 (1887)

Facingoffungi number: FoF06196

_Pileus_ 25 mm diameter, parabolic to hemispherical, with straight margin; covered with light brown to golden brown (5D5-7) granulose or crowded squamules at center, with pale orange to light orang (5A3-4) granulose to squamules towards the margin on white background; margin slightly sulcate, with white floccules or cortinate. _Lamellae_ free, crowded, ventricose, up to 4 mm wide, white, with colored eroded edge. _Stipe_ 45 × 4–6 mm, cylindrical or slightly tapering to apex; covered with white granulose or fibrillos, very crowded from base to annular zone, with pale orange to light orang (5A3-4) granulose to squamules at base zone on orange white (6A2) background. Annulus an annular zone with white fibrillos or cortinate. _Context_ white in pileus, 2 mm wide; white in stipe, hollow, with white fibrils in central cavity. _Smell and Taste_ not observed. _Spore print_ white.
Fig. 180 – Microcharacteristics of *Lepiota magnispora* (MFLU 09–0124). a Basidiospores. b Cheilocystidia. c A trichodermal pileus structure of pileus covering.

*Basidiospores* 10–14 × 3–5 μm, avl × avw = 11.8 × 4.1 μm, Q = 2.4–4.3, avQ = 2.89, in side-view ellipsoid penguin-shaped with straight adaxial side and a suprahilar depression, with slightly attenuate or rounded apex, in frontal view fusiform, hyaline, slightly thick-walled, not dextrinoid, congoophilous, cyanophilous, not metachromatic. *Basidia* 18–22 × 6–8 μm, clavate, 4-spored, sometime 2-spored, rarely 1-spored, hyaline, thick-walled. *Cheilocystidia* 18–30 × 10–12 μm, short clavate to clavate, often utriform or fusiform. *Pileus covering* a trichoderm made up of narrowly cylindrical elements with rounded apex, 110–190 × 12–15 μm, thick-walled, with pale brown parietal pigment; with concolorous short clavate elements under these long elements, 30–40 × 10–16 μm. *Stipe covering* in squamules a trichoderm similar to pileus covering. * Clamp-connections* present in all tissues.

Known distribution (based on molecular data) – China (Liang et al. 2011), Thailand (this study).

Known hosts (based on molecular data) – terrestrial on humus soil mixed with decayed leaves (Liang et al. 2011, this study).

Material examined – Thailand, Chiang Mai Province, Mae Rim District, Mae Sae Village, 18 July 2008, P. Sysouphanthong, PS89 (MFLU 09–0162).

*GenBank numbers*: ITS (MFLU 09–0162 = MN240410).

Notes – A specimen of a Thai *Lepiota thrombophora* was observed in a single young collection. It is characterized by pale orange to light orange granulose to squamules on the pileus, white and free lamellae, penguin-shaped basidiospores, clavate cheilocystidia, a trichoderm made up of narrowly cylindrical elements with rounded apex and clamp-connections (Figs 181, 182).

The microcharacters of the Thai specimen closely matches the holotype from Sri Lanka in shapes and size, including elements of the pileus covering being cylindrical with a rounded apex. However, specimens of *Lepiota thrombophora* from China has cylindrical elements with an
attenuated apex (Liang et al. 2011). In analysis of nrITS sequence data, the Thai specimen clustered with the two specimens from China (see Fig. 174).

*Lepiota thrombophora* is distributed in tropical Asia in China, India and Sri Lanka (Liang et al. 2011, Pegler & Young 1986, Natarajan & Manjula 1983, Sysouphanthong et al. 2011a). *Lepiota thrombophora* is a new record to Thailand.

![Image](image1.png)

*Fig. 181* – Fresh basidiomata of *Lepiota thrombophora* on field. (a, b = MFLU 09–0162). Scale bars: a–b= 10 mm.

![Image](image2.png)

*Fig. 182* – Microcharacteristics of *Lepiota thrombophora* (MFLU 09–0162). a = Basidiospores, b = Cheilocystidia, c = A trichodermal pileus structure of pileus covering.
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