Diaporthe is paraphyletic

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Abstract: Previous studies have shown that our understanding of species diversity within *Diaporthe* (syn. *Phomopsis*) is limited. In this study, 49 strains obtained from different countries were subjected to DNA sequence analysis. Based on these results, eight new species names are introduced for lineages represented by multiple strains and distinct morphology. Twelve *Phomopsis* species previously described from China were subjected to DNA sequence analysis, and confirmed to belong to *Diaporthe*. The genus *Diaporthe* is shown to be paraphyletic based on multi-locus (LSU, ITS and *TEF1*) phylogenetic analysis. Several morphologically distinct genera, namely *Mazzantia, Ophiodiaporthe, Pustulomyces, Phaeocytostroma,* and *Stenocarpella,* are embedded within *Diaporthe s. lat.,* indicating divergent morphological evolution. However, splitting *Diaporthe* into many smaller genera to achieve monophyly is still premature, and further collections and phylogenetic datasets need to be obtained to address this situation.

Key words: Ascomycota Diaporthales Phomopsis phylogeny taxonomy

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INTRODUCTION

Species of Diaporthe are known as important plant pathogens, endophytes or saprobes (Udayanga et al. 2011, Gomes et al. 2013). They have broad host ranges, and occur on many plant hosts, including cultivated crops, trees, and ornamentals (Diogo et al. 2010, Thompson et al. 2011, Gomes et al. 2013, Huang et al. 2015). Some Diaporthe species are responsible for severe diebacks, cankers, leaf-spots, blights, decay or wilts on different plant hosts, several of which are economically important (Mostert et al. 2001, Van Rensburg et al. 2006, Thompson et al. 2011, Gomes et al. 2013), leading to serious diseases and significant yield losses (Santos et al. 2011). For example, Diaporthe helianthi is the cause of one of the most important diseases of sunflower (Helianthus annuus) worldwide, and has reduced production by up to 40 % in Europe (Masirevic & Gulya 1992, Thompson et al. 2011). Diaporthe neoviticola and D. vitimegaspora, the causal agents of leaf-spot and swelling arm, are known as severe pathogens of grapevines (Vitis vinifera) (Van Niekerk et al. 2005). Úrbez-Torres et al. (2013) indicated that D. neoviticola was one of the most prevalent fungi isolated from grapevine perennial cankers in declining vines. Diaporthe scabra has been reported causing cankers and dieback on London plane (Platanus acerifolia) in Italy (Grasso et al. 2012). Symptoms of umbel browning and necrosis caused by D. angeliace

have been regularly observed on carrots in France, resulting in seed production losses since 2007 (Ménard et al. 2014). Avocado (Persea americana), cultivated worldwide in tropical and subtropical regions, is threatened by branch cankers and fruit stem-end rot diseases caused by D. foeniculina and D. sterilis (Guarnaccia et al. 2016). Furthermore, species of Diaporthe are commonly introduced into new areas as endophytes or latent pathogens along with plant produce. For instance, Torres et al. (2016) reported D. rudis causing stemend rot in avocados in Chile, which was imported via avocado fruit from California (USA). Some endophytes have been shown to act as opportunistic plant pathogens. Diaporthe foeniculina (syn. P. theicola), which is a common endophyte, has been shown to cause stem and shoot cankers on sweet chestnut (Castanea sativa) in Italy (Annesi et al. 2015, Huang et al. 2015). Because of this unique ecology and potential role as plant pathogens, it is of paramount importance to accurately identify species of Diaporthe to facilitate disease surveillance, control, and trade.

The initial species concept of *Diaporthe* based on the assumption of host-specificity, resulted in the introduction of more than 1000 names (http://www.indexfungorum.org/ Names/Names.asp); (Gomes *et al.* 2013, Gao *et al.* 2016). In recent years, however, a polyphasic approach employing multi-locus DNA data together with morphology and ecology has been employed for species delimitation in the genus

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(Udayanga et al. 2011, Gomes et al. 2013). The nuclear ribosomal internal transcribed spacer (ITS), the translation elongation factor 1-α (TEF1), β-tubulin (TUB), histone H3 (HIS), and calmodulin (CAL) genes are the most commonly used molecular loci for the identification of *Diaporthe* spp. (Dissanayake et al. 2015, Udayanga et al. 2015, Huang et al. 2015, Santos et al. 2017). Furthermore, molecular marker aids are being used to rapidly identify Diaporthe species which tend to be morphologically conserved (Udayanga et al. 2012, Tan et al. 2013, Lombard et al. 2014, Thompson et al. 2015, Huang et al. 2015). However, defining species boundaries remains a major challenge in Diaporthe (Huang et al. 2015), which may be a consequence of limited sampling or the use of DNA loci with insufficient phylogenetic resolution (Liu et al. 2016). It has therefore been proposed that new species in the genus should be introduced with caution, and that multiple strains from different origins should be subjected to a multi-gene phylogenetic analysis to determine intraspecific variation (Liu et al. 2016).

The generic relationships of Diaporthe with other genera in Diaporthaceae remain unclear. The family name Diaporthaceae was established by Wehmeyer (1926) to accommodate Diaporthe, Mazzantia, Melanconis, and some other genera, mainly based on morphological characters such as the position, structure, and arrangement of ascomata, stroma, and spore shapes. Castlebury et al. (2002) reported that Diaporthaceae comprised Diaporthe and Mazzantia based on LSU DNA sequence data, removing other genera to different families in Diaporthales. Additional genera subsequently placed in the Diaporthaceae include Leucodiaporthe (Vasilyeva et al. 2007), Stenocarpella (Crous et al. 2006), Phaeocytostroma (Lamprecht et al. 2011), Ophiodiaporthe (Fu et al. 2013), and Pustulomyces (Dai et al. 2014). All the above genera were represented by a few species or are monotypic. Although they appeared to be morphologically divergent from Diaporthe, their phylogenetic relationships remain unclear.

About 991 names of *Diaporthe* and 979 of *Phomopsis* have been established to date (http://www.indexfungorum. org/Names/Names.asp). Among them, many old epithets lack molecular data, and few morphological characters can be used in species delimitation, making it difficult to merge these names to advance to the one name scenario (Rossman *et al.* 2014, 2015). In China, more than 50 plant pathogenic *Phomopsis* species have been published to date (Chi *et al.* 2007). In order to stabilize these species names in the genus *Diaporthe*, here we introduce 12 new combinations for *Phomopsis* species that have been subjected to DNA sequencing, and whose phylogenetic position has been resolved in *Diaporthe* in the present study.

The objectives of this study were: (1) to examine the phylogenetic relationships of *Diaporthe* with other closely related genera in *Diaporthaceae*; (2) to introduce new species in *Diaporthe*; and (3) to transfer *Phomopsis* species described from China to *Diaporthe* based on morphological and newly generated molecular data.

MATERIAL AND METHODS

Isolates

Strains were isolated from leaves of both symptomatic and healthy plant tissues from Yunnan, Zhejiang, and Jiangxi Provinces in China. A few other strains were obtained via the Ningbo Entry-Exit Inspection and Quarantine Bureau, which were isolated from imported plants from other countries. Single spore isolations were conducted from diseased leaves with visible fungal sporulation following the protocol of Zhang et al. (2013), and isolation from surface sterilized leaf tissues was conducted following the protocol of Gao et al. (2014). Fungal endophytes were isolated according to the method described by Liu et al. (2015). The Diaporthe strains were primarily identified from the other fungal species based on cultural characteristics on PDA, spore morphology, and ITS sequence data. Type specimens of new species were deposited in the Mycological Herbarium, Microbiology Institute, Chinese Academy of Sciences, Beijing, China (HMAS), with ex-type living cultures deposited in the China General Microbiological Culture Collection Center (CGMCC).

Morphological analysis

Cultures were incubated on PDA at 25 °C under ambient daylight and growth rates were measured daily for 7 d. To induce sporulation, isolates were inoculated on PNA (pine needle agar; Smith *et al.* 1996) containing double-autoclaved (30 min, 121°C, 1 bar) healthy pine needles and incubated at a room temperature of *ca.* 25 °C (Su *et al.* 2012). Cultures were examined periodically for the development of conidiomata and perithecia. Conidia were taken from pycnidia and mounted in sterilized water. The shape and size of microscopic structures were observed and noted using a light microscope (Nikon Eclipse 80i) with differential interference contrast (DIC). At least 10 conidiomata, 30 conidiophores, alpha and beta conidia were measured to calculate the mean size and standard deviation (SD).

DNA extraction, PCR amplification and sequencing

Isolates were grown on PDA and incubated at 25 °C for 7 d. Genomic DNA was extracted following the protocol of Cubero et al. (1999). The quality and quantity of DNA was estimated visually by staining with GelRed after 1 % agarose gel electrophoresis. The primers ITS5 and ITS4 (White et al. 1990) were used to amplify the internal transcribed spacer region (ITS) of the nuclear ribosomal RNA gene operon, including the 3' end of the 18S nrRNA, the first internal transcribed spacer region, the 5.8S nrRNA gene; the second internal transcribed spacer region and the 5' end of the 28S nrRNA gene. The primers EF1-728F and EF1-986R (Carbone & Kohn 1999) were used to amplify part of the translation elongation factor 1- α gene (*TEF1*), and the primers CYLH3F (Crous et al. 2004) and H3-1b (Glass & Donaldson 1995) were used to amplify part of the histone H3 (HIS) gene. The primers T1 (O'Donnell & Cigelnik 1997) and Bt2b (Glass & Donaldson 1995) were used to amplify the beta-tubulin gene (TUB); the additional combination of Bt2a/Bt2b (Glass & Donaldson 1995) was used in case of amplification failure of the T1/Bt2b primer pair. The primer pair CAL228F/CAL737R

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Table 1. Sources of isolates and GenBank accession numbers used in the phylogenetic analyses of Diaporthaceae.

| Species names* | Culture collection | Isolation sources | Country | GenBan | k Accession | Numbers | References |
|---------------------|--------------------------|-----------------------|--------------------|----------|-------------|----------|-------------------------------|
| | no. | | | ITS | LSU | TEF1 | - |
| D. acaciigena | CBS 129521 (ex- type) | Acacia retinodes | Australia | KC343005 | - | KC343731 | Gomes <i>et al.</i> (2013) |
| D. ampelina | FAU 586 | <i>Vitis</i> sp. | USA: New York | - | AF439635 | - | - |
| D. angelicae | CBS 111592 | Heracleum sphondylium | Austria | KC343027 | - | KC343753 | Gomes <i>et al.</i> (2013) |
| | AR 3724 | Heracleum sphondylium | Austria | KC343026 | - | KC343752 | Gomes <i>et al.</i> (2013) |
| D. apiculata | LC 3418 (ex-type) | Camellia sinensis | China | KP267896 | KY011852 | KP267970 | This study |
| | LC 3452 | Camellia sinensis | China | KP267901 | KY011853 | KP267975 | This study |
| D. arecae complex | LC 4155 | Rhododendron sp. | China | KY011895 | KY011879 | KY011906 | This study |
| | LC 4159 | Rhododendron sp. | China | KY011896 | KY011880 | KY011907 | This study |
| | LC 4164 | Unknown host | China | KY011897 | KY011881 | KY011908 | This study |
| D. biguttusis | LC 1106 (ex-type) | Lithocarpus glaber | China | KF576282 | KY011878 | KF576257 | This study |
| D. compacta | LC 3078 | Camellia sinensis | China | KP267850 | KY011839 | KP267924 | This study |
| | LC 3083 (ex-type) | Camellia sinensis | China | KP267854 | KY011840 | KP267928 | This study |
| | LC 3084 | Camellia sinensis | China | KP267855 | KY011841 | KP267929 | This study |
| D. decedens | CBS 109772 | Corylus avellana | Austria | KC343059 | - | KC343785 | Gomes <i>et al.</i> (2013) |
| D. detrusa | CBS 109770 | Berberis vulgaris | Austria | KC343061 | - | KC343787 | Gomes <i>et al.</i> (2013) |
| D. discoidispora | LC 3503 | Camellia sinensis | China | KY011887 | KY011854 | KY011898 | This study |
| D. elaeagni-glabrae | LC 4802 (ex-type) | Elaeagnus glabra | China | KX986779 | KY011885 | KX999171 | This study |
| | LC 4806 | Elaeagnus glabra | China | KX986780 | KY011886 | KX999172 | This study |
| D. ellipicola | LC 0810 (ex-type) | Lithocarpus glaber | China | KF576270 | KY011873 | KF576245 | This study |
| D. eres | LC 3198 | Camellia sinensis | China | KP267873 | KY011845 | KP267947 | This study |
| | LC 3205 | Camellia sinensis | China | KP714499 | KY011846 | KP714511 | This study |
| | LC 3206 | Camellia sinensis | China | KP714500 | KY011847 | KP714512 | This study |
| | CBS 109767 | Acer campestre | Austria | KC343075 | - | KC343801 | Gomes <i>et al.</i> (2013) |
| D. fusicola | LC 1126 | Lithocarpus glaber | China | KF576281 | KY011836 | KF576256 | This study |
| | LC 0778 (ex-type) | Lithocarpus glaber | China | KF576263 | KY011877 | KF576238 | This study |
| D. hongkongensis | LC 0784 | Lithocarpus glaber | China | KC153104 | KY011876 | KC153095 | This study |
| | LC 0812 | Smilax china | China | KC153103 | KY011875 | KC153094 | This study |
| D. incompleta | LC 6706 | Camellia sinensis | China | KX986793 | KY011859 | KX999185 | This study |
| | LC 1127 (ex-type) | Lithocarpus glaber | China | KF576267 | KY011837 | KF576242 | This study |
| D. mahothocarpi | LC 0732 | Mahonia bealei | China | KC153097 | KY011872 | KC153088 | This study |
| | LC 0763 (ex-type) | Lithocarpus glaber | China | KC153096 | KY011871 | KC153087 | This study |
| D. masirevicii | Diaporthe sp. | Camellia sinensis | China | KY011888 | KY011861 | KY011899 | This study |
| D. neoarctii | CBS 109490 | Ambrosia trifida | USA: New Jersey | KC343145 | - | KC343871 | Gomes <i>et al.</i> (2013) |
| D. oncostoma | CBS 109741 | Robinia pseudoacacia | Russia | KC343161 | - | KC343887 | Gomes <i>et al.</i> (2013) |
| D. oraccinii | LC 3166 (ex-type) | Camellia sinensis | China | KP267863 | KY011843 | KP267937 | This study |
| | LC 3172 | Camellia sinensis | China | KP267864 | KY011844 | KP267938 | This study |
| | LC 3296 | Camellia sinensis | China | KP267884 | KY011849 | KP267958 | This study |
| D. ovoicicola | LC 1128 (ex-type) | Lithocarpus glaber | China | KF576264 | KY011838 | KF576239 | This study |
| D. penetriteum | LC 3215 | Camellia sinensis | China | KP267879 | KY011848 | KP267953 | This study |
| | LC 3353 (ex-type) | Camellia sinensis | China | KP714505 | KY011850 | KP714517 | This study |
| | LC 3394 | Camellia sinensis | China | KP267893 | KY011851 | KP267967 | This study |
| D. perjuncta | CBS 109745 | Ulmus glabra | Austria | KC343172 | - | KC343898 | Gomes <i>et al.</i> (2013) |

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| Table 1. (Continued). | | | | | | | |
|-----------------------------|--------------------|--|--------------|----------|-----------|----------|-----------------------------------|
| Species names* | Culture collection | Isolation sources | Country | GenBank | Accession | Numbers | References |
| | no. | | | ITS | LSU | TEF1 | |
| D. pseudophoenicicola | LC 6150 | Phoenix canariensis | China | KY011891 | KY011865 | KY011902 | This study |
| | LC 6151 | Phoenix canariensis | China | KY011892 | KY011866 | KY011903 | This study |
| D. pustulata | CBS 109742 | Acer pseudoplatanus | Austria | KC343185 | - | KC343911 | Gomes <i>et al.</i> (2013) |
| | CBS 109760 | Acer pseudoplatanus | Austria | KC343186 | - | KC343912 | Gomes <i>et al.</i> (2013) |
| | CBS 109784 | Prunus padus | Austria | KC343187 | - | KC343913 | Gomes <i>et al.</i> (2013) |
| D. rudis | LC 6147 | Dendrobenthamia japonica | USA | KY011890 | KY011864 | KY011901 | This study |
| | LC 6145 | llex aquifolium | China | KY011889 | KY011863 | KY011900 | This study |
| D. saccarata | CBS 116311 | Protea repens, cankers | South Africa | KC343190 | - | KC34391 | Gomes <i>et al.</i> (2013) |
| D. sclerotioides | CBS 296.67 | Cucumis sativus | Netherlands | KC343193 | - | KC343919 | Gomes <i>et al.</i> (2013) |
| D. tectonendophytica | LC 6623 | Unknown host | China | KX986795 | KY011857 | KX999187 | This study |
| D. tectonigena | LC 6512 | Camellia sinensis | China | KX986782 | KY011856 | KX999174 | This study |
| D. ternstroemiae | LC 0777 (ex-type) | Ternstroemia gymnanthera | China | KC153098 | KY011874 | KC153089 | This study |
| D. ueckerae | LC 3564 | Camellia sinensis | China | KP267912 | KY011855 | KP267986 | This study |
| D. undulata | LC 6624 | Unknown host | China | KX986798 | KY011858 | KX999190 | This study |
| D. velutina | LC 4414 | Lithocapus sp. | China | KX986788 | KY011882 | KX999180 | This study |
| | LC 4419 | Neolitsea sp. | China | KX986789 | KY011883 | KX999181 | This study |
| | LC 4421 (ex-type) | Neolitsea sp. | China | KX986790 | KY011884 | KX999182 | This study |
| D. xishuangbanica | LC 6707 | Camellia sinensis | China | KX986783 | KY011860 | KX999175 | This study |
| · | LC 6744 | Camellia sinensis | China | KX986784 | KY011862 | KX999176 | This study |
| D. yunnanensis | LC 6168 | Coffea sp. | China | KX986796 | KY011867 | KX999188 | This study |
| Diaporthe sp. | LC 3156 | Camellia sinensis | China | KP267861 | KY011842 | KP267935 | This study |
| , , | LC 6170 | Coffea sp. | China | KY011893 | KY011869 | KY011904 | This study |
| | LC 6171 | Solanum melongena | China | KY011894 | KY011870 | KY011905 | This study |
| | LC 6232 | Theobroma cacao | China | KX986797 | KY011868 | KX999189 | This study |
| Mazzantia napelli | AR 3498 | Aconitum vulparia | Austria | - | AF408368 | EU222017 | Castlebury et al. (2002) |
| Ophiodiaporthe cyatheae | BCRC 34961 | Cyathea lepifera | Taiwan | JX570889 | JX570891 | KC465406 | Fu <i>et al.</i> (2013) |
| Phaeocytostroma ambiguum | CPC 17071 | Zea mays | South Africa | FR748036 | - | FR748068 | Lamprecht <i>et al.</i> (2011) |
| | CPC 17072 | Zea mays | South Africa | FR748037 | FR748096 | FR748069 | Lamprecht <i>et al.</i> (2011) |
| Ph. plurivorum | CBS 113835 | Helianthus annuus | Portugal | FR748046 | FR748104 | FR748078 | Lamprecht <i>et al.</i> (2011) |
| Ph. sacchari | CBS 275.34 | - | Japan | FR748047 | FR748105 | FR748079 | Lamprecht <i>et al.</i> (2011) |
| Ph. megalosporum | CBS 284.65 | Rice-field soil | India | FR748045 | FR748103 | FR748077 | Lamprecht <i>et al.</i> (2011) |
| Pustulomyces bambusicola | MFLUCC 11-0436 | on dead culm of bamboo | Thailand | - | KF806753 | KF806755 | Dai <i>et al.</i> (2014) |
| Stenocarpella macrospora | CBS 117560 | Rain damaged Bt maize hybrid, 2003-04 season | South Africa | FR748048 | DQ377934 | - | Lamprecht <i>et al.</i> (2011) |
| S. maydis | CBS 117558 | Traditional/landrace maize from 2003/04 season | South Africa | FR748051 | DQ377936 | FR748080 | Lamprecht <i>et al.</i> (2011) |
| Valsa ambiens | CFCC 89894 | Pyrus bretschneideri | China | KR045617 | KR045699 | KU710912 | Fan <i>et al.</i> (2014) |

*New species described in this paper are shown in bold.

(Carbone & Kohn 1999) and LR0R/LR5 primer pair (Rytas & Mark 1990) were used to amplify the calmodulin gene (*CAL*) and the LSU rDNA, respectively. Amplification reactions of 25 μ L were composed of 10 × EasyTaq buffer (MgCl²⁺ included; Transgen, Beijing), 50 μ M dNTPs, 0.2 μ M of each forward and reverse primers (Transgen), 0.5 U EasyTaq DNA polymerase (Transgen) and 1–10 ng of genomic DNA. PCR parameters were as follows: 94 °C for 5 min, followed by 35 cycles of denaturation at 94 °C for 30 s, annealing at a suitable temperature for 30 s (52 °C for ITS and LSU, 56 °C for CAL, HIS, TEF1 and TUB), extension at 72 °C for 30 s and a final elongation step at 72 °C for 10 min. DNA sequencing was performed by Omegagenetics Company, Beijing.

Phylogenetic analyses

The DNA sequences generated with forward and reverse primers were used to obtain consensus sequences using MEGA v. 5.1 (Tamura et al. 2011), and subsequently aligned using MAFFT v. 6 (Katoh & Toh 2010); alignments were manually edited using MEGA v. 5.1 when necessary. Two datasets were employed in the phylogenetic analyses. LSU, ITS and TEF1 loci were selected to infer the generic relationships within Diaporthaceae (Table 1), with Valsa ambiens as outgroup. All available sequences of Diaporthe species were included in the dataset of combined ITS, HIS, TEF1, TUB, and CAL regions to infer the interspecific relationships within Diaporthe (Table 2) with Diaporthella corylina as outgroup. Maximum likelihood (ML) gene trees were estimated using the software RAxML v. 7.4.2 Black Box (Stamatakis 2006, Stamatakis et al. 2008). The RAxML software selected the GTR model of nucleotide substitution with the additional options of modelling rate heterogeneity (Γ) and proportion invariable sites (I). Bayesian analyses (critical value for the topological convergence diagnostic set to 0.01) were performed on the concatenated loci using MrBayes v. 3.2.2 (Ronguist et al. 2012) as described by (Crous et al. 2006) using nucleotide substitution models for each data partition selected by jModeltest (Darriba et al. 2012) and MrModeltest v. 2.3 (Nylander 2004). Bayesian analyses were launched with random starting trees for 10 000 000 generations, and Markov chains were sampled every 1000 generations. The first 25 % resulting trees were discarded as burn-in. The remaining trees were summarized to calculate the posterior probabilities (PP) of each clade being monophyletic. Trees were visualized in FigTree v. 1.1.2 (http://tree.bio.ed.ac.uk/ software/). New sequences generated in this study were deposited in NCBI's GenBank nucleotide database (www. ncbi.nlm.nih.gov; Table 1).

RESULTS

Collection of Diaporthe strains

Twenty-one *Diaporthe* strains including presumed plant pathogens and endophytes were isolated from 11 different host plant species (Table 2) collected from three provinces

(Jiangxi, Yunnan, Zhejiang) in the northern part of China. In addition, 28 strains were isolated from the plant samples inspected by Jiangsu Entry-Exit Inspection and Quarantine Bureau.

The paraphyly of Diaporthe

Phylogenetic analysis was conducted with 224 sequences derived from 76 ingroup taxa from *Diaporthaceae* with *Valsa ambiens* as the outgroup (Table 1). The combined alignment comprised 1 817 characters including gaps (795 for LSU, 558 for ITS, 464 for *TEF1*). Based on the results of the Mrmodeltest, the following priors were set in MrBayes for the different data partitions: GTR+G models with gamma-distributed rates were implemented for LSU and ITS, HKY+I+G model with invgamma-distributed rates were implemented for TEF1. The Bayesian analysis lasted 7 × 10⁸ generations and the consensus tress and posterior probabilities were calculated from the trees left after discarding the first 25 % generations for burn-in (Fig. 1).

The generic relationships of Mazzantia, Ophiodiaporthe, Phaeocytostroma, Pustulomyces, and Stenocarpella with Diaporthe from this analysis are shown in Fig. 1. The topology and branching order of the phylogenetic trees inferred from ML and Bayesian methods were essentially similar. Five genera from Diaporthaceae did not form discrete clades from Diaporthe species but are scattered in the latter, although the family remains monophyletic. The paraphyletic nature of Diaporthe, however, is demonstrated (Fig. 1). Ophiodiaporthe formed a well resolved and distinct clade represented by strain YMJ 1364, and clustered together with the ex-type culture of D. sclerotioides (CBS 296.67) (BPP 0.99, MLBS: 90). Stenocarpella, represented by S. maydis and S. macrospora, was well supported (BPP 1, MLBS = 96) and closely related to several species of Phaeocytostroma. Mazzantia, however, was poorly supported for its phylogenetic position in Diaporthaceae (Fig. 1).

Phylogenetic analyses of the combined datasets of *Diaporthe* species

In total, 1089 sequences derived from 273 ingroup taxa were combined and *Diaporthella corylina* was used as outgroup. A total of 2783 characters including gaps (568 for *CAL*, 554 for *HIS*, 523 for ITS, 636 for *TEF1* and 456 for *TUB*) were included in the multi-locus dataset, comprising sequences generated from this study and others downloaded from GenBank (Table 2). For the Bayesian inference, GTR+I+G model was selected for *CAL*, *HIS* and ITS, HKY+I+G for *TEF1* and *TUB* through the analysis of Mrmodeltest. The maximum likelihood tree conducted by the GTR model confirmed the tree topology and posterior probabilities of the Bayesian consensus tree.

The topology and branching order for the phylogenetic trees inferred from ML and Bayesian methods were essentially similar (Fig. 2). Based on the multi-locus phylogeny and morphology, 49 strains were assigned to 13 species, including eight taxa which we describe here as new (Fig. 2).



Fig. 1. Phylogenetic tree of the family *Diaporthaceae* from a maximum likelihood analysis based on the combined multi-locus dataset (ITS, LSU, *TEF1*). The ML bootstrap values \geq 70 %, bayesian probabilities BPP \geq 0.90 are marked above the branches. The tree is rooted with *Valsa ambiens*.



Fig. 2. Phylogenetic tree of the genus *Diaporthe* from a maximum likelihood analysis based on the combined multi-locus dataset (*CAL*, *HIS*, ITS, *TEF1*, *TUB*). The ML bootstrap values \geq 70 %, bayesian probabilities BPP \geq 0.90 are marked above the branches. The tree is rooted with *Diaporthella corylina*. The novel species are highlighted.

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Fig. 2. (Continued).

| Table 2. Sources of | solates and GenBank accession | numbers used in the phylogenetic an | alyses of Diaporth | ie. Newly sequ | enced materia | I is indicated i | n bold type. | | |
|---------------------|-------------------------------------|---|--------------------|----------------|---------------|------------------|--------------|----------|------------------------------|
| Species names* | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nun | nbers | | References |
| | | | | ПS | TEF1 | TUB | SIH | CAL | |
| D. acaciigena | CBS 129521 (ex-type) | Acacia retinodes | Mimosaceae | KC343005 | KC343731 | KC343973 | KC343489 | KC343247 | Gomes <i>et al.</i> (2013) |
| D. acerina | CBS 137.27 | Acer saccharum | Aceraceae | KC343006 | KC343732 | KC343974 | KC343490 | KC343248 | Gomes <i>et al.</i> (2013) |
| D. acutispora | CGMCC 3.18285 = LC 6161 | <i>Coffea</i> sp., endophyte | Rubiaceae | KX986764 | KX999155 | KX999195 | KX999235 | KX999274 | This study |
| | LC 6142 | Camellia sasanqua, endophyte | Theaceae | KX986762 | KX999153 | KX999193 | KX999233 | KX999272 | This study |
| | LC 6160 | Camellia sasanqua, endophyte | Theaceae | KX986800 | KX999192 | KX999232 | KX999271 | KX999293 | This study |
| D. alleghaniensis | CBS 495.72 (ex-type) | <i>Betula alleghaniensis</i> , branches | Betulaceae | KC343007 | KC343733 | KC343975 | KC343491 | KC343249 | Gomes <i>et al.</i> (2013) |
| D. alnea | CBS 146.46 (ex-type) | Alnus sp. | Betulaceae | KC343008 | KC343734 | KC343976 | KC343492 | KC343250 | Gomes <i>et al.</i> (2013) |
| | CBS 159.47 | Alnus sp. | Betulaceae | KC343009 | KC343735 | KC343977 | KC343493 | KC343251 | Gomes <i>et al.</i> (2013) |
| D. ambigua | CBS 114015 | Pyrus communis | Rosaceae | KC343010 | KC343736 | KC343978 | KC343494 | KC343252 | Gomes <i>et al.</i> (2013) |
| | CBS 117176 | Aspalathus linearis, crown | Fabaceae | KC343011 | KC343737 | KC343979 | KC343495 | KC343253 | Gomes <i>et al.</i> (2013) |
| D. ampelina | CBS 114016 | Vitis vinifera | Vitaceae | AF230751 | AY745056 | JX275452 | ı | AY745026 | Gomes <i>et al.</i> (2013) |
| | CBS 111888 | Vitis vinifera | Vitaceae | KC343016 | KC343742 | KC343984 | KC343500 | KC343258 | Gomes <i>et al.</i> (2013) |
| D. amygdali | CBS 126679 (ex-type) | Prunus dulcis | Rosaceae | KC343022 | KC343748 | KC343990 | KC343506 | KC343264 | Gomes <i>et al.</i> (2013) |
| | CBS 111811 | Vitis vinifera | Vitaceae | KC343019 | KC343745 | KC343987 | KC343503 | KC343261 | Gomes <i>et al.</i> (2013) |
| D. anacardii | CBS 720.97 (ex-epitype) | Anacardium occidentale | Anacardiaceae | KC343024 | KC343750 | KC343992 | KC343508 | KC343266 | Gomes <i>et al.</i> (2013) |
| D. angelicae | CBS 111592 (ex-epitype) | Heracleum sphondylium | Apiaceae | KC343027 | KC343743 | KC343995 | KC343511 | KC343269 | Gomes <i>et al.</i> (2013) |
| | CBS 123215 | Foeniculum vulgare | Apiaceae | KC343028 | KC353754 | KC343996 | KC343512 | KC343270 | Gomes <i>et al.</i> (2013) |
| D. apiculata | LC 4152 | <i>Camellia</i> , leaf | Theaceae | KP267915 | KP267989 | KP293495 | KP293562 | | Gao <i>et al.</i> (2016) |
| | LC 3418, (ex-type) | Camellia sinensis, leaf, endophyte | Theaceae | KP267896 | KP267970 | KP293476 | KP293550 | | Gao e <i>t al.</i> (2016) |
| D. arctii | CBS 136.25 | Arctium sp. | Arecaceae | KC343032 | KC343758 | KC344000 | KC343516 | KC343273 | Gomes <i>et al.</i> (2013) |
| D. arecae | CBS 535.75 | <i>Citrus</i> sp., fruit | Rutaceae | KC343033 | KC343759 | KC344001 | KC343517 | KC343275 | Gomes <i>et al.</i> (2013) |
| | CBS 161.64 (ex-isotype) | <i>Areca catechu</i> , fruit | Arecaceae | KC343032 | KC343758 | KC344000 | KC343516 | KC343274 | Gomes <i>et al.</i> (2013) |
| D. arengae | CBS 114979 (ex-type) | Arenga engleri | Arecaceae | KC343034 | KC343760 | KC344002 | KC343518 | KC343276 | Gomes <i>et al.</i> (2013) |
| D. asheiola | CBS 136967, CPC 16508, (ex-type) | Vaccinium ashei | Ericaceae | KJ160562 | KJ160594 | KJ160518 | ı | KJ160542 | Lombard <i>et al.</i> (2014) |
| | CBS 136968, CPC 16511 | Vaccinium ashei | Ericaceae | KJ160563 | KJ160595 | KJ160519 | | KJ160543 | Lombard <i>et al.</i> (2014) |
| D. aspalathi | CBS 117168 | Aspalathus linearis | Fabaceae | KC343035 | KC343761 | KC344003 | KC343519 | KC343277 | Gomes <i>et al.</i> (2013) |
| | CBS 117169, (ex-type) | Aspalathus linearis | Fabaceae | KC343036 | KC343762 | KC344004 | KC343520 | KC343278 | Gomes <i>et al.</i> (2013) |
| D. australafricana | CBS 111886 | Vitis vinifera | Vitaceae | KC343038 | KC343764 | KC344006 | KC343522 | KC343280 | Gomes <i>et al.</i> (2013) |
| | CBS 113487 | Vitis vinifera | Vitaceae | KC343039 | KC343765 | KC344007 | KC343523 | KC343281 | Gomes <i>et al.</i> (2013) |
| D. baccae | CBS 136971 | Vaccinium corymbosum | Ericaceae | KJ160564 | KJ160596 | | | | Lombard <i>et al.</i> (2014) |
| | CBS 136972 (ex-type) | Vaccinium corymbosum | Ericaceae | KJ160565 | KJ160597 | ı | ı | ı | Lombard <i>et al.</i> (2014) |

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| Table 2. (Continued) | | | | | | | | | |
|----------------------|-------------------------------------|---------------------------------------|----------------|----------|-----------|--------------|----------|----------|--------------------------------|
| Species names* | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nun | lbers | | References |
| | | | | ΠS | TEF1 | TUB | SIH | CAL | |
| D. batatas | CBS 122.21 | Ipomoea batatas | Convolvulaceae | KC343040 | KC343766 | KC344008 | KC343524 | KC343282 | Gomes et al. (2013) |
| D. beckhausii | CBS 138.27 | Viburnum sp. | Caprifoliaceae | KC343041 | KC343767 | KC344009 | KC343525 | KC343283 | Gomes <i>et al.</i> (2013) |
| D. beilharziae | BRIP 54792 (ex-type) | Indigofera australis | Papilionaceae | JX862529 | JX862535 | KF170921 | | | Thompson <i>et al.</i> (2015) |
| D. benedicti | CFCC 50062 (ex-type) | Juglans mandshurica | Juglandaceae | KP208847 | KP208853 | KP208855 | KP208851 | KP208849 | Fan <i>et al</i> . (2015) |
| | CFCC 50063 | Juglans mandshurica | Juglandaceae | KP208848 | KP208854 | KP208856 | KP208852 | KP208850 | Fan <i>et al</i> . (2015) |
| D. betulae | CFCC 50469 (ex-type) | Betula platyphylla | Betulaceae | KT732950 | KT733016 | KT733020 | KT732999 | KT732997 | Du <i>et al.</i> (2016) |
| | CFCC 50470 | Betula platyphylla | Betulaceae | KT732951 | KT733017 | KT733021 | KT733000 | KT732998 | Du <i>et al.</i> (2016) |
| D. betulicola | CFCC 51128 (ex-type) | Betula albosinensis | Betulaceae | KX024653 | KX024655 | KX024657 | KX024661 | KX024659 | Du <i>et al.</i> (2016) |
| | CFCC 51129 | Betula albosinensis | Betulaceae | KX024654 | KX024656 | KX024658 | KX024662 | KX024660 | Du <i>et al.</i> (2016) |
| D. bicincta | DP0659, CBS 121004 | <i>Juglans</i> sp., dead wood | Juglandaceae | KC343134 | KC343860 | KC344102 | KC343618 | | Udayanga <i>et al.</i> (2014a) |
| D. biconispora | ZJUD 60, CGMCC 3.17250 | Citrus sinensis | Rutaceae | KJ490595 | KJ490474 | KJ490416 | KJ490537 | | Huang <i>et al.</i> (2015) |
| | ZJUD 61, CGMCC 3.17251 | Fortunella margarita | Rutaceae | KJ490596 | KJ490475 | KJ490417 | KJ490538 | | Huang <i>et al.</i> (2015) |
| | ZJUD 62, CGMCC 3.17252 | Citrus grandis | Rutaceae | KJ490597 | KJ490476 | KJ490418 | KJ490539 | | Huang <i>et al.</i> (2015) |
| D. biguttulata | ZJUD 47, CGMCC 3.17248 (ex-type) | Citrus limon | Rutaceae | KJ490582 | KJ490461 | KJ490403 | KJ490524 | ı | Huang <i>et al.</i> (2015) |
| | ZJUD 48, CGMCC 3.17249 | Citrus limon | Rutaceae | KJ490583 | KJ490462 | KJ490403 | KJ490525 | | Huang <i>et al.</i> (2015) |
| D. biguttusis | CGMCC 3.17081 (ex-type) | Lithocarpus glabra | Fagaceae | KF576282 | KF576257 | KF576306 | | | Gao <i>et al.</i> (2015) |
| D. brasiliensis | CBS 133183 (ex-type) | Aspidosperma tomentosus | Apocynaceae | KC343042 | KC343768 | KC344010 | KC343526 | KC343284 | Gomes <i>et al.</i> , 2013 |
| | LGMF 926 | Aspidosperma tomentosus | Apocynaceae | KC343043 | KC343769 | KC344011 | KC343527 | KC343285 | Gomes <i>et al.</i> , 2013 |
| D. canthii | CBS 132533 (ex-type) | Canthium inerme | Rubiaceae | JX069864 | KC843120 | KC843230 | | KC843174 | Du <i>et al.</i> (2016) |
| D. carpini | CBS 114437 | Carpinus betulus | Corylaceae | KC343044 | KC343770 | KC344012 | KC343528 | KC343286 | Gomes <i>et al.</i> (2013) |
| D. caulivora | CBS 127268 (ex-neotype) | Glycine max | Fabaceae | KC343045 | KC343771 | KC344013 | KC343529 | KC343287 | Gomes <i>et al.</i> (2013) |
| | CBS 178.55 | Glycine soja | Fabaceae | KC343046 | KC343772 | KC344014 | KC343530 | KC343288 | Gomes <i>et al.</i> (2013) |
| D. celastrina | CBS 139.27 | Celastrus scandens | Celastraceae | KC343047 | KC343773 | KC344015 | KC343531 | , | Gomes <i>et al.</i> (2013) |
| D. cf. heveae 1 | CBS 852.97 | Hevea brasiliensis | Euphorbiaceae | KC343116 | KC343842 | KC344084 | KC343600 | KC343358 | Gomes <i>et al.</i> (2013) |
| D. cf. heveae 2 | CBS 681.84 | Hevea brasilliensis, leaf | Euphorbiaceae | KC343117 | KC343843 | KC344085 | KC343601 | KC343359 | Gomes <i>et al.</i> (2013) |
| D. chamaeropis | CBS 454.81 | Chamaerops humilis, dead part of leaf | Arecaceae | KC343048 | KC343774 | KC344016 | KC343532 | KC343290 | Gomes <i>et al.</i> (2013) |
| | CBS 753.70 | Spartium junceum, dead branch | Fabaceae | KC343049 | KC343775 | KC344017 | KC343533 | KC343291 | Gomes <i>et al.</i> (2013) |
| D. charlesworthii | BRIP 4884m (ex-type) | Rapistrum rugostrum | Brassicaceae | KJ197288 | KJ197250 | KJ197268 | ı | ı | Thompson <i>et al.</i> (2015) |
| D. cinerascens | CBS 719.96 | Ficus carica | Moraceae | KC343050 | KC343776 | KC344018 | KC343534 | KC343292 | Gomes <i>et al.</i> (2013) |
| D. citri | CBS 230.52 | Citrus sinensis | Rutaceae | KC343052 | KC343778 | KC344020 | KC343536 | KC343294 | Gomes <i>et al.</i> (2013) |
| | CBS 199.39 | | ŀ | KC343051 | KC343777 | KC344019 | KC343535 | KC343293 | Gomes <i>et al.</i> (2013) |
| | AR 3405 | Citrus sp. | Rutaceae | KC843311 | KC843071 | KC843187 | KJ420881 | | Udayanga <i>et al.</i> (2014b) |

| Table 2. (Continued). | | | | | | | | | |
|----------------------------|------------------------------------|---|----------------|----------|-----------|--------------|----------|----------|--------------------------------|
| Species names [*] | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nun | nbers | | References |
| | | | | ΠS | TEF1 | TUB | HIS | CAL | |
| D. citriasiana | ZJUD 30 (ex-type) | Citrus unshiu, dead wood | Rutaceae | JQ954645 | JQ954663 | KC357459 | | KC357491 | Huang <i>et al.</i> (2015) |
| | ZJUD 33 | <i>Citrus paradise</i> , stem-end rot fruit | Rutaceae | JQ954658 | JQ972716 | KC357460 | | KC357493 | Huang <i>et al.</i> (2015) |
| D. citrichinensis | ZJUD 34 | Citrus sp. | Rutaceae | JQ954648 | JQ954666 | ı | ı | KC357494 | Huang <i>et al.</i> (2015) |
| | ZJUD 35 | <i>Citrus unshiu</i> , dead wood | Rutaceae | JQ954649 | JQ954667 | KC357461 | | KC357495 | Huang <i>et al.</i> (2015) |
| | ZJUD 36 | Citrus unshiu, dead wood | Rutaceae | KC357556 | KC357525 | KC357462 | | KC357496 | Huang <i>et al.</i> (2015) |
| D. compacta | LC3083 (ex-type) | <i>Camellia sinensis</i> , leaf, endophyte | Theaceae | KP267854 | KP267928 | KP293434 | KP293508 | | Gao <i>et al.</i> (2016) |
| | LC3084 | Camellia sinensis, leaf, endophyte | Theaceae | KP267855 | KP267929 | KP293435 | KP293509 | | Gao <i>et al.</i> (2016) |
| D. convolvuli | CBS 124654 | Convolvulus arvensis | Convolvulaceae | KC343054 | KC343780 | KC344022 | KC343538 | KC343296 | Huang <i>et al.</i> (2015) |
| D. crataegi | CBS 114435 | Crataegus oxyacantha | Rosaceae | KC343055 | KC343781 | KC344023 | KC343539 | KC343297 | Gomes <i>et al.</i> (2013) |
| D. crotalariae | CBS 162.33 (ex-type) | Crotalaria spectabilis | Fabaceae | KC343056 | KC343782 | KC344024 | KC343540 | KC343298 | Gomes <i>et al.</i> (2013) |
| D. cuppatae | CBS 117499 | Aspalathus linearis | Fabaceae | KC343057 | KC343783 | KC344025 | KC343541 | KC343299 | Gomes <i>et al.</i> (2013) |
| D. cynaroidis | CBS 122676 | Protea cynaroides | Proteaceae | KC343058 | KC343784 | KC344026 | KC343542 | KC343300 | Gomes <i>et al.</i> (2013) |
| D. cytosporella | AR 5149 | Citrus sinensis | Rutaceae | KC843309 | KC843118 | KC843222 | ı | KC843143 | Udayanga <i>et al.</i> (2014b) |
| D. decedens | CBS 114281 | Corylus avellana | Corylaceae | KC343060 | KC343786 | KC344028 | KC343544 | KC343302 | Gomes <i>et al.</i> (2013) |
| | CBS 109772 | Corylus avellana | Corylaceae | KC343059 | KC343785 | KC344027 | KC343543 | KC343301 | Gomes <i>et al.</i> (2013) |
| D. detrusa | CBS 109770 | Berberis vulgaris | Berberidaceae | KC343061 | KC343787 | KC344029 | KC343545 | KC343303 | Gomes <i>et al.</i> (2013) |
| | CBS 114652 | Berberis vulgaris | Berberidaceae | KC343062 | KC343788 | KC344030 | KC343546 | KC343304 | Gomes <i>et al.</i> (2013) |
| D. discoidspora | ZJUD 87, CGMCC 3.17254 | Citrus sinensis | Rutaceae | KJ490622 | KJ490501 | KJ490443 | KJ490564 | ı | Huang <i>et al.</i> (2015) |
| | ZJUD 89, CGMCC 3.17255 | Citrus unshiu | Rutaceae | KJ490624 | KJ490503 | KJ490445 | KJ490566 | ı | Huang <i>et al.</i> (2015) |
| D. elaeagni | CBS 504.72 | <i>Elaeagnus</i> sp., twig | Elaeagnaceae | KC343064 | KC343790 | KC344032 | KC343548 | KC343306 | Gomes <i>et al.</i> (2013) |
| D. elaeagni-glabrae | CGMCC 3.18287 = LC 4802 | <i>Elaeagnus glabra</i> , pathogen | Elaeagnaceae | KX986779 | KX999171 | KX999212 | KX999251 | KX999281 | This study |
| | LC 4806 | <i>Elaeagnus glabra</i> , pathogen | Elaeagnaceae | KX986780 | KX999172 | KX999213 | KX999252 | KX999282 | This study |
| D. ellipicola | CGMCC 3.17084 (ex-type) | Lithocarpus glabra, diseased leaves | Fagaceae | KF576270 | KF576245 | KF576291 | I | ı | Gao <i>et al.</i> (2015) |
| D. endophytica | CBS 133811 (ex-type) | Schinus terebinthifolius | Anacardiaceae | KC343065 | KC343791 | KC344033 | KC343549 | KC343307 | Gomes <i>et al.</i> (2013) |
| | LGMF 911 | Schinus terebinthifolius | Anacardiaceae | KC343066 | KC343792 | KC344034 | KC343550 | KC343308 | Gomes <i>et al.</i> (2013) |
| D. eres | AR5193, CBS 13859 (ex- epitype) | Ulmus laevis | Ulmaceae | KJ210529 | KJ210550 | KJ420799 | KJ420850 | I | Udayanga <i>et al.</i> (2014a) |
| | CBS 113470 | Castanea sativa | Fagaceae | KC343146 | KC343872 | KC344114 | KC343630 | ı | Udayanga <i>et al.</i> (2014a) |
| D. eugeniae | CBS 444.82 | Eugenia aromatica, leaf | Mrytaceae | KC343098 | KC343824 | KC344066 | KC343582 | KC343340 | Gomes <i>et al.</i> (2013) |
| D. fibrosa | CBS 109751 | Rhamnus cathartica | Rhamnaceae | KC343099 | KC343825 | KC344067 | KC343583 | KC343341 | Gomes <i>et al.</i> (2013) |
| | CBS 113830 | Rhamnus cathartica | Rhamnaceae | KC343100 | KC343826 | KC344068 | KC343584 | KC343342 | Gomes et al. (2013) |

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| Table 2. (Continued) | | | | | | | | | |
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| Species names [*] | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nun | hers | | References |
| | | | | ITS | TEF1 | TUB | SIH | CAL | |
| D. foeniculina | CBS 116957 | Pyrus pyrifolia | Rosaceae | KC343103 | KC343829 | KC344071 | KC343587 | KC343345 | Gomes et al. (2013) |
| | CBS 187.27 (ex-type of <i>P. theicola</i>) | <i>Camellia sinensis</i> , leaves and branches | Theaceae | KC343107 | KC343833 | KC344075 | KC343591 | KC343349 | Gomes <i>et al.</i> (2013) |
| | CBS 123208 | Foeniculum vulgare | Apiaceae | KC343104 | KC343830 | KC344072 | KC343588 | KC343346 | Gomes <i>et al.</i> (2013) |
| D. fraxini- angustifolia | BRIP 54781 (ex-epitype) | <i>Fraxinus-angustifolia</i> subsp. oxycapa | Oleaceae | JX862528 | JX852534 | KF170920 | ı | ı | Tan <i>et al.</i> (2013) |
| D. ganjae | CBS 180.91 (ex-type) | <i>Cannabis sativa</i> , dead leaf | Cannabaceae | KC343112 | KC343838 | KC344080 | KC343596 | KC343354 | Gomes <i>et al.</i> (2013) |
| D. gardeniae | CBS 288.56 | Gardenia florida, stem | Rubiaceae | KC343113 | KC343839 | KC344081 | KC343597 | KC343355 | Gomes <i>et al.</i> (2013) |
| D. goulteri | BRIP 55657a (ex-type) | Helianthus annuus | Asteraceae | KJ197289 | KJ197252 | KJ197270 | | | Thompson <i>et al.</i> (2015) |
| D. gulyae | BRIP 54025 (ex-type) | Helianthus annuus | Asteraceae | JF431299 | JN645803 | KJ197271 | , | | Thompson <i>et al.</i> (2015) |
| D. helianthi | CBS 344.94 | Helianthus annuus | Asteraceae | KC343114 | KC343840 | KC344082 | KC343598 | KC343356 | Gomes <i>et al.</i> (2013) |
| | CBS 592.81 (ex-type) | Helianthus annuus | Asteraceae | KC343115 | KC343841 | KC344083 | KC343599 | KC343357 | Gomes <i>et al.</i> (2013) |
| D. helicis | AR 5211 | Hedera helix | Araliaceae | KJ210538 | KJ210559 | KJ420828 | KJ420875 | KJ435043 | Udayanga <i>et al.</i> (2014a) |
| D. hickoriae | CBS 145.26 (ex-epitype) | Carya glabra | Juglandaceae | KC343118 | KC343844 | KC344086 | KC343602 | KC343360 | Gomes <i>et al.</i> (2013) |
| D. hongkongensis | CBS 115448 (ex-type) | Dichroa febrifuga, fruit | Hydrangeaceae | KC343119 | KC343845 | KC344087 | KC343603 | KC343361 | Gomes <i>et al.</i> (2013) |
| D. hordei | CBS 481.92 | Hordeum vulgare | Poaceae | KC343120 | KC343846 | KC344088 | KC343604 | KC343362 | Gomes <i>et al.</i> (2013) |
| D. impulsa | CBS 114434 | Sorbus aucuparia | Rosaceae | KC343121 | KC343847 | KC344089 | KC343605 | KC343363 | Gomes <i>et al.</i> (2013) |
| | CBS 141.27 | Sorbus americana | Rosaceae | KC343122 | KC343848 | KC344090 | KC343606 | KC343364 | Gomes <i>et al.</i> (2013) |
| D. incompleta | CGMCC 3.18288 = LC 6754 | <i>Camellia sinensis</i> , pathogen | Theaceae | KX986794 | KX999186 | KX999226 | KX999265 | KX999289 | This study |
| | LC 6706 | <i>Camellia sinensis</i> , pathogen | Theaceae | KX986793 | KX999185 | | KX999264 | KX999288 | This study |
| D. inconspicua | CBS 133813 (ex-type) | <i>Maytenus ilicifolia</i> , endophytic in petiole | Celastraceae | KC343123 | KC343849 | KC344091 | KC343607 | KC343365 | Gomes <i>et al.</i> (2013) |
| D. infecunda | CBS 133812 (ex-type) | Schinus terebinthifolius | Anacardiaceae | KC343126 | KC343852 | KC344094 | KC343610 | KC343368 | Gomes <i>et al.</i> (2013) |
| | LGMF 908 | Schinus terebinthifolius | Anacardiaceae | KC343127 | KC343853 | KC344095 | KC343611 | KC343369 | Gomes <i>et al.</i> (2013) |
| D. kongii | BRIP 54031 (ex-type) | Helianthus annuus | Asteraceae | JF431301 | JN645797 | KJ197272 | ı | ı | Thompson <i>et al.</i> (2011) |
| D. lichicola | BRIP 54900 (ex-type) | Litchi chinensis | Sapindaceae | JX862533 | JX862539 | KF170925 | 1 | , | Tan <i>et al.</i> (2013) |
| D. longicicola | CGMCC 3.17089 (ex-type) | Lithocarpus glabra | Fagaceae | KF576267 | KF576242 | KF576291 | 1 | , | Gao <i>et al.</i> (2015) |
| D. longicolla | FAU 599 | Glycine max | Fabaceae | KJ590728 | KJ590767 | KJ610883 | KJ659188 | | Udayanga <i>et al.</i> (2015) |
| D. longispora | CBS 194.36 (ex-type) | <i>Ribes</i> sp. | Grossulariaceae | KC343135 | KC343861 | KC344103 | KC343619 | KC343377 | Gomes <i>et al.</i> (2013) |
| D. lusitanicae | CBS 123212 (ex-type) | Foeniculum vulgare | Apiaceae | KC343136 | KC343862 | KC344104 | KC343620 | | Gomes <i>et al.</i> (2013) |
| | CBS 123213 | Foeniculum vulgare | Apiaceae | KC343137 | KC343863 | KC344105 | KC343621 | KC343379 | Gomes <i>et al.</i> (2013) |
| D. macintoshii | BRIP 55064a | Rapistrum rugostrum | Brassicaceae | KJ197290 | KJ197251 | KJ197269 | | | Thompson <i>et al.</i> (2015) |
| D. mahothocarpus | CGMCC 3.15181 | Lithocarpus glabra | Fagaceae | KC153096 | KC153087 | | | | Gao <i>et al.</i> (2014) |
| D. manihotia | CBS 505.76 | Manihot utilissima, leaves | Euphorbiaceae | KC343138 | KC343864 | KC344106 | KC343622 | KC343380 | Gomes <i>et al.</i> (2013) |

| Table 2. (Continued) | | | | | | | | | |
|----------------------------|------------------------------------|--|----------------|----------|-----------|--------------|----------|----------|--------------------------------|
| Species names [*] | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nur | nbers | | References |
| | | | | ΠS | TEF1 | TUB | HIS | CAL | |
| D. maritima | NB 382-2E | Picea rubens needle | Pinaceae | KU552026 | KU552024 | KU574614 | | | Tanney <i>et al.</i> (2016) |
| | NB 463-3A | Picea rubens needle | Pinaceae | KU552027 | KU552022 | KU574616 | | , | Tanney <i>et al.</i> (2016) |
| | NB 365-711 (ex-type) | Picea rubens needle | Pinaceae | KU552025 | KU552023 | KU574615 | | , | Tanney <i>et al.</i> (2016) |
| D. masirevicii | BRIP 57330 | <i>Chrysanthemoides monilifera</i> subsp. <i>rotundat</i> a | Rosaceae | KJ197275 | KJ197237 | KJ197255 | ı | ı | Huang <i>et al.</i> (2015) |
| | BRIP 57892a (ex-type) | Helianthus annuus | Asteraceae | KJ197277 | KJ197239 | KJ197257 | ı | | Huang <i>et al.</i> (2015) |
| D. mayteni | CBS 133185 (ex-type) | Maytenus ilicicolia | Celastraceae | KC343139 | KC343865 | KC344107 | KC343623 | KC343381 | Gomes <i>et al.</i> (2013) |
| D. megalospora | CBS 143.27 | Sambucus canadensis | Caprifoliaceae | KC343140 | KC343866 | KC344108 | KC343624 | KC343382 | Gomes <i>et al.</i> (2013) |
| D. melonis | CBS 435.87 | Glycine soja | Fabaceae | KC343141 | KC343867 | KC344109 | KC343625 | KC343383 | Gomes <i>et al.</i> (2013) |
| | CBS 507.78 (ex-isotype) | Cucumis melo | Cucurbitaceae | KC343142 | KC343868 | KC344110 | KC343626 | KC343384 | Gomes <i>et al.</i> (2013) |
| D. middletonii | BRIP 57329 | <i>Chrysanthemoides monilifera</i> subsp. <i>rotundata</i> | Rosaceae | KJ197285 | KJ197247 | KJ197265 | ı | · | Thompson <i>et al.</i> (2015) |
| | BRIP 54884e (ex-type) | Rapistrum rugostrum | Brassicaceae | KJ197286 | KJ197248 | KJ197266 | | | Thompson <i>et al.</i> (2015) |
| D. miriciae | BRIP 55662c | Glycine max | Fabaceae | KJ197283 | KJ197245 | KJ197263 | | | Thompson <i>et al.</i> (2015) |
| | BRIP 54736j (ex-type) | Helianthus annuus | Asteraceae | KJ197282 | KJ197244 | KJ197262 | | | Thompson <i>et al.</i> (2015) |
| | BRIP 56918a | Vigna radiata | Papilionaceae | KJ197284 | KJ197246 | KJ197264 | | | Thompson <i>et al.</i> (2015) |
| D. multigutullata | ZJUD 98 | Citrus grandis | Rosaceae | KJ490633 | KJ490512 | KJ490454 | KJ490575 | | Huang <i>et al.</i> (2015) |
| D. musigena | CBS 129519; CPC 17026 (ex-type) | <i>Musa</i> sp., leaves | Musaceae | KC343143 | KC343869 | KC344111 | KC343627 | KC343385 | Gomes <i>et al.</i> (2013) |
| D. neilliae | CBS 144.27 | <i>Spiraea</i> sp. | Rosaceae | KC343144 | KC343870 | KC344112 | KC343628 | KC343386 | Udayanga <i>et al.</i> (2014a) |
| D. neoarctii | CBS 109490 (ex-type) | Ambrosia trifida | Asteraceae | KC343145 | KC343871 | KC344113 | KC343629 | KC343387 | Gomes <i>et al.</i> (2013) |
| D. neoraonikayapo- rum | MFLUCC 14-1136 | Tectona grandis | Verbenaceae | KU712449 | KU749369 | KU743988 | | KU749356 | Doilom <i>et al.</i> (2017) |
| | MFLUCC 14-1137 | Tectona grandis | Verbenaceae | KU712450 | KU749370 | KU743989 | ı | KU749357 | Doilom <i>et al.</i> (2017) |
| | MFLUCC 14-1133 | Tectona grandis | Verbenaceae | KU712448 | KU749368 | KU743987 | | KU749355 | Doilom <i>et al.</i> (2017) |
| D. nobilis | CBS 200.39 | Laurus nobilis, stem | Lauraceae | KC343151 | KC343877 | KC344119 | KC343635 | KC343393 | Gomes <i>et al.</i> (2013) |
| D. nomurai | CBS 157.29 | <i>Morus</i> sp. | Moraceae | KC343154 | KC343880 | KC344122 | KC343638 | KC343396 | Gomes <i>et al.</i> (2013) |
| D. nothofagi | BRIP 54801 (ex-type) | Nothofagus cunninghamii | Fagaceae | JX862530 | JX862536 | KF170922 | ı | ı | Tan <i>et al.</i> (2013) |
| D. novem | CBS 127269 | Glycine max | Fabaceae | KC343155 | KC343881 | KC344123 | KC343639 | KC343397 | Gomes <i>et al.</i> (2013) |
| | CBS 127270 (ex-type) | Glycine max | Fabaceae | KC343156 | KC343882 | KC344124 | KC343640 | KC343398 | Gomes <i>et al.</i> (2013) |
| D. oncostoma | CBS 100454 | <i>Robinia pseudoacacia</i> , leaf spot | Fabaceae | KC343160 | KC343886 | KC344128 | KC343644 | KC343402 | Gomes <i>et al.</i> (2013) |
| | CBS 109741 | Robinia pseudoacacia | Fabaceae | KC343161 | KC343887 | KC344129 | KC343645 | KC343403 | Gomes <i>et al.</i> (2013) |
| D. oraccinii | LC 3166 (ex-type) | Camellia sinensis, leaf, endophyte | Theaceae | KP267863 | KP267937 | KP293443 | KP293517 | ı | Gao <i>et al.</i> (2016) |
| | LC 3296 | <i>Camellia sinensis</i> , leaf, endophyte | Theaceae | KP267884 | KP267958 | KP293464 | KP293538 | | Gao <i>et al.</i> (2016) |

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| Table 2. (Continued). | | | | | | | | | |
|---------------------------------------|-------------------------|--|---------------|----------|-----------|--------------|----------|----------|-------------------------------|
| Species names [*] | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nun | nbers | | References |
| | | | | ΠS | TEF1 | TUB | SIH | CAL | |
| D. ovalispora | ZJUD 93, CGMCC 3.17256 | Citrus limon | Rosaceae | KJ490628 | KJ490507 | KJ490449 | KJ490570 | 1 | Huang <i>et al.</i> (2015) |
| D. oxe | CBS 133186 (ex-type) | Maytenus ilicifolia | Celastraceae | KC343164 | KC343890 | KC344132 | KC343648 | KC343406 | Gomes <i>et al.</i> (2013) |
| | CBS 133187 | Maytenus ilicifolia | Celastraceae | KC343165 | KC343891 | KC344133 | KC343649 | KC343407 | Gomes <i>et al.</i> (2013) |
| D. padi var. padi | CBS 114200 | Prunus padus | Rosaceae | KC343169 | KC343895 | KC344137 | KC343653 | KC343411 | Gomes <i>et al.</i> (2013) |
| | CBS 114649 | Alnus glutinosa | Betulaceae | KC343170 | KC343896 | KC344138 | KC343654 | KC343412 | Gomes <i>et al.</i> (2013) |
| D. paranensis | CBS 133184 (ex-type) | Maytenus ilicifolia | Celastraceae | KC343171 | KC343897 | Kc344139 | KC343655 | KC343413 | Gomes <i>et al.</i> (2013) |
| D. pascoei | BRIP 54847 (ex-type) | Persea americana | Lauraceae | JX862532 | JX862538 | KF170924 | | | Tan <i>et al.</i> (2013) |
| D. penetriteum | LC 3353 | Camellia sinensis, leaf | Theaceae | KP714505 | KP714517 | KP714529 | KP714493 | | Gao <i>et al.</i> (2016) |
| | LC 3394 | Camellia sinensis, leaf | Theaceae | KP267893 | KP267967 | KP293473 | KP293547 | | Gao <i>et al.</i> (2016) |
| D. perjuncta | CBS 109745 (ex-type) | Ulmus glabra | Ulmaceae | KC343172 | KC343898 | KC344140 | KC343656 | KC343414 | Gomes <i>et al.</i> (2013) |
| D. perniciosa | CBS 124030 | Malus pumila, bark | Rosaceae | KC343149 | KC343875 | KC344117 | KC343633 | KC343391 | Gomes <i>et al.</i> (2013) |
| D. perseae | CBS 151.73 | Perseae gratissima, young fruit | Lauraceae | KC343173 | KC343899 | KC344141 | KC343657 | KC343415 | Gomes <i>et al.</i> (2013) |
| D. phaseolorum | AR 4203, CBS 139281 | Phaseolus vulgaris | Fabaceae | KJ590738 | KJ590739 | KJ610893 | KJ659220 | , | Huang <i>et al.</i> (2015) |
| | CBS 116019 | Caperonia palustris | Euphorbiaceae | KC343175 | KC343901 | KC344143 | KC343659 | KC343417 | Gomes <i>et al.</i> (2013) |
| | CBS 116020 | Aster exilis | Asteraceae | KC343176 | KC343902 | KC344144 | KC343660 | KC343418 | Gomes <i>et al.</i> (2013) |
| D. podocarpi- macrophylli | CGMCC 3.18281 = LC 6155 | Podocarpus macrophyllus, endophyte | Podocarpaceae | KX986774 | KX999167 | KX999207 | KX999246 | KX999278 | This study |
| | LC 6144 | Podocarpus macrophyllus, endophyte | Podocarpaceae | KX986773 | KX999166 | KX999206 | KX999245 | ı | This study |
| | LC 6194 | Podocarpus macrophyllus, endophyte | Podocarpaceae | KX986765 | KX999156 | KX999196 | KX999236 | KX999275 | This study |
| | LC 6197 | Podocarpus macrophyllus, endophyte | Podocarpaceae | KX986777 | KX999170 | KX999210 | KX999249 | KX999279 | This study |
| | LC 6200 | Podocarpus macrophyllus, endophyte | Podocarpaceae | KX986769 | KX999161 | KX999201 | KX999240 | KX999276 | This study |
| | LC 6229 | Olea europaea, endophytes | Oleaceae | KX986771 | KX999164 | KX999204 | KX999243 | KX999277 | This study |
| D. pseudomangi í - erae | CBS 101339 (ex-type) | Mangifera indica | Anacardiaceae | KC343181 | KC343907 | KC344149 | KC343665 | KC343423 | Gomes <i>et al.</i> (2013) |
| | CBS 388.89 | Mangifera indica, peel of fruit | Anacardiaceae | KC343182 | KC343908 | KC344150 | KC343666 | KC343424 | Gomes <i>et al.</i> (2013) |
| D. pseudophoenici- cola | CBS 462.69 (ex-type) | Phoenix dactylifera, dead tops of green leaves | Anacardiaceae | KC343184 | KC343910 | KC344152 | KC343668 | KC343426 | Gomes <i>et al.</i> (2013) |
| | CBS 176.77 | <i>Mangifera indica</i> , showing dieback | Anacardiaceae | KC343183 | KC343909 | KC344151 | KC343667 | KC343425 | Gomes <i>et al.</i> (2013) |
| D. pterocarpi | MFLUCC 10-0571 | Pterocarous indicus | Papilionaceae | JQ619899 | JX275416 | JX275460 | | JX197451 | Udayanga <i>et al.</i> (2012) |
| | MFLUCC 10-0575 | Pterocarous indicus | Papilionaceae | JQ619901 | JX275418 | JX275462 | | JX197453 | Udayanga <i>et al.</i> (2012) |

| Table 2. (Continued). | | | | | | | | | |
|----------------------------|---------------------------|--|---------------|----------|-----------|--------------|----------|----------|-------------------------------------|
| Species names [*] | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nur | nbers | | References |
| | | | | ПS | TEF1 | TUB | HIS | CAL | |
| D. pterocarpicola | MFLUCC 10-0580a (ex-type) | Piterocarpus indicus | Papilionaceae | JQ619887 | JX275403 | JX275441 | | JX197433 | Udayanga <i>et al.</i> (2012) |
| | MFLUCC 10-0580b | Piterocarpus indicus | Papilionaceae | JQ619888 | JX275404 | JX275442 | ı | JX197434 | Udayanga <i>et al.</i> (2012) |
| D. pulla | CBS 338.89 | Hedera helix | Araliaceae | KC343152 | KC343878 | KC344120 | KC343636 | ı | Udayanga <i>et al.</i> (2014a) |
| D. pustulata | CBS 109742 | Acer pseudoplatanus | Aceraceae | KC343185 | KC343911 | KC344153 | KC343669 | KC343427 | Gomes <i>et al.</i> (2013) |
| | CBS 109760 | Acer pseudoplatanus | Aceraceae | KC343186 | KC343912 | KC344154 | KC343670 | KC343428 | Gomes <i>et al.</i> (2013) |
| D. raonikayaporum | CBS 133182 (ex-type) | Spondias mombin | Anacardiaceae | KC343188 | KC343914 | KC344156 | KC343672 | KC343430 | Gomes <i>et al.</i> (2013) |
| D. rhoina | CBS 146.27 | Rhus toxicodendron | Anacardiaceae | KC343189 | KC343915 | KC344157 | KC343673 | KC343431 | Gomes <i>et al.</i> (2013) |
| D. rudis | CBS 113201 (ex-type) | Vitis vinifera | Vitaceae | KC343234 | KC343960 | KC344202 | KC343718 | KC343476 | Machingambi <i>et al.</i> (2015) |
| | CBS 114011 | Vitis Vinifera | Vitaceae | KC343235 | KC343961 | KC344203 | KC343718 | KC343477 | Machingambi <i>et al.</i> (2015) |
| D. saccarata | CBS 116311 (ex-type) | Protea repens, cankers | Proteceae | KC343190 | KC343916 | KC344158 | KC343674 | KC343432 | Gomes <i>et al.</i> (2013) |
| D. sackstonii | BRIP 54669b (ex-type) | Helianthus annuus | Asteraceae | KJ197287 | KJ197249 | KJ197267 | | | Gomes <i>et al.</i> (2013) |
| D. salicicola | BRIP 54825 (ex-type) | Salix purpurea | Salicaceae | JX862531 | JX862537 | KF170923 | | | Gomes <i>et al.</i> (2013) |
| D. schini | LGMF 910, CPC 20286 | Schinus terebinthifolius, endophytic in leaf | Anacardiaceae | KC343192 | KC343918 | KC344160 | KC343676 | KC343434 | Thompson <i>et al.</i> (2015) |
| | CBS 133181 (ex-type) | Schinus terebinthifolius, endophytic in leaf | Anacardiaceae | KC343191 | KC343917 | KC344159 | KC343675 | KC343433 | Tan <i>et al.</i> (2013) |
| D. sclerotioides | CBS 296.67 (ex-type) | Cucumis sativus | Cucurbitaceae | KC343193 | KC343919 | KC344161 | KC343677 | KC343435 | Gomes <i>et al.</i> (2013) |
| | CBS 710.76 | Cucumis sativus | Cucurbitaceae | KC343194 | KC343920 | KC344162 | KC343678 | KC343436 | Gomes <i>et al.</i> (2013) |
| D. scobina | CBS 251.38 | <i>Fraxinus Excelsior</i> , living and dead twig | Oleaceae | KC343195 | KC343921 | KC344163 | KC343679 | KC343437 | Gomes <i>et al.</i> (2013) |
| D. serafiniae | BRIP 55665a (ex-type) | Helianthus annuus | Asteraceae | KJ197274 | KJ197236 | KJ197254 | ı | ı | Gomes <i>et al.</i> (2013) |
| | BRIP 54136 | Lupinus albus "Rosetta" | Fabaceae | KJ197273 | KJ197235 | KJ197253 | | | Gomes <i>et al.</i> (2013) |
| D. siamensis | MFLUCC 10_0573a | Dasymaschalon sp. | Annonaceae | JQ619879 | JX275393 | JX275429 | ı | ı | Thompson <i>et al.</i> (2015) |
| | MFLUCC 10_0573b | Dasymaschalon sp. | Annonaceae | JQ619880 | JX275395 | JX275430 | ı | ı | Thompson <i>et al.</i> (2015) |
| D. sojae | CBS 100.87 | Glycine soja | Fabaceae | KC343196 | KC343922 | KC344164 | KC343680 | KC343438 | Udayanga <i>et al.</i> (2012) |
| | CBS 116017 | Euphorbia nutans | Euphorbiaceae | KC343197 | KC343923 | KC344165 | KC343681 | KC343439 | Udayanga <i>et al.</i> (2012) |
| | FAU 635 | Glycine max | Fabaceae | KJ590719 | KJ590762 | KJ610875 | KJ659208 | ı | Gomes <i>et al.</i> (2013) |
| D. sterilis | CBS 136969 (ex-type) | Vaccinium corymbosum | Ericaceae | KJ160579 | KJ160611 | KJ160528 | I | KJ160548 | Gomes <i>et al.</i> (2013) |
| | CBS 136970 | Vaccinium corymbosum | Ericaceae | KJ160580 | KJ160612 | KJ160529 | | KJ160549 | Huang <i>et al.</i> (2015) |
| D. stewartii | CBS 193.36 | | · | FJ889448 | GQ250324 | ı | ı | ı | Lombard <i>et al.</i> (2014) |
| D. stictica | CBS 370.54 | Buxus sampervirens, dead twig | Buxaceae | KC343212 | KC343938 | KC344180 | KC343696 | KC343454 | Lombard <i>et al.</i> (2014) |
| D. subclavata | ZJUD 83, CGMCC 3.17253 | Citrus grandis cv. Shatianyou | Rosaceae | KJ490618 | KJ490497 | KJ490439 | KJ490560 | 1 | Udayanga <i>et al.</i> (2011) |

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| Table 2. (Continued) | | | | | | | | | |
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| Species names* | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nun | hers | | References |
| | | | | ΠS | TEF1 | TUB | HIS | CAL | |
| | ZJUD 95, CGMCC 3.17257 | Citrus unshiu | Rosaceae | KJ490630 | KJ490509 | KJ490451 | KJ490572 | 1 | Gomes et al. (2013) |
| D. subordinaria | CBS 101711 | Plantago lanceolata | Plantaginaceae | KC343213 | KC343939 | KC344181 | KC343697 | KC343455 | Huang <i>et al.</i> (2015) |
| | CBS 464.90 | Plantago lanceolata | Plantaginaceae | Kc343214 | KC343940 | KC344182 | KC343698 | KC343456 | Huang <i>et al.</i> (2015) |
| D. tecomae | CBS 100547 | <i>Tabebuia</i> sp. | Bignoniaceae | KC343215 | KC343941 | KC344183 | KC343699 | KC343457 | Gomes <i>et al.</i> (2013) |
| D. tectonae | MFLUCC 12-0777 | Tectona grandis | Verbenaceae | KU712430 | KU749359 | KU743977 | | KU749345 | Gomes <i>et al.</i> (2013) |
| | MFLUCC 14-1138 | Tectona grandis | Verbenaceae | KU712437 | KU749365 | KU743984 | | KU749352 | Gomes <i>et al.</i> (2013) |
| D. tectonendo- phytica | MFLUCC 13-0471 | Tectona grandis | Verbenaceae | KU712439 | KU749367 | KU743986 | | KU749354 | Doilom <i>et al.</i> (2017) |
| D. tectonigena | MFLUCC 12-0767 | Tectona grandis | Verbenaceae | KU712429 | KU749371 | KU743976 | | KU749358 | Doilom <i>et al.</i> (2017) |
| D. terebinthifolii | CBS 133180 | Schinus terebinthifolius | Anacardiaceae | KC343216 | KC343942 | KC344184 | KC343700 | KC343458 | Doilom <i>et al.</i> (2017) |
| | LGMF 907 | Schinus terebinthifolius | Anacardiaceae | KC343217 | KC343943 | KC344185 | KC343701 | KC343459 | Doilom <i>et al.</i> (2017) |
| D. thunbergii | MFLUCC 10_0756a | Thunbergia laurifolia | Acanthaceae | JQ619893 | JX275409 | JX275449 | | JX197440 | Doilom <i>et al.</i> (2017) |
| | MFLUCC 10_0756b | Thunbergia laurifolia | Acanthaceae | JQ619894 | JX275410 | JX275450 | | JX197441 | Doilom <i>et al.</i> (2017) |
| D. toxica | CBS 534.93 (ex-type) | Lupinus angustifolius, stem | Fabaceae | KC343220 | KC343946 | KC344188 | KC343704 | KC343462 | Udayanga <i>et al.</i> (2012) |
| | CBS 535.93 | Lupinus sp. | Fabaceae | KC343221 | KC343947 | KC344189 | KC343705 | KC343463 | Udayanga <i>et al.</i> (2012) |
| D. tulliensis | BRIP 62248a | Theobroma cacao | Sterculiaceae | KR936130 | KR936133 | KR936132 | | | Gomes <i>et al.</i> (2013) |
| D. ueckerae | FAU 656 | Cucumis melo | Cucurbitaceae | KJ590726 | KJ590747 | KJ610881 | KJ659215 | | Gomes <i>et al.</i> (2013) |
| | FAU 658 | Cucumis melo | Cucurbitaceae | KJ590725 | KJ590746 | KJ610880 | KJ659214 | , | Crous <i>et al.</i> (2015) |
| D. undulata | CGMCC 3.18293 = LC 6624 | Unknown host, pathogen | I | KX986798 | KX999190 | KX999230 | KX999269 | | Huang <i>et al.</i> (2015) |
| | LC 8110 | Unknown host, pathogen | ı | KY491545 | KY491555 | KY491565 | | | Huang <i>et al.</i> (2015) |
| | LC 8111 | Unknown host, pathogen | ı | KY491546 | KY491556 | KY491566 | | | This study |
| D. unshiuensis | ZJUD 51, CGMCC 3.17568 | Fortunella margarita | Rutaceae | KJ490586 | KJ490465 | KJ490407 | KJ490528 | | This study |
| | ZJUD 52, CGMCC 3.17569 | Citrus unshiu | Rosaceae | KJ490587 | KJ490466 | KJ490408 | KJ490529 | | This study |
| D. vaccinii | CBS 160.32 (ex-type) | Oxycoccus macrocarpos | Ericaceae | KC343228 | KC343954 | KC344196 | KC343712 | KC343470 | Huang <i>et al.</i> (2015) |
| | CBS 118571 | Vaccinium corymbosum | Ericaceae | KC343223 | KC343949 | KC344191 | KC343707 | KC343465 | Huang <i>et al.</i> (2015) |
| D. vawdreyi | BRIP 57887a | Psidium guajava | Sterculiaceae | KR936126 | KR936129 | KR936128 | | | Gomes <i>et al.</i> (2013) |
| D. velutina | CGMCC 3.18286 = LC 4421 | Neolitsea sp., pathogen | Lauraceae | KX986790 | KX999182 | KX999223 | KX999261 | | Gomes <i>et al.</i> (2013) |
| | LC 4419 | Neolitsea sp., pathogen | Lauraceae | KX986789 | KX999181 | KX999222 | KX999260 | KX999286 | Crous <i>et al.</i> (2015) |
| | LC 4641 | <i>Callerya cinerea</i> , pathogen | Fabaceae | KX986792 | KX999184 | KX999225 | KX999263 | KX999287 | This study |
| | LC 4788 | Unknown host, pathogen | | KX986785 | KX999177 | KX999218 | KX999256 | KX999285 | This study |
| | LC 6708 | Camellia sinensis, pathogen | Theaceae | KX986787 | KX999179 | KX999220 | KX999258 | | This study |
| D. vexans | CBS 127.14 | Solanum melongena | Solanaceae | KC343229 | KC343955 | KC344197 | KC343713 | KC343471 | This study |
| D. virgilia | CMW 40755 (ex-type) | Virgilia oroboides | Unknown | KP247573 | | KP247582 | | ı | This study |

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| Table 2. (Continued). | | | | | | | | | |
|----------------------------|-------------------------|------------------------------------|---------------|----------|-----------|--------------|----------|----------|-------------------------------|
| Species names* | Culture collection no. | Isolation sources | Host family | | GenBank A | ccession Nur | nbers | | References |
| | | | | ΠS | TEF1 | TUB | HIS | CAL | |
| | CMW 40748 | Virgilia oroboides | Unknown | KP247566 | , | KP247575 | | | Gomes <i>et al.</i> (2013) |
| D. woodii | CBS 558.93 | Lupinus sp. | Fabaceae | KC343244 | KC343970 | KC344212 | KC343728 | KC343486 | Gomes <i>et al.</i> (2013) |
| D. woolworthii | CBS 148.27 | Ulmus americana | Ulmaceae | KC343245 | KC343971 | KC344213 | KC343729 | KC343487 | Gomes <i>et al.</i> (2013) |
| D. xishuangbanica | CGMCC 3.18282= LC 6707 | Camellia sinensis, pathogen | Theaceae | KX986783 | KX999175 | KX999216 | KX999255 | | This study |
| | LC 6744 | Camellia sinensis, pathogen | Theaceae | KX986784 | KX999176 | KX999217 | | | This study |
| D. yunnanensis | CGMCC 3.18289 = LC6168 | Coffea sp., endophytes | Rubiaceae | KX986796 | KX999188 | KX999228 | KX999267 | KX999290 | This study |
| | LC 8106 | Coffea sp., endophytes | Rubiaceae | KY491541 | KY491551 | KY491561 | | KY491571 | This study |
| | LC 8107 | Coffea sp., endophytes | Rubiaceae | KY491542 | KY491552 | KY491562 | | KY491572 | This study |
| Diaporthe sp. | LC 6496 | Camellia sinensis, endophytes | Theaceae | KX986781 | KX999173 | KX999214 | KX999253 | KX999283 | This study |
| | LC 6512 | Camellia sinensis, endophyte | Theaceae | KX986782 | KX999174 | KX999215 | KX999254 | KX999284 | This study |
| | LC 6232 | <i>Theobroma cacao</i> , endophyte | Sterculiaceae | KX986797 | KX999189 | KX999229 | KX999268 | KX999291 | This study |
| | LC 8108 | <i>Theobroma cacao</i> , endophyte | Sterculiaceae | KY491543 | KY491553 | KY491563 | | KY491573 | This study |
| | LC 8109 | <i>Theobroma cacao</i> , endophyte | Sterculiaceae | KY491544 | KY491554 | KY491564 | ı | KY491574 | This study |
| | LC 6623 | Unknown host, pathogen | | KX986795 | KX999187 | KX999227 | KX999266 | | This study |
| | LC 8114 | Unknown host, pathogen | | КҮ491549 | KY491559 | KY491569 | ı | | This study |
| | LC 8115 | Unknown host, pathogen | | KY491550 | KY491560 | KY491570 | ı | | This study |
| | LGMF 947 | Glycine max, seed | Fabaceae | KC343203 | KC343929 | KC344171 | KC343687 | KC343445 | Gomes <i>et al.</i> (2013) |
| | CBS 119639 | Man, abscess | ı | KC343202 | KC343928 | KC344170 | KC343687 | KC343444 | Gomes <i>et al.</i> (2013) |
| Diaporthe sp. 1 | CGMCC 3.18292 = LC 0771 | <i>Alnus</i> sp., pathogen | Betulaceae | KX986799 | KX999191 | KX999231 | KX999270 | KX999292 | This study |
| Diaporthe sp. 2 | CGMCC 3.18291 = LC 6140 | Acer sp., endophyte | Aceraceae | KX986799 | KX999191 | KX999231 | KX999270 | KX999292 | This study |
| | LC8112 | Acer sp., endophyte | Aceraceae | KY491547 | KY491557 | KY491567 | ı | KY491575 | This study |
| | LC8113 | Acer sp., endophyte | Aceraceae | КҮ491548 | KY491558 | KY491568 | | KY491576 | This study |
| Diaporthella corylina | CBS 121124 | Corylus sp., dying stems | Corylaceae | KC343004 | KC343730 | KC343972 | KC343488 | KC343246 | Gomes <i>et al.</i> (2013) |
| P. conorum | CBS 587.79 | Penus pentaphylla | Pinaceae | KC343153 | KC343879 | KC344121 | KC343637 | KC343395 | Gomes <i>et al.</i> (2013) |
| P. emicis | BRIP 45089a (ex-type) | Emex australis | Polygonaceae | JF957784 | JX275414 | JX275458 | | JX197449 | Udayanga <i>et al.</i> (2012) |
| P. fukushii | CBS 116953 | Pyrus pyrifolia | Roseceae | KC343147 | KC343873 | KC344115 | KC343631 | KC343389 | Gomes <i>et al.</i> (2013) |
| | BRIP 45089b | Emex australis | Polygonaceae | JQ619898 | JX275415 | JX275459 | | JX197450 | Udayanga <i>et al.</i> (2012) |
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-: not provided in literatures.





Fig. 3. *Diaporthe acutispora* (CGMCC 3.18285). **A–B.** 30-d-old culture on PNA medium. **C.** Conidiomata. **D–E.** Conidiophores. **F–G.** Alpha conidia. Bars: C = 100 μm; D–G = 10 μm.

TAXONOMY

Diaporthe acutispora Y.H. Gao & L. Cai, **sp. nov.** MycoBank MB820679 (Fig. 3)

Etymology: Named after the acute spores.

Diagnosis: Diaporthe acutispora is phylogenetically distinct and morphologically differs from species reported from the host genera *Coffea* and *Camellia* in the larger conidiophores and alpha conidia (Table 3).

Type: **China**: *Yunnan Province*: Aini Farm, on healthy leaves of *Coffea* sp., 20 Sep. 2014, *W.J. Duan* (HMAS 247086 – holotype, dried culture; CGMCC 3.18285 = LC 6161 – ex-type culture).

Description: On PNA: *Conidiomata* pycnidial, globose, brownish, embedded in tissue, erumpent at maturity, 99–473 µm diam, often with a yellowish conidial cirrus exuding from the ostioles. *Conidiophores* 10–34.5 × 2–3 µm, cylindrical, hyaline, septate, branched, straight or slightly curved, tapering towards the apex. *Alpha conidia* abundant in culture, 7–10.5 × 2–3 µm (\overline{x} = 8.4 ± 0.7 × 2.6 ± 0.2, *n* = 30), aseptate, hyaline, ellipsoidal to fusoid, multi-guttulate. *Beta conidia* not observed.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 7.5 mm diam/d. Colony entirely white at surface, reverse with pale brown pigmentation, white, fluffy aerial mycelium.

Additional material examined: China: Yunnan Province: Xishuangbanna, on healthy leaves of Camellia sasanqua, 20 Sep. 2014, W.J. Duan, culture LC 6142; *ibid*. culture LC 6160.

Diaporthe elaeagni-glabrae Y.H. Gao & L. Cai, sp. nov. MycoBank MB820680 (Fig. 4)

Etymology: Named after the host species Elaeagnus glabra.

Diagnosis: *Diaporthe elaeagni-glabrae* can be distinguished from the closely related species *D. elaeagni* (96 % in ITS, 93 % in *TEF1*, 94 % in *TUB*, 96 % in *HIS*, and 94 % in *CAL*) and *D. stictica* (96 % in ITS, 95 % in *TEF*, 97 % in *TUB*, 96 % in *HIS*, and 96 % in *CAL*) (Fig. 2). *Diaporthe elaeagni-glabrae* differs from other species recorded from *Elaeagnus* in the significantly longer alpha conidia (Table 3).

Type: **China**: *Jiangxi Province*: on diseased leaves of *Elaeagnus* glabra, 5 Sep. 2013, Y.H. Gao (HMAS 247089 – holotype, dried culture; CGMCC 3.18287 = LC 4802 – ex-type culture).

| Host genera | Species | Conidiomata (µm) | Conidiophores (µm) | Alpha conidia (µm) | Beta conidia (µm) | References |
|---|---|--|--|--|--|---|
| Coffea | P. coffeae | 200–250 | 12–16 × 2 | 8–9 × 2.5 | - | Uecker (1988) |
| Camellia | D. acutispora | 99-473 | 10-34.5 × 2-3 | 6.9–10.4 × 2.1–3.1 | 1 | This study |
| | D. amygdali | 160–220 × 120–300 | 7.4–36.3 × 1.5–3.2 | (4.18–)6.27–6.32(–9.64) × (1.63–)2.36– 2.38(–3.31) | | Diogo <i>et al.</i> (2010) |
| | D. apiculata | 74–195 (–416) | 9.0-12.5 × 1.5-2.5 | 6.5–10 × 2–3 | (20.0–)25.0-39.0 × 1.0–1.5 | Gao <i>et al.</i> (2016) |
| | D. compacta | 237–350 | 6.0-12.5 × 1.5-2.5 | 6–7.5 × 2–3 | 20.0–24.5 × 1.0–1.5 | Gao <i>et al.</i> (2016) |
| | D. discoidispora | 200 × 118 | 8.9–23.4 × 1.3–2.7 | 5.6-8 × 2.1-3.2 | 21.2–38.7 × 0.9–1.6 | Huang <i>et al.</i> (2015) |
| | D. eres | 200–250 | 10–15 × 2–3 | $(6-)6.5-8.5(-9) \times 3-4$ | (18–)22–28(29) × 1–1.5 | Udayanga <i>et al</i> . (2014b) |
| | D. foeniculacea | 560 × 350 | 10–13 × 1.5–3 | (5.4–)6.8–7(–9) × (2–)2.3–2.4(–3.1) | (16.8–)19.6–21(–24.2) × (1.1–)1.3– 1.4(–1.7) | Phillips (2003) |
| | D. foeniculina | 400-700 | 9–15(–18) × 1–2 | $(7.5-)8.5-9(-9.2) \times (2-)2.3-2.5(-2.7)$ | (20–)22–28(–29) × (1.1–)1.4–1.6(–2) | Udayanga <i>et al</i> . (2014c) |
| | D. hongkongensis | to 200 | 5-12 × 2-4 | $(5-)6-7(-8) \times (2-)2.5(-3)$ | 18–22 × 1.5–2 | Gomes <i>et al.</i> (2013) |
| | D. oraccinii | 400 | 10.5–22.5 × 1–2 | $5.5-7.5 \times 0.5-2$ | $24.5-31.0 \times 1.0-1.5$ | Gao <i>et al.</i> (2016) |
| | D. penetriteum | 176–486 | 13–21.5 (–27) × 1–2 | 4.5–5.5 × 1.5–2.5 | $16.5-27.5 \times 1.0-2.0$ | Gao <i>et al.</i> (2016) |
| | D. ueckerae | 150-200 | (9–)12–28(–30) × 1.5–2.5 | (6–)6.4–8.2(–8.6) × (2–)2.3–3 | 1 | Udayanga <i>et al.</i> (2014a) |
| | D. xishuangbanica | 180–310 | $13-34.5 \times 1.5-3$ | $7-9.5 \times 2.5-3.5$ | 1 | This study |
| | D. yunnanensis | 195–880 | | 3-6.5 × 1-2.5 | 13.5–33.5 × 1–1.5 | This study |
| | P. acaciicola | | | 7–9 × 3–3.5 | 1 | Diedicke (1911) |
| | P. theae | 40 × 25 | | 6–8 × 1.5–2 | 18–24 × 0.75 | Petch (1925) |
| Elaeagnus | P. arnoldiae | 900 × 500 | 6–12 × 1–2 | 5.5–11 × 1.5–2 | 15–20 | Uecker (1988) |
| | P. elaeagni | 500-750 | 20–25 × 1–1.5 | 6-10 ×2-3 | 1 | Uecker (1988) |
| | P. elaeagnicola | 175–413 × 83–185 | 10.0-22.5 × 1.5-2.7 | 6.0-7.4 × 1.7-2.2 | 19–43 × 0.7–1.2 | Chang <i>et al.</i> (2005) |
| | D. elaeagni-glabrae | 330-1170 | 16–28 × 1.5–2.5 | 6–13 × 1.5–3 | 7.5–22.5 × 1–2 | This study |
| | D. incompleta | 207–650 | 8–22 × 1–2.5 | | 19-44 × 0.5-1.5 | This study |
| Neolitsea | D. velutina | 69-428 | 10-23 × 1-2.5 | 5.5–10 × 2–2.5 | 11–27.5 × 0.5–1.5 | This study |
| AR, DP, FAU BRIP: Austra China. CGM maintained at at Institute of | : Isolates in culture co lian plant pathogen cu CC: China General Mi t the Westerdijk Institu Microbiology, CAS, CP | Illection of Systematic N Inture collection, Queen crobiological Culture Co te; LGMF: Culture colle nina; MFLUCC: Mae Fa | lycology and Microbiology La sland, Australia; CBS : Weste ollection; CMW : culture colle ction of Laboratory of Geneti h Luang University Culture C | boratory, USDA-ARS, Beltsville, Maryland, I ardijk Fungal Biodiversity Institute, Utrecht, 1 ction (CMW) of the Forestry and Agricultura cto Microorganisms, Federal University of collection; ZJUD : Zhe Jiang University, Chine | JSA; BCRC : Bioresource Collection and The Netherlands; CFCC : China Forestry ¹ al Biotechnology Institute; CPC : working Parana, Curitiba, Brazil; LC : Working col a. | Research Center, Taiwan; Cutture Collection Center, collection of Pedro Crous bilection of Lei Cai, housed |

Table 3. Synoptic characters of Diaporthe spp. referred to in this study.



Fig. 4. *Diaporthe elaeagni-glabrae* (CGMCC 3.18287). **A–B.** 14-d-old culture on PDA; **C.** Conidiomata; **D–H.** Conidiophores; **I.** Alpha conidia; **J.** Beta conidia. Bars: C = 100 µm; D–J = 10 µm.

Description: On PDA: Conidiomata globose, to 330–1170 µm, erumpent, with slightly elongated black necks, yellowish or dirty white, spiral conidial cirri extruding from ostioles. Conidiophores 16–28 × 1.5–2.5 µm, cylindrical, phialidic, septate, branched, sometimes inflated. Alpha conidia 6–13 × 1.5–3 µm ($\overline{X} = 8.3 \pm 1.4 \times 2.2 \pm 0.3$, n = 30), hyaline, fusiform or oval, usually biguttulate. Beta conidia 7.5–22.5 × 1–2 µm ($\overline{X} = 15.1 \pm 3.5 \times 1.2 \pm 0.2$, n = 40), hyaline, filiform, smooth, curved, base truncate.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 7 mm diam/d. Colony pale yellowish, greenish to brownish at the centre, reverse pale yellowish and brownish at the centre with age. Aerial mycelium white, sparse, fluffy, with irregular margin and visible conidiomata at maturity.

Additional material examined: China: Jiangxi Province: on diseased leaves of *Elaeagnus glabra*, 5 Sep. 2013, Y.H. Gao, culture LC 4806.

Diaporthe helianthi Munt.-Cvetk. *et al.*, *Nova Hedwigia* 34: 433 (1981). (Fig. 5)

Description: Sexual morph not produced. Conidiomata pycnidial globose to subglobose, dark brownish to black,

erumpent or immersed in medium, translucent conidia exuded from the ostioles, 110–380 µm diam. *Conidiophores* cylindrical, straight or sinuous, apical or base sometimes swelling, 11.5–23.5 × 1.8–3.5 µm ($\overline{X} = 16 \pm 3 \times 2.5 \pm 0.5$, n = 30). *Beta conidia* filiform, hamate or slightly curved, base truncate, tapering towards one apex, 11.5–32 × 0.5–2 µm ($\overline{X} = 20 \pm 7.5 \times 1 \pm 0.4$, n = 20). *Alpha conidia* not observed.

Culture characters: Cultures on PDA at 25 °C in dark, with 12/12 h alternation between daylight and darkness pure white (surface) and pale yellow to cream (reverse). Colony pellicular, forming less pigmented sectors, with concentric rings of gummy mycelium. Growth rate was 10.5 mm diam/d.

Material examined: Ukraine: from seeds of Helianthus annuus, 30 Oct. 2015, W.J. Duan culture LC 6173. – Japan: Lagerstroemia indica, 30 Oct. 2015, W.J. Duan, culture LC 6185.

Notes: Diaporthe helianthi, responsible for stem canker and grey spot disease of sunflower (*Helianthus annuus*) (Muntanola-Cvetkovic *et al.* 1981), has been listed in the Chinese quarantine directory. There is increasing evidence that this serious sunflower pathogen is being quickly and globally disseminated with international trade. The cases reported here were intercepted from imported sunflower seeds from Ukraine and *Lagerstroemia indica* from Japan.



Fig. 5. *Diaporthe helianthi* (LC 6185). **A–B.** 7-d-old culture on PDA; **C.** Conidiomata; **D–F.** Conidiophores; **G–H.** Beta conidia. Bars: C = 100 μm; D–H = 10 μm.

Diaporthe incompleta Y.H. Gao & L. Cai, **sp. nov.** MycoBank MB820681 (Fig. 6)

Etymology: Named after the absence of alpha conidia.

Diagnosis: Diaporthe incompleta is phylogenetically distinct and differs morphologically from other species recorded from *Elaeagnus* and *Camellia* in the much longer beta conidia (Table 3).

Type: **China**: *Yunnan Province*: Xishuangbanna, on diseased of *Elaeagnus glabra*, 19 Apr. 2015, *F. Liu* (HMAS 247088 – holotype, dried culture; CGMCC 3.18288 = LC 6754 – ex-type culture).

Description: Conidiomata pycnidial, subglobose to globose, brownish to black, 207–650 µm diam, cream to pale luteous

conidial droplets exuding from the central ostioles. Conidiophores 8–22 × 1–2.5 µm, cylindrical, hyaline, septate, unbranched, smooth, slightly curved, tapering towards apex. Alpha conidia not observed. Beta conidia 19–44 × 0.5–1.5 µm ($\overline{X} = 30.5 \pm 8.7 \times 1.1 \pm 0.4$, n = 30), smooth, hyaline, filiform, base subtruncate, straight or curved.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 16.5 mm diam/d. Colony entirely white, flat, reverse pale yellowish, becoming brownish zoned at the centre with age. Aerial mycelium white, cottony, margin lobate, conidiomata visible at maturity.

Additional material examined: **China**: Yunnan Province: Xishuangbanna, on diseased leaves of *Camellia sinensis*, 19 Apr. 2015, *F. Liu*, culture LC 6706.



Fig. 6. *Diaporthe incompleta* (CGMCC 3.18288). **A.** Leaves of host plant; **B–C.** 7-d-old culture; **D.** Conidiomata; **E–F.** Conidiophores; **G.** Beta conidia. Bars: D = 100 μm; E–G = 10 μm.

Diaporthe podocarpi-macrophylli Y.H. Gao & L. Cai, sp. nov. MycoBank MB820682 (Fig. 7)

Etymology: Named after the host plant *Podocarpus macrophyllus*.

Diagnosis: *Diaporthe podocarpi-macrophylli* can be distinguished from the phylogenetically closely related species *D. pseudophoenicicola* (97 % identity in ITS, 90 % in *TEF1*, 98 % in *TUB*, 97 % in *HIS*, and 97 % in *CAL*). Morphologically, *D. podocarpi-macrophylli* differs from other species occurring on the host genera *Podocarpus* and *Olea*, i.e. *D. cinerascens* and *Phomopsis podocarpi* in its wider and shorter alpha co-



Fig. 7. *Diaporthe podocarpi-macrophylli* (CGMCC 3.18281). A–B. 30-d-old culture on PDA; C. Conidiomata; D–F. Conidiophores; G–I. Alpha and beta conidia. Bars: C = 100 µm; D–I = 10 µm.

nidia and the presence of beta conidia (Chang *et al.* 2005, Gomes *et al.* 2013; https://nt.ars-grin.gov/fungaldatabases/).

Type: **Japan**: on healthy leaves of *Podocarpus macrophyllus*, 20 Sep. 2014, *W.J. Duan* (HMAS 247084 – holotype, dried culture; CGMCC 3.18281 = LC 6155 – ex-type culture).

Description: Conidiomata pycnidial in culture on PDA, solitary or aggregated, deeply embedded in the PDA, erumpent, dark brown to black, 222–699 µm diam, yellowish translucent conidial drops exuding from the ostioles. Alpha conidiophores $6-18 \times 1.5-3 \text{ µm}$ ($\overline{X} = 12.3 \pm 2.6 \times 2.1 \pm 0.3, n = 30$), hyaline, septate, branched, cylindrical, straight to sinuous, sometimes inflated, occurring in dense clusters. Beta conidiophores

10.5–27 × 1.5–2.5 µm (\overline{x} = 15.3 ± 4.3 × 2.1 ± 0.3, *n* = 30), cylindrical to clavate, hyaline, septate, branched, smooth, straight. *Alpha conidia* 3.5–8.5 × 1–3 µm (\overline{x} = 6.3 ± 1.7 × 2.1 ± 0.7, *n* = 50), unicellular, aseptate, fusiform, hyaline, usually biguttulate and acute at both ends. *Beta conidia* 8.5–31.5 × 0.5–2 µm (\overline{x} = 19.5 ± 7.1 × 1.1 ± 0.4, *n* = 30), hyaline, aseptate, eguttulate, filiform, curved, tapering towards both ends, base truncate.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 12.5 mm diam/d. Colony at first white, becoming cream to yellowish, flat, with dense and felted mycelium, reverse pale brown with brownish dots with age, with visible solitary or aggregated conidiomata at maturity.



Fig. 8. Diaporthe undulata (CGMCC 3.18293). A. Leaves of host plant; B–C. 30-d-old culture on PNA medium; D. Conidiomata; E. Conidiophores; F–G. Alpha conidia. Bars: D = 100 μm; E–G = 10 μm.

Additional material examined: Japan: on healthy leaves of *Podocarpus macrophyllus*, 20 Sep. 2014, *W.J. Duan*, culture LC 6141; *ibid*. culture LC 6144; *ibid*. culture LC 6156; *ibid*. culture LC 6157. – **China:** *Zhejiang Province*: on healthy leaves of *P. macrophyllus*, 10 Jul. 2015, *W.J. Duan*, culture LC 6194; *ibid*. culture LC 6195; *ibid*. culture LC 6196; *ibid*. culture LC 6197; *ibid*. culture LC 6198; *ibid*. culture LC 6199; *ibid*. culture LC 6200; *ibid*. culture LC 6201; *ibid*. culture LC 6202; *ibid*. culture LC 6235. – **Italy:** on healthy leaves of *Olea europaea*, 20 Sep. 2014, *W.J*. Duan, culture LC 6229.

Diaporthe undulata Y.H. Gao & L. Cai, **sp. nov.** MycoBank MB820683 (Fig. 8)

Etymology: Named after the colony's undulate margin.

Diagnosis: Diaporthe undulata differs from the most closely related species, *D. biconispora*, in several loci (94 % in ITS, 84 % in *TEF1*, and 93 % in *TUB*), and from other *Diaporthe* species in the obpyriform conidiophores and shorter and wider alpha conidia (Table 3).

Type: **China-Laos border**: on diseased leaves of unknown host, 19 Apr. 2014, *F. Liu* (HMAS 247091 – holotype, dried culture; CGMCC 3.18293 = LC 6624 – ex-type culture).

Description: Conidiomata pycnidial, irregular, embedded in the needle, erumpent, necks, hairy, 282–543 µm long, coated with short hyphae, one to several necks forming from a single pycnidium. *Conidiophores* obpyriform, hyaline, phiailidic, septate, branched, 5–17.5 × 2–3 µm ($\overline{X} = 9.7 \pm 4.0 \times 2.4 \pm 0.5$, n = 20). *Alpha conidia* ellipsoid, hyaline, biguttulate, rounded at both ends, 5–6.5 × 2–3 ($\overline{X} = 5.8 \pm 0.4 \times 2.3 \pm 0.3$, n = 50). *Beta conidia* not observed.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 10.5 mm diam/d. Colony entirely white, reverse pale yellowish and dark brownish at the centre with age. Aerial mycelium white, cottony, dense, with undulate margin and visible conidiomata at maturity.

Additional material examined: China-Laos border: unknown host, 19 Apr. 2014, F. Liu, culture LC 8110; *ibid*. culture LC 8111.

Diaporthe velutina Y.H. Gao & L. Cai, **sp. nov.** MycoBank MB820684 (Fig. 9)

Etymology: Named after the felted colony.

Diagnosis: *Diaporthe velutina* is distinguished from *D. anacardii* in the ITS, *TEF1*, *TUB* and *HIS* loci (99 % in ITS, 95 % in *TEF1*, 99 % in *TUB*, and 98 % in *HIS*), and from





Fig. 9. *Diaporthe velutina* (CGMCC 3.18286). **A.** Diseased leaves; **B–C.** 30-d-old culture on PDA; **D.** Conidiomata; **E.** Conidiophores; **E.** Alpha and beta conidia. Bars: D = 100 μm; E–F = 10 μm.

other *Diaporthe* species reported from *Camellia sinensis* in the more variable size of the alpha conidia (Table 3).

Type: **China**: *Jiangxi Province*: on diseased leaves of *Neolitsea* sp., 5 Sep. 2013, *Y.H. Gao* (HMAS 247087 – holotype, dried culture; CGMCC 3.18286 = LC 4421 – ex-type culture).

Description: Conidiomata pycnidial, globose, black, embedded in PDA, aggregated in clusters, 69–428 µm diam, cream translucent drop of conidia exuded from the central ostioles. *Conidiophores* 10–23 × 1–2.5 µm, cylindrical, hyaline, branched, densely aggregated, slightly tapering towards the apex, sometimes slightly curved. *Alpha conidia* 5.5–10 × 2–2.5 µm ($\overline{X} = 6.9 \pm 0.9 \times 2.2 \pm 0.2$, n = 50), unicellular, aseptate, hyaline, fusoid to ellipsoid or clavate, bi-guttulate or multi-guttulate. *Beta conidia* 11–27.5 × 0.5–1.5 µm ($\overline{X} = 16.1 \pm 5.0 \times 0.8 \pm 0.4$, n = 30), smooth, hyaline, apex acutely rounded, curved.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 18.75 mm diam/d. Colony entirely white, surface mycelium greyish to brownish at the centre,

dense, felted, conidiomata erumpent at maturity, reverse centre yellowish to brownish.

Additional material examined: **China**: Jiangxi Province: Yangling, on diseased leaves of *Neolitsea* sp., 5 Sep. 2013, *Y.H. Gao*, culture LC 4419; *ibid*. culture LC 4422; Gannan Normal University, unknown host, 23 Apr. 2013, *Q. Chen*, culture LC 4788; Fengshan, on diseased leaves of *Callerya cinerea*, 5 Sep. 2013, *Y.H. Gao*, culture LC 4641. *Yunnan Province*: Xishuangbanna, on diseased leaves of *Camellia sinensis*, 19 Apr. 2015, *F. Liu*, culture LC 6708; *loc. cit.*, on healthy leaves of *C. sinensis*, 21 Apr. 2015, *F. Liu*, culture LC 6519.

Diaporthe xishuangbanica Y.H. Gao & L. Cai, sp. nov. MycoBank MB820685 (Fig. 10)

Etymology: Named after the locality, Xishuangbanna.

Diagnosis: *Diaporthe xishuangbanica* can be distinguished from the phylogenetically closely related *D. tectonigena* in several loci (98 % in ITS, 90 % in *TEF1*, and 96 % in *TUB*)





Fig. 10. Diaporthe xishuangbanica (CGMCC 3.18283). A–B. 7-d-old culture on PDA; C–D. 30-d-old culture on PNA medium; E. Conidiomata;
F–K. Conidiophores; L–N. Alpha conidia. Bars: E = 100 μm; F–N = 10 μm.

(Fig. 2), and from other *Diaporthe* species reported from *Camellia* in the longer conidiophores and alpha conidia (Table 3).

Type: **China**: *Yunnan Province*: Xishuangbanna, on diseased leaves of *Camellia sinensis*, 19 Apr. 2015, *F. Liu* (HMAS 247083 – holotype, dried culture; CGMCC 3.18283 = LC 6744 – ex-type culture).

Description: Conidiomata pycnidial, globose, 180–310 µm diam, scattered on the pine needle. Conidiophores cylindrical, 13–34.5 × 1.5–3 µm (\overline{x} = 20.9 ± 5.2 × 2.1 ± 0.3, *n* = 40), branched, septate, straight, sometimes sinuous or lateral. Alpha conidia 7–9.5 × 2.5–3.5 µm (\overline{x} = 8.3 ± 0.7 × 2.8 ± 0.3, *n* = 30), fusiform, hyaline, multi-guttulate. Beta conidia not observed.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 17.5 mm diam/d. Colony entirely white, reverse pale yellowish to greenish. Aerial mycelium white, velvety, margin well defined, with visible conidiomata at maturity.

Additional material examined: China: Yunnan Province: Xishuangbanna, on diseased leaves of Camellia sinensis, 19 Apr. 2015, F. Liu, culture LC 6707 (CGMCC 3.18282).

Diaporthe yunnanensis Y.H. Gao & L. Cai, **sp. nov.** MycoBank MB820686 (Fig. 11)

Etymology: Named after the location where the fungus was collected, Yunnan Province.

Diagnosis: Diaporthe yunnanensis can be distinguished from the phylogenetically closely related *D. siamensis* (96 % in ITS, 91 % in *TEF1*, and 94 % in *TUB*) (Fig. 2), and from other *Diaporthe* species reported on the genus *Camellia* in the smaller alpha conidia (Table 3).

Type: **China**: *Yunnan Province*: Xishuangbanna, on healthy leaves of *Coffea* sp., 20 Sep. 2014, *W.J. Duan* (HMAS 247096 – holotype, dried culture; CGMCC 3.18289 = LC 6168 – ex-type culture).

Description: Conidiomata pycnidial, 195–880 µm diam, globose or irregular, erumpent, solitary or aggregated together, dark brown to black. *Conidia* exuding from the pycnidia in white to cream drops. *Conidiophores* cylindrical, straight or slightly curved. *Alpha conidia* 3–6.5 × 1–2.5 µm ($\overline{x} = 5.5 \pm 1 \times 2 \pm 0.5$, n = 30), fusiform, hyaline, biguttulate, with one end obtuse and the other acute. *Beta conidia* 13.5–33.5 × 1–1.5 µm ($\overline{x} = 27.5 \pm 5.5 \times 1.5 \pm 0.3$, n = 30), hyaline,



Fig. 11. *Diaporthe yunnanensis* (fCGMCC 3.18289). **A–B.** 7-d-old culture on PDA; **C.** Conidiomata; **D.** Conidiophores; **E.** Alpha and beta conidia; **F.** Beta conidia. Bars: C = 100 μm; D–F = 10 μm.

aseptate, hamate or curved, base truncate.

Culture characters: Colonies on PDA flat, with a moderate growth rate of 5.5 mm diam/d, with abundant dirty white and yellowish pigmented mycelium, dry, felted, extensive thin, and in reverse the centre cream, with zone rings of pale to dark brownish pigmentation.

Additional material examined: China: Yunnan Province: Xishuangbanna, on healthy leaves of *Coffea* sp., 20 Sep. 2014, *W.J. Duan*, culture LC 8106; *ibid*. culture LC 8107.

Diaporthe sp. 1 (Fig. 12)

(119.12)

Description: Conidiomata pycnidial, subglobose to globose, dark brown to black, deeply embedded in the substrate, scattered on the substrate surface, embedded in PDA, clusters in group of 2–7 pycnidia, 268–509 µm, yellowish drop of conidia diffusing from the central ostioles. Conidiophores 6.5–19.5 × 1–3 µm, cylindrical, hyaline, septate, branched, straight to sinuous, base inflated, slightly tapering towards the apex. Alpha conidia 7.5–13.5 × 2–3.5 µm ($\overline{X} = 9.9 \pm 1.4 \times 2.8 \pm 0.4$, n = 30), unicellular, hyaline, fusoid to ellipsoid or clavate, two or several large guttulae

observed, base subtruncate. *Beta conidia* 15–40.5 × 1–2.5 μ m (\overline{x} = 26.0 ± 5.8 × 1.8 ± 0.5, *n* = 30), smooth, hyaline, curved, base subtruncate, tapering towards one apex.

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate 7.83 mm diam/day. Colony entire, white to dirty pink, cottony, sparse, brownish to black conidiomata erumpent at maturity, coated with white hypha, granular at margin, reverse pale brown, with brownish dots when maturity.

Material examined: **China**: *Zhejiang Province*: Gutianshan Nature Reserve (29°20' N 18°14' E), on leaves of *Alnus mill*, Jan. 2010, *Y.Y. Su* (culture CGMCC 3.18292 = LC 0771).

Notes: The present culture belongs to the *Diaporthe eres* complex, which is reported from a very wide range of host plants and includes mostly opportunistic pathogens or secondary invaders on saprobic host substrata (Udayanga *et al.* 2014a, Gao *et al.* 2016). Species delimitation in this complex is currently unclear. Udayanga *et al.* (2015) accepted nine phylogenetic species in the *D. eres* complex, including *D. alleghaniensis*, *D. alnea*, *D. bicincta*, *D. celastrina*, *D. eres*, *D. helicis*, *D. neilliae*, *D. pulla*, and *D. vaccinia*. Gao *et al.* (2016) examined 17 isolates belonging to the *D. eres*



Fig. 12. *Diaporthe* sp. 1 (CGMCC 3.18292). **A.** Leaves of host plant; **B–C.** 30-d-old culture on PDA; **D.** Conidiomata; **E–F.** Conidiophores; **G.** Beta conidia; **H–I.** Alpha conidia. Bars: D = 100 μm; E–I = 10 μm.

complex, and reported that many presented intermediate morphology among "species" and the phylogenetic analyses often resulted in ambiguous clades with short branch and moderate statistical support. The identification of taxa in this group remains unresolved.

Diaporthe sp. 2

Culture characters: Cultures incubated on PDA at 25 °C in darkness, growth rate, slow, 3.83 mm diam/d. Colony low, convex, entire white to yellowish, reverse brownish. Aerial mycelia white, dry, downy, with near-circular margin.

Material examined: Japan: on leaves of *Acer* sp., 20 Sep. 2014, *W.J. Duan*, culture CGMCC 3.18291 = LC 6140, culture LC 8112; *ibid.* culture LC 8113.

Notes: Although three isolates clustered in a clade distinctly different from known species in the genus included, they are not formally described because they were sterile. *Diaporthe* sp. 2 shares a low homology to the most closely related species, *D. rhoina* (95 % in ITS, 87 % in *TEF1*, 97 % in *TUB*, 94 % in *HIS*, and 95 % in *CAL*). Five *Diaporthe* species are so far only known from the sterile state, including *D. endophytica*, *D. inconspicua*, *D. infecunda*, *D. asheicola*, and *D. sterilis* (Gomes *et al.* 2013, Lombard *et al.* 2014).

Diaporthe averrhoae (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821437

Basionym: Phomopsis averrhoae C.Q. Chang et al., Mycosystema 24: 6 (2005).

Type: **China**: *Fujian Province*: on living branches of *Averrhoa carambola*, *Y.H. Cheng* (SCHM 3605 – holotype; AY618930, ITS sequence derived from the holotype SCHM 3605).

Diaporthe camptothecae (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821438

Basionym: Phomopsis camptothecae C.Q. Chang et al., Mycosystema 24: 145 (2005).

Type: **China**: *Hunan Province*: on living branches of *Camptotheca acuminate*, *L.J. Luo* (SCHM 3611 – holotype; AY622996, ITS sequence derived from the holotype SCHM 3611).

Diaporthe chimonanthi (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821439

Basionym: Phomopsis chimonanthi C.Q. Chang et al., Mycosystema 24: 146 (2005).

Type: **China**: *Hunan Province*: on living branches of *Chimonanthus praecox*, *C.Q. Chang* (SCHM 3614 – holotype; AY622993, ITS sequence derived from the holotype SCHM 3614).

Diaporthe eucommiae (F.X. Cao *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821440

Basionym: Phomopsis eucommiae F.X. Cao et al., J. Middle-South China Forestry Coll. **10**: 34 (1990); as 'eucommi'.

Type: **China**: *Guangdong Province*: from leaves of *Eucommia ulmoides*, *F.X. Cao* (SCHM 0020 – holotype; AY601921, ITS sequence derived from the holotype SCHM 0020).

Diaporthe eucommiicola (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821441

Basionym: Phomopsis eucommiicola C.Q. Chang et al., Mycosystema 24: 147 (2005).

Type: **China**: *Hunan Province*: on living branches of *Eucommia ulmoides* and *Styrax hypoglauca*, *L.J. Luo* (SCHM 3607 – holotype; AY578071, ITS sequence derived from the holotype SCHM 3607).

Diaporthe glabrae (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821443

Basionym: Phomopsis glabrae C.Q. Chang et al., Mycosystema 24: 8 (2005).

Type: **China**: *Fujian Province*: on living branches of *Bougainvillea glabra*, *Y.H. Cheng* (SCHM 3622 – holotype; AY601918, ITS sequence derived from the holotype SCHM 3622).

Diaporthe lagerstroemiae (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821444

Basionym: Phomopsis lagerstroemiae C.Q. Chang et al., Mycosystema 24: 148 (2005).

Type: **China**: *Hunan Province*: on living branches of *Lagerstroemia indica*, *C.Q. Chang* (SCHM 3608 – holotype; AY622994, ITS sequence derived from the holotype SCHM 3608).

Diaporthe liquidambaris (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821446

Basionym: Phomopsis liquidambaris C.Q. Chang et al., Mycosystema 24: 9 (2005).

Type: **China**: *Fujian Province*: on living branches of *Liquidambar formosana*, Y.H. Cheng (SCHM 3621 – holotype; AY601919, ITS sequence derived from the holotype SCHM 3621).

Diaporthe loropetali (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821448

Basionym: Phomopsis loropetali C.Q. Chang et al., Mycosystema 24: 148 (2005).

Type: **China**: Hunan Province: on living branches of *Loropetalum chinense*, *C.Q. Chang* (SCHM 3615 – holotype; AY601917, ITS sequence derived from the holotype SCHM 3615).

Diaporthe magnoliicola Y.H. Gao & L. Cai, nom. nov. MycoBank MB821459

Replaced name: Phomopsis magnoliae M.M. Xiang et al., My-cosystema **21**: 501 (2002). *Type*: **China**: *Guangdong Province*: on leaves of *Magnolia coco*, *Z.D. Jiang* (SCHM 3001 – holotype; AY622995, ITS sequence derived from the holotype SCHM 3001).

Note: The epithet *magnoliae* is occupied, so *Diaporthe magnoliicola* is proposed as a replacement name.

Diaporthe michelina (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, **comb. nov.**

MycoBank MB821460

Basionym: Phomopsis michelina C.Q. Chang et al., Mycosystema 24: 9 (2005); as 'micheliae'.

Type: **China**: *Fujian Province*: on living branches of *Michelia alba*, *Y.H. Cheng* (SCHM 3603 – holotype; AY620820, ITS sequence derived from the holotype SCHM 3603).

Diaporthe phyllanthicola (C.Q. Chang *et al.*) Y.H. Gao & L. Cai, comb. nov.

MycoBank MB821461

Basionym: Phomopsis phyllanthicola C.Q. Chang et al., Mycosystema 24: 10 (2005).

Type: **China**: *Fujian Province*: on living branches of *Phyllanthus emblica*, *Y.H. Cheng* (SCHM 3680 – holotype; AY620819, ITS sequence derived from the holotype SCHM 3680).

DISCUSSION

In this study, eight new species of *Diaporthe* are introduced, having been isolated from various plant hosts collected in different countries. Twelve *Phomopsis* species described from China were subjected to molecular analysis, and transferred to *Diaporthe* to conform to the "one fungus one name" rule (Udayanga *et al.* 2011, Rossman *et al.* 2016). To address the taxonomy of the other *Phomopsis* species described from China, neo- or epitypes will need to be designated to resolve their position and confirm their placement in *Diaporthe*.

Previous taxonomic studies in *Diaporthe* (syn. *Phomopsis*) have been primarily based on morphology, which has been shown to be unnatural in reflecting evolutionary history due to the simple and plastic morphological characters (Gao et al. 2015). The same applies to many other genera of ascomycetes. For example, species referred to Phoma have been shown to be highly polyphyletic and scattered throughout at least six families within Pleosporales (Aveskamp et al. 2010, Chen et al. 2015). Although Diaporthe was previously thought to be monophyletic based on its typical and unique Phomopsis asexual morph and diaporthalean sexual morph (Gomes et al. 2013), a paraphyletic nature is revealed in the present study (Fig. 1). Several genera, notably Ophiodiaporthe (Fu et al. 2013), Pustulomyces (Dai et al. 2014), Phaeocytostroma, and Stenocarpella (Lamprecht et al. 2011), are shown to be embedded in Diaporthe s. lat., none of which present an independent lineage from Diaporthe as currently circumscribed (Fig. 1). These genera were established based on their morphological characteristics (Vasilyeva et al. 2007, Lamprecht et al. 2011, Fu et al. 2013, Dai et al. 2014). For example, Ophiodiaporthe produces only one type of globose or subglobose conidia that differs from the dimorphic (fusiform and filiform) conidia of Diaporthe (Fu et al. 2013); Phaeocytostroma and Stenocarpella produce pigmented alpha conidia which differ from the hyaline conidia of Diaporthe (Lamprecht et al. 2011); Pustulomyces produces larger, straight or sigmoid conidia (Dai et al. 2014). Phaeocytostroma and Stenocarpella were originally suspected to be members of Botryosphaeriaceae (Botryosphaeriales) because of their pigmented alpha conidia and diplodia-like morphology (Crous et al. 2006). However, they were subsequently allocated to Diaporthales based on phylogenetic analysis (Lamprecht et al. 2011), which is confirmed in this study.

The large "Diaporthe" clade embedded with the heterogeneous genera Ophiodiaporthe, Pustulomyces, Phaeocytostroma, and Stenocarpella is probably a typical example of divergent evolution in morphological characters. Such an evolution could have been driven by host and/or environmental adaptations. For example, the monotypic Ophiodiaporthe is associated with Cyathea lepifera (a fern), while Pustulomyces is bambusicolous (Dai et al. 2014). On the contrary, none of the previously named over 1 900 Diaporthe / Phomopsis species was recorded from a fern or Bambusaceae (https:// nt.ars-grin.gov/fungaldatabases/). It is therefore reasonable to speculate that the speciation of Ophiodiaporthe and Pustulomyces, as well as the distinctly different morphologies from their close Diaporthe allies, are the consequences of evolutionary adaption to new hosts. Similarly, Phaeocytostroma and Stenocarpella are mainly restricted to maize (Zea mays), causing root stalk and cob rot (Stovold et al. 1996, Lamprecht et al. 2011).

Splitting *Diaporthe* into many smaller genera would achieve monophyletic groupings, but would also create many additional problems. The "new genera" split from *Diaporthe* would have no recognisable morphological distinctions in either sexual or asexual morphs. In addition, splitting *Diaporthe* into many smaller genera will result in numerous name changes, which is certainly an unfavourable option for both mycologists and plant pathologists.

Diaporthe has long been well-known to include plant pathogens, some on economically important hosts, such as Helianthus annuus (sunflower; Thompson et al. 2011) and Glycine max (soybean; Santos et al. 2011). However, the number of known endophytic Diaporthe species has increased rapidly in recent years (Huang et al. 2015, Gao et al. 2016). Wang et al. (2013) concluded that our current knowledge of the ecology and biology of endophytic Diaporthe species is just the "tip of the iceberg". In 2013, a new sterile endophytic species, Diaporthe endophytica, was formally named (Gomes et al. 2013). The research on Citrus conducted by Huang et al. (2015) recorded seven apparently undescribed endophytic Diaporthe species. Inspection of Diaporthe species on Camellia sinensis resulted in the description of four new and five known species, all occurring as endophytes (Gao et al. 2016). Because many of these plant pathogenic Diaporthe species are commonly encountered as sterile endophytes, a multigene DNA database will be essential to aid in their future identification.

Accurate identification of fungal pathogens is the basis of quarantine and disease control (Udayanga *et al.* 2011).

Thompson *et al.* (2011) reported significant damage to sunflower in Australia caused by *Diaporthe helianthi* which was originally only known from Europe (former Yugoslavia), and is apparently an invasive species in Australia. *Diaporthe helianthi* is listed in the Chinese quarantine directory, and has long been considered a predominant disease limiting production in Europe (Desanlis *et al.* 2013). Duan *et al.* (2016) reported this pathogen on sunflower seeds imported from Ukraine into China. Here, we report another interception of *D. helianthi* from *Lagerstroemia indica* imported from Japan to China. This serves as additional evidence of how quickly serious pathogens such as *Diaporthe* species can be distributed as endophytes or latent pathogens with global trade.

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