Three new species of Dicephalospora from China as revealed by morphological and molecular evidences

Huan-Di Zheng¹, Wen-Ying Zhuang¹

¹ State Key Laboratory of Mycology, Institute of Microbiology, Chinese Academy of Sciences, Beijing 100101, China

Corresponding author: Wen-Ying Zhuang (zhuangwy@im.ac.cn)

Abstract
Three new species of Dicephalospora are introduced based on morphological characters and DNA sequence analyses (maximum parsimony and neighbor-joining methods), viz. D. albolutea, D. shennongjiana, and D. yunnanica. All of them lack mucilaginous caps at ascospore poles. Dicephalospora albolutea is distinguished by cream to yellowish white apothecia and slightly curved ascospores. Dicephalospora shennongjiana is characterized by yellow apothecia, elliptical-fusoid ascospores 19–22 × 7–8.8 μm, and J+ asci 130–150 × 14–16.5 μm. Dicephalospora yunnanica is distinguished by orange apothecia and fusoid ascospores 16.5–25.3 × 3.3–3.5 μm. Descriptions and illustrations of the new species as well as a key to the known species in the genus are provided.

Keywords
Morphology, phylogeny, species diversity, taxonomy

Introduction
Dicephalospora Spooner is a small genus established by Spooner (1987) with D. calochroa (Syd. & P. Syd.) Spooner as the type species. The poles of ascospores with a mucilaginous cap and J+ asci were treated as two important features to delimitate the genus, but a later study proved they are not reliable features at the generic level (Zhuang et al. 2016). The emended diagnostic characters of the genus are that apothecia erumpent or superficial, stipitate, yellow, orange, red to blackish, ectal excipulum of textura prismatica with refractive walls, medullary excipulum of textura intricata, asci J+ or J- in Melzer’s reagent, ascospores hyaline, subellipsoid to fusoid, guttulate, poles...
either with a mucilaginous cap or not, paraphyses filiform, straight or slightly curved at apex, and occurring on rotten wood, twigs, and leaf petioles (Zhuang et al. 2016). The genus was once treated as a member of Rutstroemiaceae (Kirk et al. 2008), Helotiaceae (Wijayawardene et al. 2017, 2018), or Sclerotiniaceae (Index Fungorum 2019). Including *Dicephalospora* in Helotiaceae is more reasonable in view of the phylogenetic studies of related groups in recent years (Han et al. 2014; Zhao et al. 2016).

Zhuang et al. (2016) carried out a comprehensive study on taxonomy of *Dicephalospora* in China and provided a key to the known species of the genus. Approximately, 10 species are currently accepted in the genus and nine of them have been found in China (Zhuang 1995a, 1995b, 1999; Verkley 2004; Zhuang et al. 2016). Dicephalosterol was discovered from the culture of *D. rufocornea* (Hosoya et al. 1999). This compound is a new testosterone $5\alpha$-reductase inhibitor and has a potential to be developed as a drug to prevent and cure prostatic hypertrophy (Hosoya et al. 1999). Additional information about utilization of the *Dicephalospora* spp. was rarely published maybe due to the minimal biomass in nature, difficulty of getting pure culture, and slow-growth if cultured.

During the examinations of helotialean fungi from China, three species fit well with the emended generic concept of *Dicephalospora* (Zhuang et al. 2016). However, new collections are found to differ from hitherto known species of *Dicephalospora*. To confirm their affinities and investigate their relationships with other species, phylogenetic analyses were conducted based on the internal transcribed spacers and 5.8S of nuclear ribosomal DNA (ITS). The results support their placement within the genus and their distinctions from any known species.

**Materials and methods**

Specimens were collected, recorded, and photographed by a Canon PowerShot G16 digital camera in the field. Descriptions of gross morphology and substrate were according to field notes and photos. Dried apothecia were rehydrated with distilled water and sectioned at a thickness of 15–20 μm with a Yidi YD-1508A freezing microtome (Jinhua, China). Measurements were taken from longitudinal sections and squash mounts in lacto-phenol cotton blue solution using an Olympus BH-2 microscope (Tokyo, Japan). Iodine reactions of ascal apparatus were tested with or without 3% KOH solution pretreatment in Melzer’s reagent and Lugol’s solution (Baral 2009). Microscopic images were taken using a Canon G5 digital camera (Tokyo, Japan) attached to a Zeiss Axioskop 2 Plus microscope (Göttingen, Germany). Voucher specimens were deposited in the Herbarium Mycologicum Academiae Sinicae (HMAS). Names of the new species were formally registered in the database Fungal Names (http://www.fungalinfo.net/fungalname/fungalname.html).

Pure cultures were obtained from some specimens following the method provided by Webster and Weber (2001) and preserved in the State Key Laboratory of Mycology, Institute of Microbiology, Chinese Academy of Sciences.

Genomic DNA was extracted from dried apothecia or pure culture, using Plant Genomic DNA Kit (TIANGEN Biotech. Co., Beijing, China). ITS region was amplified
and sequenced using the primer pair ITS1/ITS4 (White et al. 1990). PCR reactions had a final volume of 30 μl, containing 15 μl 2×Taq MasterMix (Beijing CWBiotech, China), 1.5 μl of each primer (10 mM), 2 μl DNA, and 10 μl deionized water. PCR reactions were carried out in an Applied Biosystems 2720 thermocycler (Foster City, CA, USA) under the following conditions: 94 °C for 5 min, followed by 35 cycles of 94 °C for 30 s, 53 °C for 30 s and 30 s at 72 °C, and a final extension of 72 °C for 10 min. The PCR products were purified and sequenced at Beijing Tianyi Huiyuan Bioscience and Technology, China.

Newly generated sequences were assembled and edited using BioEdit 7.0.5.3 (Hall 1999) or SeqMan (DNASTAR, Lasergene 7.1.0). The new sequences were deposited in GenBank and additional sequences were downloaded from GenBank (Table 1).

### Table 1. Sequences used in this study.

| Species                                | Specimen/strain | ITS          |
|----------------------------------------|-----------------|--------------|
| *Chlorosplenium chlora* (Schwein.) M.A. Curtis | HMAS 266518     | MK425599     |
|                                        | HMAS 279692     | MK425600     |
| *Ciboria bachiata* (Zopf) N.F. Buchw.  | CBS 312.37      | KF859931     |
| *Ciborinia folicola* (E.K. Cash & R.W. Davidson) Wherzel | 1932.H         | Z80892       |
| *Dicephalospora albolutea* H.D. Zheng & W.Y. Zhuang | HMAS 279693     | MK425601     |
| *Dicephalospora aurantiaca* (W.Y. Zhuang) W.Y. Zhuang & Z.Q. Zeng | HMAS 61850      | DQ986486     |
| *Dicephalospora chrysotricha* (Berk.) Verkley | ICMP:19950      | KF727410     |
|                                        | ICMP:19952      | KF727411     |
| *Dicephalospora dentata* Xiao X. Liu & W.Y. Zhuang | HMAS 266694     | KP204263     |
| *Dicephalospora huangshanica* (W.Y. Zhuang) W.Y. Zhuang & Z.Q. Zeng | HMAS 74836      | DQ986485     |
|                                        | HMAS 81364      | DQ986844     |
|                                        | HMAS 279694     | MK425602     |
| *Dicephalospora rufocornea* (Berk. & Broome) Spooner | HMAS 75518      | DQ986480     |
|                                        | 10106           | KU668565     |
|                                        | HMAS 279695     | MK425603     |
|                                        | HMAS 279696     | MK425604     |
|                                        | HMAS 279697     | MK425605     |
| *Dicephalospora shennongjiana* H.D. Zheng & W.Y. Zhuang | HMAS 279698     | MK425606     |
| *Dicephalospora yunnanica* H.D. Zheng & W.Y. Zhuang | HMAS 279699     | MK425607     |
|                                        | HMAS 279700     | MK425608     |
|                                        | HMAS 279701     | MK425609     |
| *Hymenoscyphus fructigenus* (Bull.) Gray | CBS650.92       | GU586933     |
|                                        | HMAS 75893      | JX977144     |
| *Lachnum pygmaeum* (Fr.) Bres.          | ARON 2924.S     | AJ430215     |
| *Lachnum spartinae* S.A. Cantrel        | SAP 138         | AF422970     |
| *Lambertella corni-maris* Höhn.         | CLX 3892        | KC958560     |
|                                        | CLX 4075        | KC958562     |
| *Lanzia allantospora* (Dennis) Spooner  | PRJ D804        | AFY755334    |
| *Lanzia luteovirens* (Roberge ex Desm.) Dumont & Korf | 1823            | KC533545     |
| *Moellerodiscus lentus* (Berk. & Broome) Dumont | 7818            | KU668564     |
|                                        | 10544           | KU668566     |
| *Monilinia fructicola* (G. Winter) Honey | MO-3D           | JN001480     |
|                                        | RS10            | JF325841     |
| *Rutstroemia firma* (Pers.) P. Karst.   | 2089.1          | Z80893       |
|                                        | 2089            | KC533547     |
| *Sclerotinia sclerotiorum* (Lib.) de Bary | 2              | KF148605     |
|                                        | 6              | KF148609     |

* Numbers in bold indicate sequences produced by this study.
**Lachnum pygmaeum** (Fr.) Bres. and *L. spartinae* S.A. Cantrel were chosen as outgroup taxa. The ITS sequence matrix was aligned and manually edited using BioEdit 7.0.5.3 (Hall 1999). Phylogenetic analyses were performed using maximum parsimony (MP) and neighbor-joining (NJ) methods with PAUP* 4.0b10 and parameters were set according to Zheng and Zhuang (2015). The topological confidence of the NJ and MP trees was assessed with bootstrap analysis using 1,000 replications, each with 10 replicates of random stepwise addition of taxa. The resulting trees were viewed via TreeView 1.6.6 (Page 1996).

**Results**

**Phylogenetic analyses**

The ITS dataset included 37 sequences from eight *Dicephalospora* species, 11 related fungi and two outgroup taxa. The final alignment resulted in 634 characters including gaps, of which 252 were parsimony-informative, 38 were variable and parsimony-uninformative, and 344 were constant. In the MP analysis, eight most parsimonious trees were generated (tree length = 790, consistency index = 0.5899, homoplasy index = 0.4101, retention index = 0.8126, rescaled consistency index = 0.4793) and one of them was shown in Figure 1. MP and NJ bootstrap proportions (BP) greater than 50% were labeled at the nodes.

From topology of the phylogenetic tree (Fig. 1), *Dicephalospora* species clustered together with a medium supporting value (56% MPBP). The three putative new species were clearly distinct from the known and sequenced species of the genus. *Dicephalospora albolutea* appeared as an independent lineage distinct from any other members of the genus. *Dicephalospora shennongjiana* was resolved as a sibling species of *D. huangshanica* (97% MPBP and 99% NJBP). ITS sequences of the three collections of *D. yunnanica* were identical and formed a well-supported group with *D. aurantiaca* (100% MPBP and 100% NJBP).

**Taxonomy**

*Dicephalospora albolutea* H.D. Zheng & W.Y. Zhuang, sp. nov.

Fungal Names FN570602

Figure 2

**Etymology.** The specific epithet refers to the color of apothecia.

**Holotype.** CHINA. Yunnan Province, Binchuan County, Jizu Mountain, alt. 2500 m, on rotten leaf veins, 21 September 2017, H.D. Zheng, X.C. Wang, Y.B. Zhang & Y. Zhang 11613 (HMAS 279693, ITS GenBank accession number: MK425601).

**Description.** *Apothecia* scattered, discoid, stipitate, with even margin, 1–2.5 mm in diameter; hymenium surface cream to yellowish white; receptacle surface concolorous.
Three new species of *Dicephalospora*

Ectal excipulum of textura prismatica, 20–70 μm thick, cells somewhat thick- and glassy-walled, 16.5–40 × 5.5–11 μm. Medullary excipulum of textura porrecta and textura intricata, 25–275 μm thick, hyphae hyaline, thin-walled, 2.5–5 μm wide. Subhymenium not distinguishable. Hymenium 165–175 μm thick. Asci unitunicate, arising from simple septa, 8-spored, cylindric-clavate, J+ in Melzer’s reagent and Lugol’s solution without KOH pretreatment, visible as two blue lines, 140–156 ×

**Figure 1.** One of the MP trees inferred from ITS sequences. Bootstrap support values (≥50%) of MP and NJ are shown at nodes from left to right. New proposed species are shown in bold. New species are in bold. Sequences derived from holotypes are marked with an asterisk (*).
Figure 2. Dicephalospora albolutea (HMAS 279693, holotype). a fresh apothecia on natural substrate b longitudinal section of apothecium c structure of margin and hymenium d structure of flank e asci f IKI reaction of apical rings g ascospores. Mounting media: b–e, g lacto-phenol cotton f lugol’s solution. Scale bars: 5 mm (a); 200 μm (b); 20 μm (c, d); 10 μm (e, f); 5 μm (g).

9.5–10.5 μm. Ascospores sausage-shaped to subfusoid, with anterior end rounded and posterior end narrower, slightly curved, aseptate, hyaline, smooth, lacking a gel cap at each end, multiguttulate, with a dark-stained area when mounted in cotton blue solution, biseriate, 26–31 × 3.8–5.0 μm. Paraphyses filiform, straight, slightly enlarged
Three new species of *Dicephalospora* at apex, hyaline, septe, 3–3.5 μm broad at upper portion and 1.5–2 μm below, equal to or very slightly exceeding the asci.

**Notes.** The diagnostic features of *D. albolutea* are cream to yellowish white apothecia and sausage-shaped ascospores. The apothecial color of earlier known *Dicephalospora* species varied from yellow, orange, red to dark, but never as pale as that in *D. albolutea*. *Dicephalospora calochroa* (Syd. & P. Syd.) Spooner is somewhat similar in length of asci and ascospores, but differs by vivid orange apothecia, wider asci (125–150 × 12–15 μm) and ascospores (20–25 × 6–8 μm), which are pointed at both ends (Spooner 1987). *Dicephalospora albolutea* differs from any investigated species by at least 45 bp in sequences of ITS region, and appeared as an independent lineage in the phylogenetic tree (Fig. 1), which further confirmed its distinction from others in the group.

**Dicephalospora shennongjiana** H.D. Zheng & W.Y. Zhuang, sp. nov.
Fungal Names FN570603
Figure 3

**Etymology.** The specific epithet refers to the type locality of the fungus.

**Holotype.** CHINA. Hubei Province, Shennongjia, Shennongyuan, alt. 2250 m, on stromatized dead vine, 15 Sept 2014, H.D. Zheng, Z.Q. Zeng, W.T. Qin & K. Chen 9589 (HMAS 279698, ITS GenBank accession number: MK425606).

**Description.** *Apothecia* scattered, discoid to flat, stipitate, with even margin, 0.5–0.8 mm in diameter; hymenium surface greenish yellow; receptacle surface slightly darker. *Ectal excipulum* of textura prismatica, 15–40 μm thick, cells hyaline to pale brownish, somewhat thick- and glassy-walled, 10–20 × 4–11 μm. *Medullary excipulum* of textura intricata, 25–110 μm thick, hyphae hyaline, thin-walled, 2–4 μm wide. *Subhymenium* about 15 μm thick. *Hymenium* 170–180 μm thick. *Asci* arising from simple septa, unitunicate, 8-spored, clavate, J+ in Melzer’s reagent and Lugol’s solution without KOH pretreatment, visible as two blue lines, 130–150 × 14–16.5 μm. *Ascospores* elliptical-subfusoid, aseptate, hyaline, smooth, lacking a gel cap at each end, multiguttulate, with a dark-stained area when mounted in cotton blue solution, uniseriate, 19–22 × 7–8.8 μm. *Paraphyses* filiform, slightly enlarged at apex, hyaline, septe, branched and tangled near apex, 3–3.5 μm broad at upper portion and 1.5–2 μm below, exceeding the asci by 10–20 μm.

**Notes.** The new species can be distinguished from other species by shape of ascospores and tangled paraphyses apices. *Dicephalospora damingshanica* has a similarly shaped ascospore, but larger (22–32 × 9–12.7 μm), and with a hyaline mucilaginous cap at both ends (Zhuang 1999). Phylogenetically, *D. shennongjiana* is closely related to *D. huangshanica*, but the latter differs by red apothecia, smaller asci (89–96 × 9.5–11 μm), fusoid ascospores (18–26 × 4–5 μm) (Zhuang 1995a), and 22 bp divergence in ITS region.
**Figure 3.** *Dicephalospora shennongjiana* (HMAS 279698, **holotype**) a fresh apothecia on natural substrate b dried apothecia c longitudinal section of apothecium d structure of margin, flank and hymenium e asci f IKI reaction of apical rings g ascospores in an ascus h ascospores. Mouting media: c–e, g, h lacto-phenol cotton f fugol’s solution. Scale bars: 2 mm (a); 0.4 mm (b); 200 μm (c); 20 μm (d); 10 μm (e–h).

*Dicephalospora yunnanica* H.D. Zheng & W.Y. Zhuang, sp. nov.
Fungal Names FN570604
Figure 4

**Etymology.** The specific epithet refers to the type locality of the fungus.

**Holotype.** CHINA. Yunnan Province, Maguan County, Dabao Village, alt. 1565 m, on rotten leaf rachis, 13 August 2016, X.H. Wang, S.H. Li, H.D. Zheng & S.C. Li YN16-108 (HMAS279699, ITS GenBank accession number: MK425607).

**Description.** *Apothecia* scattered, discoid, stipitate, with even margin, 0.8–2.0 mm in diameter; hymenium surface bright yellow to orange; receptacle surface paler. *Ectal excipulum*
Three new species of *Dicephalospora*

**Figure 4.** *Dicephalospora yunnanica* (HMAS 279699, holotype) a fresh apothecia on natural substrate b dried apothecia c longitudinal section of apothecium d structure of margin, flank and hymenium e, f asci g IKI reaction of apical rings h ascospores. Mouting media: c–e, h lacto-phenol cotton f, g lugol’s solution. Scale bars: 5 mm (a); 2 mm (b); 200 μm (c); 20 μm (d); 10 μm (e–g); 5 μm (h).

of textura prismatica, 22–60 μm thick, cells hyaline, somewhat thick- and glassy-walled, 7–20 × 5–7 μm. *Medullary excipulum* of textura intricata, 30–230 μm thick, hyphae thin-walled, 2–5 μm wide. *Subhymenium* not distinguishable. *Hymenium* 100–115 μm thick. *Asci* arising from simple septa, unitunicate, 8-spored, cylindric-clavate, J+ in Melzer’s reagent and Lugol’s solution without KOH pretreatment, visible as two faint blue lines, 85–100 × 7.5–8.5 μm. *Ascospores* fusoid, aseptate, with one side very slightly flattened and pointed at ends, hyaline, smooth, lacking a gel cap at each end, multiguttulate, biseriate, 16.5–25.3 × 3.3–3.5 μm. *Paraphyses* filiform, slightly enlarged at apex, straight or sometimes slightly curved at the apical portion, hyaline, septate, 2.5–4 μm broad at upper portion and 1.5–2 μm below, slightly exceeding the asci by about 5 μm.
Additional specimens examined. CHINA. Yunnan Province, Maguan County, Xiaobaozi Town, alt. 1550 m, on rotten leaf rachis, 13 August 2016, X.H. Wang, S.H. Li, H.D. Zheng & S.C. Li YN16-135 (HMAS 279700); Maguan County, Pojiao Village, alt. 1450 m, on rotten leaf rachis, 14 August 2016, X.H. Wang, S.H. Li, H.D. Zheng & S.C. Li YN16-165 (HMAS 279700).

Notes. Dicephalospora yunnanica shares similar gross morphology with and appeared to be sister of D. aurantiaca in the phylogenetic tree (Fig. 1). However, D. aurantiaca has larger asci (93–103 × 9.7–10.5 μm) and ascospores (21–26 × 4–4.8 μm), as well as obviously curved paraphysis apices (Zhuang 1995a; Zhuang et al. 2016). Concerning the DNA sequence data, the three collections of D. yunnanica share exactly the same sequences, while the closest species D. aurantiaca showed 75 bp divergence (including 32 gaps) for ITS.

A taxonomic key to the known species of Dicephalospora

1 Receptacle surface covered with hairs ........................................... D. chrysotricha
– Receptacle surface without hairs .................................................. 2
2 Apothecial margin dentate ............................................................. D. dentata
– Apothecial margin even ................................................................. 3
3 Hymenium surface cream to yellowish white when fresh ............. D. albolutea
– Hymenium surface darker in color ................................................. 4
4 Hymenium surface red or dark red ............................................... D. huangshanica
– Hymenium surface lacking of a red tint ........................................ 5
5 Paraphyses with darkly pigmented contents ............................... D. phaeoparaphysis
– Paraphyses without darkly pigmented contents ................................ 6
6 Ascospores with a gel cap at each end .......................................... 7
– Ascospores lacking of a gel cap at each end ...................................... 10
7 Ascospores 9–12.7 μm wide ......................................................... D. damingshanica
– Ascospores less than 9 μm wide ................................................... 8
8 Asci J−, ascospores 20–28 × 4.5–5.7 μm ....................................... D. pinglongshanica
– Asci J+ ......................................................................................... 9
9 Ascospores 23–27(–29) × 6.5–7.5 μm ............................................. D. calochroa
– Ascospores (27–)32–39 × 4–5.5(–6) μm .......................................... D. rufocornea
10 Ascospores constricted in the middle, 20–27 × 4–5 μm .............. D. contracta
– Ascospores not constricted in the middle ......................................... 11
11 Ascospores 19–22 × 7–8.8 μm ....................................................... D. shennongjiana
– Ascospores less than 7 μm wide .................................................... 12
12 Ascospores 16.5–25.3 × 3.3–3.5 μm, paraphyses straight ........... D. yunnanica
– Ascospores 21–26 × 4–4.8 μm, paraphyses curved at apex .......... D. aurantiaca
Three new species of Dicephalospora

Discussion

Identification of Dicephalospora species is mainly based on morphological features, such as color of apothecia, anatomic structure, and characteristics of asci and ascospores. DNA sequence data are sometimes considered, which play an important role in the delineation of fungal species (Hibbett et al. 2016; Jeewon and Hyde 2016). In the present study, three new species were introduced based on morphology and ITS phylogeny. So far, the genus comprises 13 species, of which 12 have been reported from China. Dicephalospora chrysotricha (Berk.) Verkley originally described from, and endemic to, New Zealand, is the only exception and known only from the type locality (Verkley 2004).

In the phylogenetic analyses, only some species possessing fusoid to sausage-shaped and elliptic-subfuscoid ascospores were involved due to limitation of the available sequences. The ITS barcodes seem to be useful for distinguishing Dicephalospora species, as they grouped as well-separated clades (Fig. 1). Seven of the eight species were together receiving moderate statistic supports (86% MPBP and 80% BIPP) and formed the core group. However, D. chrysotricha joined them as a distantly separated lineage with very low support (Fig. 1, 56% MPBP). Dicephalospora chrysotricha is distinct from any other taxa of the genus in having hair-like projections on receptacle surface. Dicephalospora chrysotricha was previously treated as a member of Trichopeziza Fuckel (Saccardo 1889) and then Chlorosplenium Fr. (Dennis 1961). The transfer of this species to Dicephalospora might have been because of presence of polar mucilaginous caps of ascospores and the more or less similar ectal excipulum structure except for hairs (Verkley 2004). However, it does not fit well the generic concept of Dicephalospora. Further study is required to clarify the taxonomic position of this fungus.

As to the phylogenetic position of Dicephalospora, Figure 1 shows its close relationship with Hymenoscyphus Gray, which agrees with the treatment of Wijayawardene et al. (2017). Similar results were also achieved in other recent studies (Han et al. 2014; Zhao et al. 2016). In the phylogenetic study of Hyaloscyphaceae and related helotialean cup-fungi, D. huangshanica and D. rufocornea grouped together with some genera of Helotiaceae, such as Hymenoscyphus, Crocierea Fr. and Cudoniella Sacc., as a highly supported clade in the maximum-likelihood tree inferred from combined sequence data of ITS, the large subunit nrDNA gene (LSU), the second largest subunit of RNA polymerase II gene (RPB2), and mitochondrial small subunit (mtSSU) (Han et al. 2014). Zhao et al. (2016) carried out phylogenetic analyses of Lambertella Höhn. and allied genera including Dicephalospora and Hymenoscyphus, as inferred from ITS, LSU and RPB2 sequence data. In their phylogenetic trees, D. rufocornea was also associated with the clade consisting of Hymenoscyphus species. In view of the above results, close relationship of Dicephalospora with genera of Helotiaceae is obvious. Comprehensive work containing more genera and more genes are required to obtain an accurate conclusion on phylogenetic placement of Dicephalospora.
Acknowledgements

This work was supported by the National Natural Science Foundation of China (nos. 31770019, 31570018). The authors thank all co-collectors of specimens examined in this study for their invaluable help during the field work.

References

Baral H-O (2009) Iodine reaction in Ascomycetes: why is Lugol's solution superior to Melzer's reagent? http://www.gbif-mycology.de/HostedSites/Baral/IodineReaction.htm [Accessed on: 2011-3-18]

Dennis RWG (1961) Some inoperculate Discomycetes from New Zealand. Kew Bulletin 15: 293–320. https://doi.org/10.2307/4109373

Hall TA (1999) BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium Series 41: 95–98.

Han JG, Hosoya T, Sung GH, Shin HD (2014) Phylogenetic reassessment of Hyaloscyphaceae sensu lato (Helotiales, Leotiomycetes) based on multigene analyses. Fungal Biology 118: 150–167. https://doi.org/10.1016/j.funbio.2013.11.004

Hibbett D, Abarenkov K, Kõljalg U, Öpik M, Chai B, Cole J, Wang Q, Crous P, Robert V, Helgason T, Herr JR, Kirk P, Lueschow S, O'Donnell K, Nilsson RH, Oono R, Schoch C, Smyth C, Walker DM, Porras-Álfaro A, Taylor JW, Geiser DM (2016) Sequence-based classification and identification of Fungi. Mycologia 108: 1049–1068. https://doi.org/10.3852/16-130

Hosoya T, Hamano K, Sugano M, Ogura Y, Hatano E, Hamada T (1999) Discovery of dicephalosterol, a new testosterone 5α-reductase inhibitor, and some new mycological aspects of its producer, Dicephalospora rufocornea (Sclerotiniaceae, Discomycetes). Mycoscience 40: 525–529. https://doi.org/10.1007/BF02461030

Index Fungorum (2019) Index Fungorum. http://www.indexfungorum.org/names/Names.asp [Accessed on: 2019-2-12]

Jeewon R, Hyde KD (2016) Establishing species boundaries and new taxa among fungi: recommendations to resolve taxonomic ambiguities. Mycosphere 7: 1669–1677. https://doi.org/10.5943/mycosphere/7/11/4

Kirk PM, Cannon PF, Minter DW, Stalpers JA (2008) Dictionary of the Fungi (10th edn). CABI, Wallingford, 771 pp.

Page RDM (1996) Treeview: an application to display phylogenetic trees on personal computers. Computer Applications in the Biosciences 12: 357–358. https://doi.org/10.1093/bioinformatics/12.4.357

Saccardo PA (1889). Discomyceteae et Phymatosphaeriaceae. Sylloge Fungorum 8: 1–1143.

Spooner BM (1987) Helotiales of Australasia: Geoglossaceae, Orbiliaceae, Sclerotiniaceae, Hyaloscyphaceae. Bibliotheca Mycologica 116: 1–711.

Verkley GJM (2004) Redisposition of Chlorosplenium chrysotrichum to the genus Dicephalospora (Sclerotiniaceae, Ascomycota). Sydowia 56: 343–348.
Three new species of *Dicephalospora*

Webster J, Weber RWS (2001) Teaching techniques for mycology: 15. Fertilization and apothecium development in *Pyronema domesticum* and *Ascobolus furfuraceus* (Pezizales). Mycologist 15: 126–131. https://doi.org/10.1016/S0269-915X(01)80035-5

White T, Bruns TD, Lee A, Taylor JW (1990) Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. In: Innis MA, Gelfand DH, Sninsky JJ, White TJ (Eds) PCR Protocols: A Guide to Methods and Applications. Academic Press, San Diego, 315–322. https://doi.org/10.1016/B978-0-12-372180-8.50042-1

Wijayawardene NN, Hyde KD, Rajeshkumar KC, Hawksworth DL, Madrid H, Kirk PM, Braun U, Singh RV, Croux M, Kukwa M, Lücking R, Kurtzman CP, Yurkov A, Haelewaters D, Aptroot A, Lumbsch HT, Timdal E, Ertez D, Etayo J, Phillips AJL, Groenewald JZ, Papizadeh M, Sellmann L, Dayarathe MC, Weerakoon G, Jones EBG, Sutrong S, Tian Q, Castañeda-Ruiz RF, Bahkali AH, Pang K-L, Tanaka K, Dai DQ, Sakayaroj J, Hujslová M, Lombard L, Shenoy BD, Suja A, Maharachchikumbura SSN, Thambugala KM, Wannasinghe DN, Sharma BO, Gaikwad S, Pandit G, Zucconi L, Onofri S, Egidi E, Raja HA, Kodsueb R, Cáceres MES, Pérez-Ortega S, Fiuza PO, Monteiro JS, Vasilyeva LN, Shivas RG, Prieto M, Wedin M, Olariaga I, Lateef AA, Agrawal Y, Fazeli SAS, Amoozegar MA, Zhao GZ, Pfiegl er WP, Sharma G, Oset M, Abdel-Wahab MA, Takamatsu S, Bensch K, de Silva NI, De Kesel A, Karunarathna A, Boonmee S, Pfister DH, Lu Y-Z, Luo Z-L, Boonyuen N, Daranagama DA, Senanayake IC, Jayasiri SC, Samarako nn MC, Zeng X-Y, Doilm M, Quijada L, Rampadarath S, Heredia G, Dissanayake AJ, Jayawardana RS, Perera RH, Tang LZ, Phukhamsakda C, Hernández-Restrepo M, Ma X, Tibpromma S, Gusmao LFP, Weerahewa D, Karunarathna SC (2017) Notes for genera: Ascomycota. Fungal Diversity 86: 1–594. https://doi.org/10.1007/s13225-017-0386-0

Wijayawardene NN, Hyde KD, Lumbsch HT, Liu JK, Maharachchikumbura SSN, Ekanayaka AH, Tian Q, Phookamsak R (2018) Outline of *Ascomycota*: 2017. Fungal Diversity 88(1): 167–263. https://doi.org/10.1007/s13225-018-0394-8

Zhao YJ, Hosaka K, Hosoya T (2016) Taxonomic re-evaluation of the genus *Lambertella* (Rustroemiaceae, Helotiales) and allied stroma-forming fungi. Mycological Progress 15: 1215–1228. https://doi.org/10.1007/s11557-016-1225-5

Zheng HD, Zhuang WY (2015) Five new species of *Hymenoscyphus* (Helotiaceae, Ascomycota) with notes on the phylogeny of the genus. Mycotaxon 130: 1017–1038. https://doi.org/10.5248/130.1017

Zhuang WY (1995a) A few petiole-inhabiting discomycetes in China. Mycosistema 7: 13–17.

Zhuang WY (1995b) Some new species and new records of discomycetes in China. V. Mycotaxon 56: 31–40.

Zhuang WY (1999) Discomycetes of tropical China. VI. Additional species from Guangxi. Fungal Diversity 3: 187–196.

Zhuang WY, Zeng ZQ, Liu XX (2016) Taxonomic revision of the genus *Dicephalospora* (Helotiales) in China. Mycosistema 35: 791–801. https://doi.org/10.13346/j.mycosystema.160047