Morphological and phylogenetic analyses reveal three new species of *Diaporthe* from Yunnan, China

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Abstract
Species of *Diaporthe* have often been reported as plant pathogens, endophytes or saprobes, commonly isolated from a wide range of plant hosts. Sixteen strains isolated from species of ten host genera in Yunnan Province, China, represented three new species of *Diaporthe*, *D. chrysalidocarpi*, *D. machili* and *D. pometiae* as well as five known species *D. arecae*, *D. hongkongensis*, *D. middletonii*, *D. osmanthi* and *D. pandanicola*. Morphological comparisons with known species and DNA-based phylogenies based on the analysis of a multigene (ITS, TUB, TEF, CAL and HIS) dataset support the establishment of the new species. This study reveals that a high species diversity of *Diaporthe* with wide host ranges occur in tropical rainforest in Yunnan Province, China.

Keywords
Diaporthaceae, Diaporthales, phylogeny, taxonomy, three taxa new to science

Introduction

The genus *Diaporthe* (Diaporthaceae Diaporthales) with asexual morphs previously known as *Phomopsis* spp. is based on the type species *Diaporthe eres* Nitschke (1870) from *Ulmus* sp. in Germany. Rossman et al. (2015) proposed to use the name *Diaporthe* over *Phomopsis* in the context of the one fungus – one name initiative, be-

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cause it was described first, is encountered commonly in literature and includes the majority of known species. The sexual morph of *Diaporthe* is characterised by immersed ascomata and an erumpent pseudostroma with elongated perithecial necks; asci are unitunicate, clavate to cylindrical; and ascospores are fusoid, ellipsoidal to cylindrical, hyaline, biseriate to uniseriate in the ascus, sometimes with appendages (Udayanga et al. 2011; Senanayake et al. 2017, 2018). The asexual morph is characterised by ostiolate pycnidia with cylindrical phialides often producing three types of hyaline, aseptate conidia called α-conidia, β-conidia and γ-conidia (Udayanga et al. 2011; Gomes et al. 2013). The α-conidia and β-conidia are produced frequently, but the γ-conidia are rarely observed (Gomes et al. 2013; Guarnaccia and Crous 2017; Guo et al. 2020).

Currently, more than 1100 epithets of *Diaporthe* are listed in Index Fungorum (http://www.indexfungorum.org/; accessed 1 Nov. 2020), but only one-fifth of these taxa have been well-studied with ex-type cultures and supplementary DNA barcodes (Guo et al. 2020; Yang et al. 2020; Zapata et al. 2020). Species of *Diaporthe* are widely distributed and have a broad range of hosts including economically significant agricultural crops and ornamental plants such as species of *Camellia, Castanea, Citrus, Glycine, Helianthus, Juglans, Persea, Pyrus, Vaccinium, Vitis* and many more (van Rensburg et al. 2006; Santos and Phillips 2009; Crous et al. 2011a, b, 2016; Santos et al. 2011; Thompson et al. 2011; Grasso et al. 2012; Huang et al. 2013; Lombard et al. 2014; Gao et al. 2015, 2016, 2017; Udayanga et al. 2012, 2015; Guarnaccia et al. 2016; Dissanayake et al. 2017; Guarnaccia and Crous 2017; Fan et al. 2018; Senanayake et al. 2018; Guo et al. 2020). *Diaporthe* species have been reported as destructive plant pathogens, harmless endophytes or saprobes (Murali et al. 2006; Udayanga et al. 2012; Gomes et al. 2013; Ménard et al. 2014; Guarnaccia et al. 2016; Torres et al. 2016; Senanayake et al. 2018). However, the biology and lifestyle of some of these fungi remain unclear (Vilka and Volkova 2015).

In the past, methods of species identification of *Diaporthe* had previously been based only on host as well as morphological characters such as the size and shape of ascomata and conidiomata. Nowadays, molecular phylogenetic studies demonstrate that determining species boundaries only by morphological characters is not possible due to lack of host specificity and their variability under changing environmental conditions (Gomes et al. 2013). Phylogenetic analysis using a five-locus dataset (ITS-TUB-TEF-CAL-HIS) has been determined to be the optimal combination to identify species of *Diaporthe* species, as revealed by Santos et al. (2017). Many *Diaporthe* species are described based on a polyphasic approach together with morphological characterisation (Rehner and Uecker 1994; Udayanga et al. 2011; Gao et al. 2017; Guarnaccia and Crous 2017; Yang et al. 2018a, 2020; Crous et al. 2020; Dayarathne et al. 2020; Guo et al. 2020; Hyde et al. 2020; Li et al. 2020; Zapata et al. 2020).

The aim of this study was to explore the diversity of *Diaporthe* species from symptomatic leaves of plants in Yunnan Province. We present three novel species and five known species of *Diaporthe*, collected from species belonging to ten host genera, based on morphological characters and phylogenetic analysis.
Materials and methods

Isolation and morphological studies

Leaves of samples were collected in Yunnan Province, China. Isolations from surface sterilized leaf tissues were conducted following the protocol of Gao et al. (2014). Tissue fragments (5 × 5 mm) were taken from the margin of leaf lesions and surface-sterilized by immersing them in 75% ethanol solution for 1 min, 5% sodium hypochlorite solution for 30 s, and then rinsing in sterile distilled water for 1 min. The pieces were dried with sterilized paper towels and placed on potato dextrose agar (PDA) (Cai et al. 2009). PDA plates (90 mm) were incubated in an incubator at 25 °C for 2–4 days, and hyphae were picked out of the periphery of the colonies and inoculated onto new PDA plates.

Following 2–3 weeks of incubation, photographs of colonies were taken at 7 days and 15 days using a Powershot G7X mark II digital camera. Colour notations was done using the colour charts of Rayner (1970). Micromorphological characters were observed using an Olympus SZX10 stereomicroscope and Olympus BX53 microscope, both fitted with Olympus DP80 high definition colour digital cameras to document fungal structures. All fungal strains were stored in 10% sterilized glycerin at 4 °C for further studies. Voucher and type specimens were deposited in the Herbarium of Plant Pathology, Shandong Agricultural University (HSAUP). Living cultures were deposited in the Shandong Agricultural University Culture Collection (SAUCC). Taxonomic information of the new taxa was submitted to MycoBank (http://www.mycobank.org).

DNA extraction and amplification

Genomic DNA was extracted from fungal mycelium on PDA, using a modified cetyltrimethylammonium bromide (CTAB) protocol as described in Guo et al. (2000). The internal transcribed spacer regions with intervening 5.8S nrRNA gene (ITS), part of the beta-tubulin gene region (TUB), partial translation elongation factor 1-alpha (TEF), histone H3 (HIS) and calmodulin (CAL) genes were amplified and sequenced by using primers pairs ITS4/ITS5 (White et al. 1990), Bt2a/Bt2b (Glass and Donaldson 1995), EF1-728F/EF1-986R (Carbone and Kohn 1999), CAL-228F/CAL-737R (Carbone and Kohn 1999) and CYLH3F/H3-1b (Glass and Donaldson 1995; Crous et al. 2004), respectively.

PCR was performed using an Eppendorf Master Thermocycler (Hamburg, Germany). Amplification reactions were performed in a 25 μL reaction volume, which contained 12.5 μL Green Taq Mix (Vazyme, Nanjing, China), 1 μL of each forward and reverse primer (10 μM) (Biosune, Shanghai, China), and 1 μL template genomic DNA in amplifier, and were adjusted with distilled deionized water to a total volume of 25 μL.

PCR parameters were as follows: 95 °C for 5 min, followed by 35 cycles of denaturation at 95 °C for 30 s, annealing at a suitable temperature for 30 s, extension at 72 °C for 1 min and a final elongation step at 72 °C for 10 min. Annealing temperature for each gene were 55 °C for ITS, 60 °C for TUB, 52 °C for TEF, 54 °C for CAL
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and 57 °C for HIS. The PCR products were visualised on 1% agarose electrophoresis gel. Sequencing was done bi-directionally, conducted by the Biosune Company Limited (Shanghai, China). Consensus sequences were obtained using MEGA 7.0 (Kumar et al. 2016). All sequences generated in this study were deposited in GenBank (Table 1).

**Phylogenetic analyses**

Novel sequences generated from the sixteen strains in this study, and all reference sequences of *Diaporthe* species downloaded from GenBank, were used for phylogenetic analyses. Alignments of the individual locus were determined using MAFFT v. 7.110 by default settings (Katoh et al. 2017) and manually corrected where necessary. To establish the identity of the isolates at species level, phylogenetic analyses were conducted first individually for each locus and then as combined analyses of five loci (ITS, TUB, TEF, CAL and HIS regions). Phylogenetic analyses were based on maximum likelihood (ML) and Bayesian inference (BI) for the multi-locus analyses. For BI, the best evolutionary model for each partition was determined using MrModeltest v. 2.3 (Nylander 2004) and incorporated into the analyses. ML and BI were run on the CIPRES Science Gateway portal (https://www.phylo.org/) (Miller et al. 2012) using RaxML-HPC2 on XSEDE (8.2.12) (Stamatakis 2014) and MrBayes on XSEDE (3.2.7a) (Huelsenbeck and Ronquist 2001; Ronquist and Huelsenbeck 2003; Ronquist et al. 2012), respectively. For ML analyses the default parameters were used and BI was carried out using the rapid bootstrapping algorithm with the automatic halt option. Bayesian analyses included five parallel runs of 5,000,000 generations, with the stop rule option and a sampling frequency of 500 generations. The burn-in fraction was set to 0.25 and posterior probabilities (PP) were determined from the remaining trees. The resulting trees were plotted using FigTree v. 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree) and edited with Adobe Illustrator CS5.1. New sequences generated in this study were deposited at GenBank (https://www.ncbi.nlm.nih.gov; Table 1) and the alignments and trees were deposited in TreeBASE: S27479 (http://treebase.org/treebase-web/home.html).

**Results**

**Phylogenetic analyses**

Sixteen strains of *Diaporthe* isolated from plant hosts from Yunnan, China, were grown in culture and used for analyses of molecular sequence data. *Diaporthe* spp. were analysed by using multilocus data (ITS, TUB, TEF, CAL and HIS) from 115 isolates of *Diaporthe* spp. and *Diaporthella corylina* (CBS 121124) as the outgroup taxon. A total of 3005 characters including gaps were obtained in the phylogenetic analysis, viz. ITS: 1–656, TUB: 657–1329, TEF: 1330–1860, CAL: 1861–2444,
| Species                  | Voucher | Host/Substrate     | GenBank accession number | Reference                  |
|-------------------------|---------|--------------------|--------------------------|----------------------------|
| Diaporthe asusta       | PSGC 046 | Pycya pyrifolia     | M1626958                 |                           |
| Diaporthe asusta       | PSGC 047*| Pycya pyrifolia     | M1626957                 |                           |
| Diaporthe asusta       | LC1601  | Camellia asamena    | M1626956                 |                           |
| Diaporthe asusta       | LC1601  | Caffe sp.           | M1626954                 |                           |
| Diaporthe asusta       | MAFF 246400 | Amanthus tricolor | M1626952                 |                           |
| Diaporthe asusta       | MAFF 246400 | Amanthus tricolor | M1626953                 |                           |
| Diaporthe asusta       | CBS 111952*| Henicoleum phoxephyllum | M1626951               |                           |
| Diaporthe asusta       | CNucci 201901* | Camellia laeneculata | M1626950                 |                           |
| Diaporthe asusta       | CNucci 201902* | Camellia laeneculata | M1626949                 |                           |
| Diaporthe asusta       | DP0482* | Acetum sp.          | M1626948                 |                           |
| Diaporthe asusta       | CBS 16164* | Aracea cathetu     | M1626947                 |                           |
| Diaporthe asusta       | CBS 535.75* | Citrus sp.          | M1626946                 |                           |
| Diaporthe asusta       | SAUCC194.18* | Persoe americana  | MT1626945                 |                           |
| Diaporthe asusta       | CBS 111979* | Avena angleri      | M1626944                 |                           |
| Diaporthe asusta       | MFLUCC 12-0299a* | On dead leaves      | M1626943                 |                           |
| Diaporthe asusta       | BRIP 54792* | Indigofera australi | M1626942                 |                           |
| Diaporthe asusta       | JZUD 60 | Citrus sinensis     | M1626941                 |                           |
| Diaporthe asusta       | JZUD 61 | Fortunella marigaria | M1626940                 |                           |
| Diaporthe asusta       | JZUD 62 | Citrus grandis      | M1626939                 |                           |
| Diaporthe asusta       | CBS 131813* | Apidocarpus tomentosus | M1626938               |                           |
| Diaporthe asusta       | URM 7486* | Tacina insana       | M1626937                 |                           |
| Diaporthe asusta       | JZB320143 | Urtica dioica       | M1626936                 |                           |
| Diaporthe asusta       | NIBM-ABIP | Carica papaya       | M1626935                 |                           |
| Diaporthe asusta       | CFCC 52563 | Carya ilicinensis  | M1626934                 |                           |
| Diaporthe asusta       | CFCC 52564 | Carya ilicinensis  | M1626933                 |                           |
| Diaporthe asusta       | CFCC 52565 | Cevica chinenis    | M1626932                 |                           |
| Diaporthe asusta       | D. crenulicai | Chrysalidocarpus lutescens | M1626931               |                           |
| Diaporthe asusta       | MFLUCC 17-1023* | Ciceribus sitillus | M1626930                 |                           |
| Diaporthe asusta       | LC0058* | Camellia viscosa    | M1626929                 |                           |
| Diaporthe asusta       | CBS 136.25 | Cucumis sativus    | M1626928                 |                           |
| Diaporthe asusta       | CBS 117499 | Aspalathus linearis | M1626927                 |                           |
| Diaporthe asusta       | CBS 109772 | Coriollus avelana  | M1626926                 |                           |
| Diaporthe asusta       | CBS 444.82 | Eugenia aromatica  | M1626925                 |                           |
| Diaporthe asusta       | BRIP 54781* | Frazinus angustifolius | M1626924               |                           |
| Diaporthe asusta       | CBS 180913 | Pycya pyrifolia     | M1626923                 |                           |
| Diaporthe asusta       | JZBH 20094* | Vitis vinifera     | M1626922                 |                           |

**Table 1.** Species and Genbank accession numbers of DNA sequences used in this study. New species in bold.
| Species                      | Voucher          | Host/Substrate          | ITS         | TUB          | TEF          | GeneBank accession number | CAL         | HIS         | Reference                   |
|------------------------------|------------------|-------------------------|-------------|--------------|--------------|--------------------------|-------------|-------------|------------------------------|
| D. gulyae                    | MF-Ha 17-042*    | Heteranthus anomus      | MK024252    | MK033488     | MK039420     | 666439498               |             |             | Thompson et al. 2011        |
| D. hongkongensis             | CBS 115448*      | Dischma fodiiseta       | KC343119    | KC344087     | KC343845     | KC343361                | KC343603    |             | Gomes et al. 2013           |
| D. huangshanensis            | CNCC 201903      | Camellia oleifera       | MN120729    | MN227010     | MN224670     | –                        |             |             | Zhou and Hou 2019           |
| D. infectoroides             | CBS 133812*      | Schinus titubifolius    | KC343126    | KC344094     | KC343852     | KC343368                | KC343610    |             | Gomes et al. 2013           |
| D. krahiana                  | MFLUCC 17-2481*  | Bruguiera sp.           | MN007101    | MN431495     | MN043215     | –                       |             |             | Dayarathe et al. 2020       |
| D. litchioidea               | BRIJ 5400*       | Litchi chinensis        | JX862531    | JX170925     | JX862539     | –                       |             |             |                              |
| D. limonia                   | CPC 28200*       | Citrus limon            | MF418423    | MF418582     | MF418501     | MF418256                | MF418342    |             | Guarnaccia and Crous 2017   |
| D. luechaniae                | CBS 123121*      | Fometricum vulgare      | KC343136    | KC344104     | KC343862     | KC343378                | KC343620    |             | Phillips and Santos 2009    |
| D. machili                   | SAUCC194.69      | Pometia pinnata         | MT822597    | MT855794     | MT855909     | MT855677                | MT855565    |             | This study                  |
| D. mahorum                   | CAA752*          | Malus domestica         | KY345645    | KY345671     | KY345630     | KY345661                | KY345651    |             | Santos et al. 2017          |
| D. malorum                   | CAA740           | Malus domestica         | KY34662     | KY345670     | KY343562     | KY345660                | KY345650    |             | Santos et al. 2017          |
| D. manihotia                 | CBS 505.76       | Manihot urilurutaxa     | KC343138    | KC344106     | KC343864     | KC343380                | KC343622    |             | Gomes et al. 2013           |
| D. maytenii                  | CBS 133185*      | Maytenus kircikovi      | KC343139    | KC344107     | KC343856     | KC343381                | KC343623    |             | Gomes et al. 2013           |
| D. melitensis                | CPC 27873*       | Citrus limon             | MF418425    | MF418584     | MF418503     | MF418258                | MF418344    |             | Guarnaccia and Crous 2017   |
| D. multidentata              | BRIJ 54846*      | Rupitrum rugosum        | KJ197286    | KJ197267     | KJ197248     | –                       |             |             | Thompson et al. 2015        |
| D. nilintii                  | SAUCC194.27      | Litchi chinensis        | MT822555    | MT855752     | MT855868     | MT855639                | MT855524    |             | This study                  |
| D. nigricanellae             | SAUCC194.45      | Lithocarpus galeber      | MT822573    | MT855770     | MT855886     | MT855654                | MT855542    |             | This study                  |
| D. nigrocardiaceae           | SAUCC194.46      | Lithocarpus galeber      | MT822574    | MT855771     | MT855887     | MT855655                | MT855543    |             | This study                  |
| D. nilintii                  | SAUCC194.48      | Lithocarpus cainilicola  | MT822576    | MT855773     | MT855889     | MT855657                | MT855545    |             | This study                  |
| D. milleitiae                | GUC199167*       | Miletta chinensis       | MK398675    | MK502089     | MK502086     | –                       |             |             | Long et al. 2019            |
| D. musgangulata              | ZJUD 98*         | Citrus grandis           | KJ490635    | KJ490454     | KJ490512     | –                       |             |             | Huang et al. 2015           |
| D. myrsina                   | CBS 129519*      | Musa sp.                | KC343143    | KC344111     | KC343869     | KC343385                | KC343627    |             | Crous et al. 2011           |
| D. neesetii                  | CBS 110499*      | Ambrosia trifida        | KC343145    | KC344113     | KC343871     | KC343387                | KC343629    |             | Silva et al. 2019           |
| D. newii                     | CBS 127270*      | Glyceine max             | KC343156    | KC344124     | KC343882     | KC343398                | KC343640    |             | Santos et al. 2011          |
| D. osmanthi                  | GUC199165*       | Osmantus fragrans       | MK398675    | MK502091     | MK502086     | –                       |             |             | Long et al. 2019            |
| D. oxyceratulii              | SAUCC194.21      | Litchi chinensis        | MT822549    | MT855746     | MT855862     | MT855634                | MT855518    |             | This study                  |
| D. pantanii                  | CBS 133186*      | Maytenus ilicifolia     | KC343164    | KC344132     | KC343890     | KC343406                | KC343648    |             | Gomes et al. 2013           |
| D. persica                   | MFLUCC 17-0607*  | Pandanus sp.            | MG694679    | MG696930     | –            | –                       |             |             | Gomes et al. 2013           |
| D. persicoidea               | MFLUCC 16-0105*  | Prunus persico          | KU557555    | KU557579     | KU557623     | KU557603                |             |             | Dissanyake et al. 2017      |
| Species                        | Voucher       | Host/Substrate       | GenBank accession number | Reference                  |
|-------------------------------|---------------|----------------------|--------------------------|-----------------------------|
| *D. pseudomangiferae*         | CBS 101339*   | Mangifera indica     | KC343458                 | Gomes et al. 2013           |
| *D. pseudophoenicola*         | CBS 462.69*   | Phoenix dactylifera   | KC343456                 | Gomes et al. 2013           |
| *D. pyracanthae*              | MFLUCC 10-0580a* | Piarus pyracanthae  | JQ619887                 | Udawanga et al. 2012       |
| *D. racemosae*                | CAA 448*      | Euclea racemosa      | KY435626, KY435636, KY43567, KY43569 | Santos et al. 2017        |
| *D. salinicola*               | MFLU 7-0553*  | Xylophilus sp.       | MN071196                 | Dayaratne et al. 2020      |
| *D. salinicola*               | MFLU 17-2592  | Xylophilus sp.       | MN071196                 | Dayaratne et al. 2020      |
| *D. scharni*                  | CBS 133181*   | Schinus terebinthifolius | KC343431, KC343439, KC343458 | Gomes et al. 2013           |
| *D. schloeni*                 | MFLU 15-2609  | Schinus terebinthifolius | KY964229, KY964112, KY964185 | Disnayake et al. 2017     |
| *D. serpens*                  | CFCC 51636*   | Senna bicapsularis    | KY203724, KY223891, KY223885, KY223887 | Yang et al. 2017       |
| *D. spinosa*                  | BRIP 55665*   | Helianthus annuus    | KJ197274, KJ197267, KJ197274, KJ197274 | Thompson et al. 2015 |
| *D. spiralis*                 | PSGS 383*     | Pyrus pyrifolia      | MK666849, MK669123, MK669112, MK669112, MK669112, MK669112 | Guo et al. 2020        |
| *D. stenosticta*              | CBS 193.36*   | Cocos nucifera       | FJ899448, JX275421, GG250324, JX197415 | Santos et al. 2018, Udawanga et al. 2018 |
| *D. subordinaria*             | CBS 101711    | *Plantago lanceolata* | KC343213, KC343418, KC343418, KC343418, KC343418, KC343418 | Gomes et al. 2013           |
| *D. tanicola*                 | CBS 466.90    | *Plantago lanceolata* | KC343214, KC343418, KC343418, KC343418, KC343418, KC343418 | Gomes et al. 2013           |
| *D. tachonousi*               | PSGG485       | Prunus persica       | MK66689, MK669122, MK669127, MK669128, MK669128, MK669128 | Disnayake et al. 2017     |
| *D. tecktonigena*             | MFLUC 12-0767* | *Tectona grandis*  | KU712429, KU743976, KU749371, KU749358 | Doki et al. 2016       |
| *D. terebinthifolii*          | CBS 131810*   | *Schinus terebinthifolius* | KC343216, KC343418, KC343418, KC343418, KC343418, KC343418 | Gomes et al. 2013           |
| *D. uniflora*                 | LG662*        | Unknown host         | KX986798, KX999230, KX999190 | Gao et al. 2017          |
| *D. vulgaris*                 | LG8110*       | Unknown host         | KY491545, KY491565, KY491565, KY491585 | Gao et al. 2017          |
| *D. xanthomelas*              | BRIP 57887*   | Psidium guajava      | KR396126, KR396128, KR396129 | Gao et al. 2015           |
| *D. xinjiangensis*            | JZBH 320071   | Vitis vinifera      | MK341550, MK500112, MK500107, MK500119 | Manawasighe et al. 2019 |
| *D. xinjiangensis*            | JZBH 320072   | Vitis vinifera      | MK341550, MK500112, MK500108, MK500107, MK500108 | Manawasighe et al. 2019 |
| *D. xinguanensis*             | LG707*        | Camellia sinensis    | KX986783, KX999216, KX999175, KX999255 | Gao et al. 2017          |
| *Diaporthe corylina*          | CBS 121124    | Corylus sp.          | KC343004, KC343397, KC343370, KC343426, KC343488 | Gomes et al. 2013           |

Isolates marked with **star** are ex-type or ex-epitype strains.
Figure 1. Phylogram of *Diaporthe* spp. based on combined sequence data of ITS, TUB, TEF, CAL and HIS genes. The ML and BI bootstrap support values above 50% and 0.90 BYPP are shown at the first and second position, respectively. Strains marked with “*” are ex-type or ex-epitype. Codes referring to strains from the current study are written in red. Some branches were shortened to fit them to the page as indicated by two diagonal lines with the number of times a branch was shortened indicated.
Five new species and three known species of *Diaportha* from Yunnan, China

**Figure 1.** Continued.
Figure 1. Continued.

HIS: 2445–3005. Of these characters, 1349 were constant, 453 were variable and parsimony-uninformative, and 1203 were parsimony-informative. For the BI and ML analyses, the substitution model GTR+I+G for ITS, TUB, TEF and HIS, HKY+I+G for and CAL were selected and incorporated into the analyses. The ML tree topology confirmed the tree topologies obtained from the BI analyses, and therefore, only the ML tree is presented (Fig. 1).

ML bootstrap support values (≥ 50%) and Bayesian posterior probability (≥ 0.90) are shown as first and second position above nodes, respectively. Based on the five-locus phylogeny and morphology, nine isolates were assigned to five species, including *Diaporthe arecae* (1), *D. hongkongensis* (2), *D. middletonii* (4), *D. osmanthi* (1) and *D. pandanicola* (1), whereas seven isolates formed distinct well supported clades, which refer to novel species named *D. chrysalidocarpi* (2), *D. machili* (2) and *D. pometiae* (3), respectively.
Taxonomy

**Diaporthe arecae** (H.C. Srivast., Zakia & Govindar.) R.R. Gomes, Glienke & Crous, *Persoonia* 31: 16. (2013)

Figure 2

**Subramanella arecae** H.C. Srivast., Zakia & Govindar., in Srivastava, Banu and Govindarajan (1962). Basionym.

**Description.** Asexual morph: Conidiomata pycnidial, several pycnidia grouped together, globose, black, erumpent, exuding creamy to yellowish conidial droplets from ostioles. Conidiophores hyaline, septate, branched, cylindrical, straight to sinuous, 25.0–32.0 × 1.4–2.5 μm. Conidiogenous cells 10.5–20.7 × 1.4–2.0 μm, phialidic, cylindrical, swollen at base, tapering towards apex, slightly curved. Alpha conidia hyaline, smooth, aseptate, ellipsoidal, guttulate, apex subobtuse, base subtruncate, 7.5–10.0 × 1.8–3.0 μm (mean = 8.2 × 2.4 μm, n = 20). Beta conidia hyaline, aseptate, filiform, slightly curved, tapering towards base, 18.5–26.5 × 1.0–1.8 μm (mean = 24.3 × 1.4 μm, n = 20). Gamma conidia not observed. Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 11.2–13.3 mm diam/day. Aerial mycelium white, cottony, feathery, abundant in center, sparse in margin, white on surface, reverse yellowish to tan.

**Specimen examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of *Persea americana* (Lauraceae). 19 April 2019, S.T. Huang, HSAUP194.18, living culture SAUCC194.18.

**Notes.** *Diaporthe arecae* (CBS 161.64) was originally described as *Subramanella arecae* on fruit of *Areca catechu* in India (Srivastava et al. 1962) and placed in *Diaporthe* by Gomes et al. (2013). The *Diaporthe* isolate from fruits of *Citrus* sp. (CBS 535.75) in Suriname was also placed in *D. arecae* by Gomes et al. (2013). In the present study, strain (SAUCC194.18) from symptomatic leaves of *Persea americana* was congruent with *D. arecae* based on morphology and DNA sequences data (Fig. 1). We therefore consider the isolated strain as *D. arecae*.

**Diaporthe chrysalidocarpi** S.T. Huang, J.W. Xia, W.X. Sun, & X.G. Zhang, sp. nov.

MycoBank No: 837812

Figure 3

**Etymology.** Named after the host genus on which it was collected, *Chrysalidocarpus lutescens*.

**Diagnosis.** *Diaporthe chrysalidocarpi* can be distinguished from the phylogenetically most closely related species *D. spinosa* by longer beta conidia (28.0–32.5 × 1.2–1.6 vs. 18.5–30.5 × 1.0–1.5 μm), and from other species *D. fulvicolor* by the types of conidia (*D. chrysalidocarpi* produces only beta conidia, while *D. fulvicolor* produces
Figure 2. *Diaporthe arecae* (SAUCC194.18) a infected leaf of *Persea americana* b, c surface and reverse of a colony after 15 days on PDA d conidiomata e–g conidiophores and conidiogenous cells h beta conidia i alpha conidia j alpha conidia and beta conidia. Scale bars: 10 μm (e–j).

only alpha conidia) and several loci (25/491 in the ITS region, 18/471 TUB, 4/298 TEF, 28/458 CAL and 13/441 HIS).

**Type.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of *Chrysalidocarpus lutescens* (Palmae). 19 April 2019, S.T. Huang, HSAUP194.35 holotype, ex-type living culture SAUCC194.35.

**Description.** Asexual morph: Leaf spots irregular, pale brown in center, brown to tan at margin. Conidiomata pycnidial, scattered or aggregated, black, erumpent, raising above surface of culture medium, subglobose, exuding white or yellowish creamy conidial droplets from central ostioles after 30 days in light at 25 °C; pycnidial wall
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Consists of black to dark brown, thin-walled cells. Conidiophores 27.5–35.0 × 1.4–2.0 μm, hyaline, slightly branched, swelling at base, subcylindrical, septate, smooth, straight or curved. Conidiogenous cells 10.5–23.0 × 1.4–1.8 μm, phialidic, cylindrical, terminal, straight to sinuous, tapering towards apex. Beta conidia 28.0–32.5 × 1.2–1.6 μm (mean = 30.3 × 1.3 μm, n = 20), filiform, hyaline, straight or slightly curved, aseptate, base subtruncate, tapering towards the base. Alpha conidia and gamma conidia not observed. Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 13.3–15.2 mm diam/day, initially white, becoming greyish, reverse pale brown, with concentric rings of dense, sparse hyphae, irregular margin, fluffy aerial mycelium at center, pycnidia forming after 15 days.

*Figure 3. Diaporthe chrysalidocarpi* (SAUCC194.35) **a** diseased leaf of *Chrysalidocarpus lutescens* **b, c** surface and reverse of a colony after 15 days on PDA **d, e** conidiomata **f, g** conidiophores and conidiogenous cells **h, i** beta conidia. Scale bars: 10 μm (**f–i**).
**Additional specimen examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of *Chrysalidocarpus lutescens* (Palmae). 19 April 2019, S.T. Huang, HSAUP194.33 paratype; living culture SAUCC194.33.

**Notes.** Phylogenetic analysis of a combined five gene showed that *D. chrysalidocarpi* formed an independent clade (Fig. 1) and is phylogenetically distinct from *D. spinosa* and *D. fulvicolor*. This species can be distinguished from *D. spinosa* by 61 different nucleotides in the concatenated alignment (13/492 in the ITS region, 17/471 TUB, 4/298 TEF, 17/458 CAL and 10/441 HIS), and *D. fulvicolor* by 88 nucleotides (25/491 in the ITS region, 18/471 TUB, 4/298 TEF, 28/458 CAL and 13/441 HIS). Morphologically, *D. chrysalidocarpi* differs from *D. spinosa* in having longer beta conidia (28.0–32.5 × 1.2–1.6 vs. 18.5–30.5 × 1.0–1.5 μm) (Guo et al. 2020). Furthermore, *Diaporthe chrysalidocarpi* produces only beta conidia, while *D. spinosa* produces alpha conidia and beta conidia and *D. fulvicolor* produces only alpha conidia (Guo et al. 2020). Therefore, we establish this fungus as a novel species.

*Diaporthe hongkongensis* R.R. Gomes, Glienke, Crous, *Persoonia* 31: 23. (2013)

**Figure 4**

**Description.** Asexual morph: Conidiomata pycnidial, subglobose or globose, solitary, black, erumpent, coated with white hyphae, thick-walled, exuding creamy conidial droplets from central ostioles. Conidiophores hyaline, smooth, septate, unbranched, densely aggregated, cylindrical or clavate, straight to sinuous, swollen at base, tapering towards apex, 32.0–42.0 × 2.0–2.9 μm. Conidiogenous cells 20.0–24.2 × 1.3–2.3 μm, phialidic, cylindrical, terminal, slightly tapering towards apex. Alpha conidia, hyaline, smooth, aseptate, ellipsoidal or oval, 0–2 guttulate, apex subobtuse, base subtruncate, 5.5–7.0 × 2.0–2.5 μm (mean = 6.2 × 2.2 μm, n = 20). Beta conidia hyaline, aseptate, filiform, hamate, tapering towards both ends, mostly J-shaped, 21.5–27.0 × 1.4–1.8 μm (mean = 25.6 × 1.3 μm, n = 20). Gamma conidia not observed. Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 19.0–21.5 mm diam/day, cottony, radial with abundant aerial mycelium, sparse at margin, with an obvious pale brown concentric ring of dense hyphae, white to grayish on surface with age, white to pale brown on the reverse side.

**Specimens examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, 19 April 2019, S.T. Huang. On diseased leaves of *Millettia reticulata* (Fabaceae) HSAUP194.81, living culture SAUCC194.81; on diseased leaves of *Camellia sinensis* (Theaceae) HSAUP194.87, living culture SAUCC194.87.

**Notes.** In the present study, two strains (SAUCC194.81 and SAUCC194.87) from symptomatic leaves of *Millettia reticulata* and *Camellia sinensis* were similar to *Diaporthe hongkongensis* (CGMCC 3.17102) (Gomes et al. 2013) and *D. salinicola* (MFLU 18-0553) (Dayarathne et al. 2020) based on DNA sequences data (Fig. 1). Morphologically, our strains were similar to *Diaporthe hongkongensis*, which was originally described with an asexual morph on fruits of *Dichroa febrifuga* in China,
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**Figure 4.** *Diaporthe hongkongensis* (SAUCC194.87) a diseased leaf of *Camellia sinensis* b, c surface and reverse of colony after 15 days on PDA d conidiomata e–g conidiophores and conidiogenous cells h beta conidia i alpha conidia. Scale bars: 10 μm (e–i).

but the asexual morph of *D. salinicola* was undetermined. We therefore identify our strains as *D. hongkongensis*.

**Diaporthe machili** S.T. Huang, J.W. Xia, W.X. Sun, & X.G. Zhang, sp. nov.
MycoBank No: 837814
Figure 5

**Etymology.** Named after the host genus on which it was collected, *Machilus pingii*.

**Diagnosis.** *Diaporthe machili* differs from *D. caryae* and *D. sackstonii* in the types of conidia (*D. machili* only produces beta conidia, while *D. caryae* produces alpha
conidia and beta conidia, and *D. sackstonii* only produces alpha conidia), and from *D. caryae* in longer beta conidia (29.0–39.0 × 1.3–1.5 vs. 15.5–34.0 × 1.1–1.4 μm).

**Type.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of *Machilus pingii* (Lauraceae). 19 April 2019, S.T. Huang, HSAUP194.111 holotype, ex-holotype living culture SAUCC194.111.

**Description.** Asexual morph: Conidiomata pycnidial, aggregated, black, erumpent, subglobose to globose, exuding creamy conidial droplets from central ostioles after 30
days in light at 25 °C. Conidiophores 7.0–11.4 × 1.8–2.8 μm, hyaline, unbranched, densely aggregated, mostly ampulliform, cylindrical, guttulate, septate, straight or slightly curved, swelling at base, tapering towards apex. Beta conidia 29.0–39.0 × 1.3–1.5 μm (mean = 32.5 × 1.4 μm, n = 20), filiform, hyaline, aseptate, mostly curved, J-shaped, swelling in middle, tapering towards both ends. Alpha and gamma conidia not observed. Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 16.3–17.5 mm diam/day, aerial mycelium abundant, white on surface, reverse white to pale yellow, with an obvious concentric zonation, pycnidia forming after 15 days.

**Additional specimen examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of *Pometia pinnata* (Sapindaceae). 19 April 2019, S.T. Huang, HSAUP194. 69 paratype; living culture SAUCC194. 69.

**Notes.** In the phylogenetic tree, *Diaporthe machili* forms an independent clade and is phylogenetically distinct from *D. caryae* and *D. sackstonii* (Fig. 1). *Diaporthe machili* can be distinguished from *D. caryae* in ITS, TUB, TEF, CAL and HIS loci by 67 nucleotide differences in concatenated alignment (5/459 in ITS, 10/416 in TUB, 15/334 in TEF, 7/454 in CAL and 30/455 in HIS), and from *D. sackstonii* in ITS, TUB and TEF loci by 58 nucleotide differences (12/559 in ITS, 23/486 in TUB and 23/348 in TEF). Moreover, *Diaporthe machili* differs from *D. caryae* in having longer beta conidia (29.0–39.0 × 1.3–1.5 vs. 15.5–34.0 × 1.1–1.4 μm). *Diaporthe machili* only produces beta conidia, while *D. caryae* produces alpha conidia and beta conidia, and *D. sackstonii* only produces alpha conidia (Thompson et al. 2015; Yang et al. 2018b).

**Diaporthe middletonii** R.G. Shivas, L. Morin, S.M. Thomps. & Y.P. Tan, *Persoonia* **35**: 45. (2015)

Figure 6

**Description.** Asexual morph: Leaf spots discoid to irregular. Conidiomata pycnidial, scattered or aggregated in groups of 3–5 pycnidia, globose, black, erumpent, coated with white to greyish hyphae, thick-walled, exuding creamy translucent conidial droplets from central ostioles. Conidiophores hyaline, smooth, septate, unbranched, densely aggregated, cylindrical, straight to sinuous, tapering towards apex, 10.0–14.0 × 1.3–2.3 μm. Conidiogenous cells 5.0–9.5 × 1.3–1.7 μm, phialidic, cylindrical, terminal, slightly tapering towards apex. Alpha conidia hyaline, smooth, aseptate, biguttulate, ellipsoidal, oval, apex subobtuse, base subtruncate, 5.5–7.0 × 2.5–3.2 μm (mean = 6.3 × 2.8 μm, n = 20). Beta conidia hyaline, aseptate, filiform, mostly curved by 90–180°, tapering towards both ends, 26.0–36.5 × 1.0–1.6 μm (mean = 21.5 × 1.2 μm, n = 20). Gamma conidia not observed. Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 22.5–24.0 mm diam/day, fluffy with abundant aerial mycelium, margin fimbriate, white on surface, white to pale yellow on reverse.
Figure 6. *Diaporthe middletonii* (SAUCC194.46) a infected leaf of *Lithocarpus glaber* b, c surface and reverse of colony after 15 days on PDA d, e conidiomata f–i conidiophores and conidiogenous cells j beta conidia k, l alpha conidia and beta conidia. Scale bars: 10 μm (f–l).

**Specimens examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, 19 April 2019, S.T. Huang. On diseased leaves of *Litchi chinensis* (Sapindaceae), HSAUP194.27, living culture SAUCC194.27; on diseased leaves of *Lithocarpus glaber* (Fagaceae), HSAUP194.45, living culture SAUCC194.45; on diseased leaves of *Lithocarpus glaber* (Fagaceae), 19 April 2019, S.T. Huang, HSAUP194.46, living culture SAUCC194.46; on diseased leaves of *Lithocarpus craibianus* (Fagaceae), HSAUP194.48, living culture SAUCC194.48.

**Notes.** *Diaporthe middletonii* was originally described from the stem of *Rapistrum rugosum* (BRIP 54884e) (Brassicaceae) and *Chrysanthemoides monilifera* subsp. *rotundata* (BRIP 57329) (Asteraceae) in Australia (Thompson et al. 2015). In the present study, four strains (SAUCC194.27, SAUCC194.45, SAUCC194.46 and SAUCC194.48) are closely related to *D. middletonii* in the combined phylogenetic tree (Fig. 1). The differences between nucleotides in the concatenated alignment (17/565 in ITS, 9/494 in TUB and 10/340 in TEF) were minor. Morphologically, our strains were similar to *D. middletonii* by slightly shorter and wider alpha conidia (5.0–7.0 ×
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2.5–3.2 vs. 6.0–7.5 × 2.0–2.5 μm), and longer beta conidia (26.0–36.5 × 1.0–1.6 vs. 20.0–35.0 × 1.0–1.5 μm) (Thompson et al. 2015). We therefore identify our strains as *Diaporthe middletonii*.

**Diaporthe osmanthi** H. Long, K.D. Hyde, & Yong Wang bis, MycoKeys 57: 120. (2019)

**Figure 7**

**Description.** Conidiomata pycnidial, globose, 5–10 pycnidia grouped together, dark brown to black, exuding creamy to yellowish conidial droplets from central ostioles. Conidiophores hyaline, smooth, densely aggregated, branched, cylindric-clavate, 20.5–32.0 × 1.8–2.4 μm. Conidiogenous cells phialidic, hyaline, terminal, cylindrical, straight, 14.0–20.5 × 1.5–2.0 μm, tapered towards apex. Alpha conidia hyaline, aseptate, fusiform, tapering towards both ends, guttulate, 7.3–9.3 × 1.8–2.3 μm (mean = 8.5 × 2.0 μm, n = 20). Beta conidia hyaline, aseptate, filiform, curved, 22.0–28.5 × 1.0–2.0 μm (mean = 27.2 × 1.3 μm, n = 20). Gamma conidia not observed. Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 12.0–13.5 mm diam/day, cottony with abundant aerial mycelium, sparse at margin. With several concentric rings of dense hyphae, white on surface, white to pale brown on reverse.

**Specimen examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, 19 April 2019, S.T. Huang. On diseased leaves of *Litchi chinensis* (Sapindaceae) HSAUP194.21, living culture SAUCC194.21.

**Notes.** *Diaporthe osmanthi* was originally described from the leaves of *Osmanthus fragrans* (Oleaceae) in Guangxi province, China (Long et al. 2019). In the present study, phylogenetic analyses (Fig. 1) indicated that the strain SAUCC194.21 is closely related to *Diaporthe osmanthi* and *D. podocarpi-macrophylli* (Gao et al. 2017). Morphological comparison indicated that this strain was most similar to *D. osmanthi* by the size of alpha conidia and beta conidia. We therefore identify this strain as belonging to *D. osmanthi*.

**Diaporthe pandanicola** Tibpromma & K.D. Hyde, MycoKeys 33: 44 (2018)

**Figure 8**

**Description.** Asexual morph: Conidiomata pycnidial, 3–5 pycnidia grouped together, superficial to embedded on PDA, erumpent, thin-walled, dark brown to black, globose or subglobose, exuding white creamy conidial mass from ostioles. Conidiophores hyaline, aseptate, cylindrical, smooth, straight to sinuous, unbranched, aggregated, 17.0–26.5 × 2.0–3.0 μm. Conidiogenous cells phialidic, cylindrical, terminal, 10.0–20.0 × 1.5–1.8 μm. Alpha conidia hyaline, smooth, aseptate, ellipsoidal, eguttulate, apex subobtuse, base subtruncate, 6.5–9.0 × 1.8–2.5 μm (mean = 7.5 × 2.0 μm, n = 20). Beta conidia hyaline, aseptate, filiform, curved, tapering towards apex, base truncate,
26.0–32.8 × 1.0–1.6 μm (mean = 29.0 × 1.3 μm, n = 20). Gamma conidia infrequent, aseptate, smooth, straight, hyaline, 12.5–14.5 × 1.3–1.8 μm (mean = 13.5 × 1.6 μm, n = 6). Sexual morph not observed.

**Culture characteristics.** Cultures incubated on PDA at 25 °C in darkness, growth rate 12.8–15.0 mm diam/day, flat, cottony in centre, with aerial mycelium sparse toward margin, white on surface, white to pale yellow on reverse.

**Specimen examined.** China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of *Millettia reticulata* (Fabaceae). 19 April 2019, S.T. Huang, HSAUP194.82, living culture SAUCC194.82.
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**Figure 8.** *Diaporthe pandanicola* (SAUCC194.82) **a** infected leaf of *Millettia reticulata* **b, c** surface and reverse of colony after 15 days on PDA **d** conidiomata **e–g** conidiophores and conidiogenous cells **h** beta conidia **i** alpha conidia and gamma conidia **j** alpha conidia, beta conidia and gamma conidia. Scale bars: 10 μm (**e–j**).

**Notes.** *Diaporthe pandanicola* was originally described by Tibpromma et al. (2018) on healthy leaves of *Pandanus* sp. (Pandanaceae) as an endophytic fungus. Our strain (SAUCC194.82) is closely related to *Diaporthe pandanicola* based on phylogenetic analyses (Fig. 1). The differences of nucleotides in the concatenated alignment (19/533 in the ITS region and 11/351 in the TUB region) are less than 3%. Morphologically, our strain produces alpha conidia, beta conidia and gamma conidia, while *Diaporthe pandanicola* did not sporulate. We therefore identify our strains as *Diaporthe pandanicola*.
Diaporthe pometiae S.T. Huang, J.W. Xia, W.X. Sun, & X.G. Zhang, sp. nov.
MycoBank No: 837815

Figure 9

Etymology. Named after the host genus on which it was collected, Pometia pinnata.

Diagnosis. Diaporthe pometiae is similar to D. biconispora but differs in having smaller alpha conidia (5.7–8.3 × 2.2–3.0 vs. 6.0–10.5 × 2–3.5 μm) and types of conidia (D. pometiae produces beta conidia unlike D. biconispora).

Type. China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, on diseased leaves of Pometia pinnata (Sapindaceae). 19 April 2019, S.T. Huang, HSAUP194.72 holotype, ex-type living culture SAUCC194.72.

Figure 9. Diaporthe pometiae (SAUCC194.72) a infected leaf of Pometia pinnata b, c surface and reverse of colony after 15 days on PDA d conidiomata e, f conidiophores and conidiogenous cells g beta conidia h alpha conidia and beta conidia. Scale bars: 10 μm (e–h).
Description. Asexual morph: Leaf spots subcircular, fawn to dark brown. Conidiomata pycnidial, subglobose to globose, aggregated in groups, black, coated with white hyphae, thick-walled, exuding creamy droplets from ostioles. Conidiophores hyaline, smooth, slightly septate, branched, densely aggregated, cylindric-clavate, straight to slightly sinuous, 22.5–32.5 × 1.0–2.0 μm. Conidiogenous cells 15.0–22.5 × 1.0–1.5 μm, phialidic, cylindrical, multi-guttulate, terminal, tapering towards apex. Alpha conidia abundant in culture, 2–4 guttulate, hyaline, smooth, aseptate, ellipsoidal to oblong ellipsoidal, with both ends obtuse, 5.7–8.3 × 2.2–3.0 μm (mean = 6.7 × 3.1 μm, n = 20). Beta conidia, hyaline, aseptate, filiform, multi-guttulate, slightly curved, tapering towards to apex, 27.8–34.5 × 1.0–1.7 μm (mean = 21.7 × 1.4 μm, n = 20). Gamma conidia not observed. Sexual morph not observed.

Culture characteristics. Cultures incubated on PDA at 25 °C in darkness, growth rate 11.5–13.0 mm diam/day, cottony with abundant aerial mycelium, with a concentric zonation, white on surface, white to grayish on reverse.

Additional specimens examined. China, Yunnan Province: Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, 19 April 2019, S.T. Huang. On diseased leaves of *Persea americana* (Lauraceae), HSAUP194.19 paratype, ex-paratype culture SAUCC194.19; on diseased leaves of *Heliconia metallica* (Musaceae), HSAUP194.73 paratype, ex-paratype culture SAUCC194.73.

Notes. *Diaporthe pometiae* is introduced based on the multi-locus phylogenetic analysis, with three isolates clustering separately in a well-supported clade (ML/BI = 100/1). *Diaporthe pometiae* is most closely related to *D. biconispora*, but distinguished based on ITS, TUB, TEF and HIS loci by 74 nucleotide differences in the concatenated alignment, in which 2/492 are distinct in the ITS region, 8/353 in the TUB region, 49/370 in the TEF region and 15/471 in the HIS region. Morphologically, *Diaporthe pometiae* differs from *D. biconispora* in its smaller alpha conidia (5.7–8.3 × 2.2–3.0 vs. 6.0–10.5 × 2–3.5 μm). Furthermore, *Diaporthe pometiae* produces beta conidia unlike *D. biconispora* (Huang et al. 2015).

Discussion

The Yunnan Province in southeastern China has a unique geography where three climatic regions meet: the eastern Asia monsoon region, the Tibetan plateau region, and the tropical monsoon region of southern Asia and Indo-China. The environment is conducive to growth of unusual microbial species. Species diversity in Yunnan Province is high compared to other parts of China.

Previously, species identification of *Diaporthe* relied on the assumption of host-specificity, leading to the proliferation of names. The morphological characters of *Diaporthe* could be changeable, as most taxa in culture do not produce all spore states of the asexual (alpha, beta and gamma conidia) or the sexual morph (Gomes et al. 2013). Based on a polyphasic approach and morphology, more than one species of
Diaporthe can colonize a single host, while one species can be associated with several hosts (Gomes et al. 2013; Gao et al. 2017; Guarnaccia and Crous 2017; Guarnaccia et al. 2018; Guo et al. 2020). These studies revealed a high diversity of Diaporthe species from different hosts. Our study supports this phenomenon. For example, Diaporthe arecae (SAUCC194.18) and D. pometiae (SAUCC194.19) were collected from Persea americana; in addition, isolates of D. middletonii were obtained from three hosts (Litchi chinensis, Lithocarpus craibianus, L. glaber). As for host specificity, in our study, four species of Diaporthe, D. machili (SAUCC194.69), D. middletonii (SAUCC194.27), D. osmanthi (SAUCC194.21), and D. pometiae (SAUCC194.72) were isolated from Litchi chinensis and Pometia pinnata belong to the Sapindaceae, and D. litchiicola also was reported from Litchi chinensis in Queensland (Tan et al. 2013); however, D. machili (SAUCC194.111) also was isolated from Machilus pingii (Lauraceae), D. middletonii (SAUCC194.45) from Lithocarpus glaber (Fagaceae), D. osmanthi (GUCC 9165) from leaves of Osmanthus fragrans (Oleaceae) (Long et al. 2019), and D. pometiae (SAUCC194.19 and SAUCC194.73) from Persea americana (Lauraceae) and Heliconia metallic (Musaceae). These results provide evidence that many species are able to colonise diverse hosts and several different species could co-occur on the same host. It seems obvious that specificity does not occur at the family level.

For the current study, sixteen strains isolated from ten host genera represented three new species and five known species, based on morphological characters and phylogenetic analyses of the five combined loci (ITS, TUB, TEF, CAL and HIS). The descriptions and molecular data for species of Diaporthe represent an important resource for plant pathologists, plant quarantine officials and taxonomists.

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