Lasiodiplodia syzygii sp. nov.
(Botryosphaeriaceae) causing post-harvest water-soaked brown lesions on Syzygium samarangense in Chiang Rai, Thailand

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Abstract

Background

Syzygium samarangense (Wax apple) is an important tropical fruit tree with high economic and nutrient value and is widely planted in the tropics or subtropics of Asia. Post-harvest water-soaked brown lesions were observed on mature fruits of ornamental wax apples in Chiang Rai Province, Thailand. A fungus with morphological characters, similar to Lasiodiplodia, was consistently isolated from symptomatic fruits. Phylogenetic analyses, based on ITS, LSU, TEF1-a and tub2, revealed that our isolates were closely related to, but phylogenetically distinct from, Lasiodiplodia rubropurpurea.
New information

Morphological comparisons indicated that pycnidia and conidiogenous cells of our strains were significantly larger than *L. rubropurpurea*. Comparisons of base-pair differences in the four loci confirmed that the species from wax apple was distinct from *L. rubropurpurea* and a new species, *L. syzygii* sp. nov., is introduced to accommodate it. Pathogenicity tests confirmed the newly-introduced species as the pathogen of this post-harvest water-soaked brown lesion disease on wax apples.

Keywords

Botryosphaeriaceae, fruit disease, new pathogen, wax apple

Introduction

Wax apple [*Syzygium samarangense* (Blume) Merrill and Perry] belongs to the *Myrtaceae* and was naturalised in the Philippines thousands of years ago (Lim 2012, Shen et al. 2012). As a kind of juicy tropical fruit like watermelon with economic importance, it has been commonly and widely cultivated in many Asian countries (Nesa et al. 2014). Every part of *S. samarangense* also has potential medicinal values (Shen et al. 2012).

Due to the fruit characteristics, such as thin peel and tender pulp with high respiratory intensity, wax apples are prone to damage by pathogens and cannot be stored for a long time (Yang et al. 2009). This causes a significant post-harvest loss. Many studies suggest that wax apple is mainly threatened by fungal diseases. For example, a new fruit rot of wax apple caused by *Phytophthora palmivora* was reported in southern Taiwan during the rainy periods in 1982 (Lin et al. 1984). Yang et al. (2009) and Che et al. (2015) reported *Lasiodiplodia theobromae* as the causal agent of black spot disease on harvested wax apple fruits. *Pestalotiopsis samarangensis* was isolated from the fruit rot in wax apples from markets in Thailand (Maharachchikumbura et al. 2013). *Chrysoporthe deuterocubensis* caused cankers on wax apple and branches in Taiwan (Fan et al. 2013).

The present study reports a new post-harvest water-soaked brown lesion disease on wax apples caused by *Lasiodiplodia* sp. in Chiang Rai, Thailand. Morphological and multi-locus phylogenetic analyses revealed that our strain represented a novel species. A pathogenicity test on fruits confirmed the pathogenic relationship between *L. syzygii* and *Syzygium samarangense*. 
Materials and methods

Sample collection, isolation and morphology

Rotten wax apple fruits were occasionally collected from a food market near Mae Fah Luang University in Chiang Rai, Thailand. On the third day after the wax apple fruits were collected, it was observed that there were conidiomata bulges on the surface of the fruit, white hyphae and the fruit turned black, rotted and had cytoplasmic extravasation. Diseased samples were conserved in self-sealing bags and then taken back to the laboratory and photographed. Before isolation, diseased fruits were surface disinfected with 70% ethanol for 30 s, 1% sodium hypochlorite (NaClO) for 1 min and repeatedly twice rinsed in sterile distilled water for 30 s. Pure cultures were obtained by single-conidium isolation following a modified method outlined by Chomnunti et al. (2011) and Maharachchikumbura et al. (2013). The morphology of fungal colonies was recorded following the method of Hu et al. (2007). Fungal mycelium and spores were observed under a light microscope and photographed. The holotype specimen is deposited in the Herbarium of the Department of Plant Pathology, Agricultural College, Guizhou University (HGUP). The ex-type and isotype cultures are deposited in the Culture Collection at the Department of Plant Pathology, Agriculture College, Guizhou University, P.R. China (GUCC) and the Mae Fah Luang University Culture Collection (MFLUCC) in Thailand.

DNA extraction, PCR reaction and sequencing

Fungal cultures were grown on PDA at 28°C. When colonies nearly covered the entire Petri dish (90 mm diam.), fresh mycelia were scraped from the agar surface with sterilised scalpels. Genomic DNA was extracted using a BIOMIGA Fungus Genomic DNA Extraction Kit (GD2416) following the manufacturer’s protocol. DNA amplification was performed in a 25 μl reaction volume following Liang et al. (2018). Primers ITS1 and ITS4 (White et al. 1990) were used to amplify the internal transcribed spacer regions and intervening 5.8S rRNA region (ITS) and LR0R and LR5 for 28S rRNA (LSU) region (Vilgalys and Hester 1990, Rehner and Samuels 1994). Two protein-coding gene fragments, the β-tubulin (tub2) and translation elongation factor 1-alpha (TEF1-a) were amplified with primer pairs BT2A/BT2B (Glass and Donaldson 1995, O'Donnell and Cigelnik 1997) and EF1-688F/EF1-986R, respectively (Carbone and Kohn 1999, Alves et al. 2008). Purification and sequencing of the PCR amplicons were done by SinoGenoMax, Beijing. The DNA sequences are deposited in the GenBank and their accession numbers are provided in Table 1. The DNA base differences of the four loci amongst our strains and ex-type or representative strains of relative taxa are shown (Table 2).
| Species                        | Isolate no. | GenBank no. |
|-------------------------------|-------------|-------------|
|                               |             | ITS  | LSU | tef 1 | tub2 |
| *Lasiodiplodia americana*     | CFCC50065T  | KP217059 | MF410052 | KP217067 | KP217075 |
| *L. avicenniae*               | CMW 414673T | KP860835 | –   | KP860680 | KP860758 |
| *L. brasiilense*              | CMM 4015T   | JX464063 | –   | JX464049 | –     |
| *L. brassilense*              | CMW 35884   | KS887094 | –   | KS886972 | KS887466 |
| *L. bruguiereae*              | CMW 41470T  | KP860833 | –   | KP860678 | KP860756 |
| *L. caatinguensis*            | CMM 1325T   | KT154760 | –   | KT008006 | KT154767 |
| *L. caatinguensis*            | IBL 40      | KT154762 | –   | KT154755 | KT154769 |
| *L. chinenis*                 | CMW 35884   | JX464063 | –   | JX464049 | –     |
| *L. citricola*                | IRAN 1522C  | GU945354 | –   | GU945340 | KS887505 |
| *L. crassispora*              | WAC12533T   | DQ103550 | DQ377901 | EU673303 | KS887506 |
| *L. euphorbicola*             | CMM 3609T   | KS234543 | –   | KS234689 | KS254926 |
| *L. exigua*                   | CBS 137785T | KJ638317 | –   | KJ638336 | KS887509 |
| *L. gilanensis*               | IRAN 1523T  | GU945351 | –   | GU945342 | KS887511 |
| *L. gilarensis*               | CMM 4564T   | KT250949 | –   | KT250950 | –     |
| *L. gilarensis*               | IRAN 1500C  | GU945355 | –   | GU945343 | KS887515 |
| *L. hyalina*                  | CMM 4564T   | KT250949 | –   | KT250950 | –     |
| *L. indica*                   | IBL 01      | KJ376151 | –   | KJ376152 | –     |
| *L. iranensis*                | IRAN 1520C  | GU945348 | –   | GU945336 | KS887516 |
| *L. laeliocattleyae*          | CBS 167.28T | KU507487 | DQ377892 | KU507454 | –     |
| *L. lignicola*                | CBS 134112  | JX646797 | JX646814 | KS887003 | JX646845 |
| *L. macrospora*               | CMM 3833T   | KS234543 | –   | KS234689 | KS254941 |
| *L. mahajangana*              | CMW 27801T  | FJ900595 | –   | FJ900641 | KS900630 |
| *L. margaritacea*             | CMW 26162T  | EU144050 | KX464354 | EU144065 | KS887520 |
| *L. mediterranea*             | CMM 137783T | KJ638312 | –   | KJ638331 | KS887521 |
| *L. missouriana*              | UCD2193MO   | HQ288225 | –   | HQ288267 | HQ288304 |
| *L. mitidjana*                | ALG111T     | MN104115 | –   | MN104115 | –     |
| *L. parva*                    | CBS 456.78T | EF622083 | KF676362 | EF622063 | KS887523 |
| *L. parva*                    | CBS 494.78  | EF622084 | EU673258 | EF622064 | EU673114 |
| *L. plurivora*                | CBS 120832T | EF445362 | KX464356 | EF445395 | KS887524 |
| *L. pontae*                   | CMW 1277T   | KT151794 | –   | KT151791 | KT151797 |
| *L. pseudomamillata*          | CBS 116459T | EF622077 | EU673256 | EF622057 | EU673111 |

Table 1. GenBank accession numbers of isolates included in this study. Ex-type isolates are labelled with superscript T.
| Species                  | Isolate no. | GenBank no.          | ITS      | LSU      | tef1    | tub2    |
|-------------------------|-------------|----------------------|----------|----------|---------|---------|
| *L. pyriformis*         | CMW 25414T  | EU101307             | –        | EU101352 | KU887527|
| *L. rubropurpurea*      | WAC 12535T  | DQ103553             | DQ377903 | DQ103571 | EU673136|
| *L. sterculiae*         | CBS 342.78T | KX464140             | JX681073 | KX464634 | KX464908|
| *L. subglobosa*         | CMM 3872T   | KF234558             | –        | KF226721 | KF254942|
| *L. syzygii*            | MFLUCC 19-0219.1T | MT990531       | MT990548 | MW016943 | MW014331|
| *L. syzygii*            | GUCC 9719.2 | MW081991             | MW081988 | MW087101 | MW087104|
| *L. syzygii*            | GUCC 9719.3 | MW081992             | MW081989 | MW087102 | MW087105|
| *L. syzygii* sp. nov.   | GUCC 9719.4 | MW081993             | MW081990 | MW087103 | MW087106|
| *L. thailandica*        | CPC 22795T  | KJ193637             | –        | KJ193681 | –       |
| *L. theobromae*         | CBS 164.96T | AY640255             | EU673253 | AY640258 | KU887532|
| *L. venezuelensis*      | WAC 12539T  | DQ103547             | DQ377904 | DQ103568 | KU887533|
| *L. viticola*           | UCD 2553AR  | HQ288227             | –        | HQ288269 | HQ288306|
| *L. vitis*              | CBS 124060T | KX464148             | KX464367 | KX464642 | KX464917|
| *Botryosphaeria dothidea* | CMW 8000T  | AY236949             | AY928047 | AY236998 | AY236927|
| *B. fabicerciana*       | CBS 127193T | HQ332197             | MF410028 | HQ332213 | KF779068|

Table 2.
DNA base pair differences between *Lasiodiplodia syzygii* and *L. rubropurpurea* in four separate loci. 

| *L. syzygiumae* strains | *Lasiodiplodia rubropurpurea* WAC 12535T |
|-------------------------|----------------------------------------|
|                         | ITS (1–530) | LSU (531–1421) | TEF1-a (1422–1748) | β-tubulin (1749–2177) |
| MFLUCC 19-0257=GUCC 9719.1T | 7 | 5 | 34 | 9 |
| GUCC 9719.2            | 7 | 5 | 34 | 9 |
| GUCC 9719.3            | 7 | 5 | 34 | 9 |
| GUCC 9719.4            | 7 | 5 | 34 | 9 |
| Total number of differences | 55 |

**Phylogenetic analyses**

Sequences of 45 *Lasiodiplodia* isolates, representing all species known from culture, were aligned using the online version of MAFFT v. 7.307 (Katoh and Standley 2016) and manually improved, where necessary, using MEGA v. 6.06 (Koichiro et al. 2013). Mesquite v. 2.75 (Maddison 2008) was used to concatenate the aligned sequences of the different loci. Ambiguous regions were excluded from analyses using AliView (Larsson 2014), gaps were treated as missing data and optimised manually with *Botryosphaeria dothidea* (CMW8000) and *B. fabicerciana* (CBS 127193) as the outgroups (Table 1). The alignment document has been deposited in TreeBASE ([www.treebase.org](http://www.treebase.org)) and the accession
number is 27461. Phylogenetic analyses were constructed by Maximum Parsimony (MP), Maximum Likelihood (ML) and Bayesian Inference methods. First, the ambiguous regions were excluded from the alignment and gaps were treated as missing data. The MP analysis was done with PAUP v. 4.0b10 (Swofford 2002), using the heuristic search option with 1,000 random taxa addition and tree bisection and reconnection (TBR) as the branch swapping algorithm. Maxtrees was set to 5000. Tree length (TL), consistency index (CI), retention index (RI), rescaled consistency index (RC) and homoplasy index (HI) were calculated for each tree generated. The Maximum Likelihood (ML) analysis was performed using IQ-tree (Nguyen et al. 2015, Chernomor et al. 2016). Nucleotide substitution models were selected under the Akaike Information Criterion (AIC) by jModelTest2 (Darriba et al. 2012) on XSEDE in the CIPRES web portal (Miller et al. 2010). For the ITS dataset, the TPM3uf+I model was selected (-lnL = 1316.7068), for LSU, the TrN+I (-lnL = 1643.7273), for TEF1-a, the HKY+I+G (-lnL = 2399.0528) and for β-tubulin, the TIM3+G (-lnL = 1161.0392). ML was inferred under partitioned models. Non-parametric bootstrap analysis was implemented with 1000 replicates. Bayesian Inference (BI) analyses was conducted in MrBayes 3.2 (Ronquist et al. 2012). MrModeltest v.2.3 (Nylander 2004) was used to estimate the best evolutionary models under the Akaike Information Criterion (AIC). HKY+I was selected as the best model for ITS, for LSU, HKY+I+G, for TEF1-a, HKY+I+G and for β-tubulin, GTR+G was selected as the best model. Six Markov Chain Monte Carlo runs were launched with random starting trees for 1,000,000 generations and sampling every 1,000 generations. The first 25% resulting trees were discarded as burn-in.

Pathogenicity tests

One isolate of the new *Lasiodiplodia* species (GUCC 9719.1) was grown on PDA and when the cultures covered the entire surface of the Petri dish, mycelia were scraped off with a sterilised blade. Conidiomata were crushed with a glass rod to prepare a spore suspension of 1 x 10^5 spores/ml. Pathogenicity testing was carried out on five healthy fruits of wax apple bought from the market. Inoculations were carried out in April 2020. The surface of the fruits was wiped with 70% ethanol and allowed to air-dry. Three fruits were slightly wounded by pin-pricking and 3 ml of spores suspension was sprayed on to the wound. The other two wounded fruits were maintained as control and inoculated with 2 ml of sterile deionised water. All inoculated fruits were placed in plastic bags, labelled and a high level of humidity was maintained for seven days by the addition of wet sterile cotton wool in each bag in an illuminated incubator at 28 ± 3°C. Daily observations were made on the development of disease symptoms. When fruits developed the symptoms, they were removed from the bags. Two isolates obtained from the diseased tissue were grown on PDA and then sequenced with primer pairs of the above four DNA markers to confirm the identity.
Taxon treatment

*Lasiodiplodia syzygii* C.R. Meng, Qian Zhang & Yong Wang bis, sp. nov.

- MycoBank [837701]

**Materials**

**Holotype:**
- scientificName: *Lasiodiplodia syzygii*; kingdom: Fungi; class: Dothideomycetes; order: Botryosphaeriales; family: Botryosphaeriaceae; genus: *Lasiodiplodia*; country: Thailand; stateProvince: Chiang Rai; catalogNumber: HGUP 9719; recordedBy: Wang Yong; identifiedBy: Chao-Rong Meng; dateIdentified: 2020; type: ex-type living culture GUCC 9719.1; MFLU 19-0565, isotype, isotype living culture MFLUCC 19-0257.

**Other material:**
- scientificName: *Lasiodiplodia syzygii*; kingdom: Fungi; class: Dothideomycetes; order: Botryosphaeriales; family: Botryosphaeriaceae; genus: *Lasiodiplodia*; country: China; stateProvince: Guiyang; catalogNumber: HGUP 9720 and HGUP 9721; recordedBy: Wang Yong; identifiedBy: Chao-Rong Meng; dateIdentified: 2020; type: living cultures GUCC 9719.2, GUCC 9719.3 and GUCC 9719.4

**Description**

Pathogenic on *Syzygium samarangense*. **Sexual morph:** Undetermined. **Asexual morph** (Fig. 2): *Conidiomata* up to 2 mm diam., pycnidial, covered with hyphae, black, globose, ostiolate, solitary, separate, uniloculate, immersed to semi-immersed. *Conidiomatal wall* composed of thick-walled, dark brown cells of textura angularis, becoming thin-walled and hyaline towards the inner region. *Paraphyses* cylindrical, aseptate, hyaline. *Conidiophores* reduced to conidiogenous cells. *Conidiogenous cells* 10–14.5 × 3.5–4.5 μm (average = 11 × 3.7 μm, n = 20), hyaline, smooth, holoblastic forming conidia at their tips. *Conidia* thick-walled, wall up to 1 μm wide, ovoid with both ends rounded, hyaline and remaining so for a long time, becoming pale brown with obsolete striations and occasionally with 1-septate after discharging from the conidioma, (27–)30–32(–36) × (13–)15–17(–20) μm (average = 31.3 × 16.4 μm, n = 50), L/W = 1.9.

**Culture characteristics:** Conidia germinate on PDA within 24 hours at room temperature (25–30°C) with germ tubes produced from both ends of the conidia. Colonies with white fluffy mycelium on PDA, after 7 days become olivaceous-grey at the centre, white at the edge, raised, fluffy, dense filamentous.

**Notes:** *Lasiodiplodia syzygii* strains are closely related to *L. rubropurpurea*, but formed a distinct, well-supported clade in the phylogenetic analyses. Base-pairs comparisons between *L. syzygii* ex-type strain (GUCC 9719.1) and ex-type strain of *L. rubropurpurea* (WAC 12535) found seven base differences (1.3%) in ITS region and five differences (0.6%) on LSU, but nine differences (2.1%) in *tub2* and 34 in TEF1-a (10.4%) (Table 2). *Lasiodiplodia syzygii* produced larger pycnidia (up to 2 mm) and larger conidiogenous cells (10–14.5 × 3.5–4.5 μm) than *L. rubropurpurea* (0.5–1.5 mm and 7–13 × 3–5 μm) (Burgess et al. 2006).
**Etymology**

In reference to the host from which the fungus was first isolated.

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**Figure 1.**

One of 850 most parsimonious trees obtained from a combined analyses of the ITS, LSU, TEF1-a and β-tubulin sequence dataset. Bootstrap values > 50% and BPP values > 0.90 are provided at the nodes and separated by “/”. Bootstrap values < 50% and Bayesian posterior probability (BPP) values < 0.90 were labelled with “-”. The tree was rooted with *Botryosphaeria fabriciana* (CBS 127193) and *B. dothidea* (CMW 8000). The branch of the new *Lasidiodiplodia* species is highlighted with pink.

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Analysis

Phylogenetic analyses

Four *Lasiodiplodia* strains isolated from *Syzygium samarangense* were sequenced. The final alignment of ITS, LSU, TEF1-a and *tub2* comprised of 2177 characters, viz. ITS: 1–530, LSU: 533–1423, TEF1-a: 1426–1752 and β-tubulin: 1755–2183. Of these, 1843 characters were constant and 73 were parsimony-uninformative. Maximum parsimony analysis of the remaining 261 parsimony-informative characters resulted in 850 most parsimonious trees (TL = 676, CI = 0.64, RI = 0.81, RC = 0.52 and HI = 0.36) and the first
one is shown as Fig. 1. The ML and Bayesian analyses resulted in trees with similar topologies. Strains GUCC 9719.1, GUCC 9719.2, GUCC 9719.3 and GUCC 9719.4 formed an independent well-supported clade sister to *Lasiodiplodia rubropurpurea* (MP: 100%, ML: 100% and Bayesian posterior probability: 1) Comparison of the DNA base-pair differences between our strains and *L. rubropurpurea* species in four gene regions (Table 2) confirmed the presence of two species; therefore, a new species is introduced for those isolates from wax apple.

**Pathogenicity test on the fruits of wax apple**

At the third day after inoculation, water-soaked areas with a few white hyphae began to appear on all inoculated fruits similar to the naturally-infected wax apples (Fig. 2a and Fig. 3a). The water-soaked symptom of diffusion with abundant hyphae producing mycelium further appeared on inoculated *Syzygium samarangense* fruits after five days (Fig. 3b). At the 7th day after inoculation, the symptoms spread throughout the fruit (Fig. 3c), together with many white mycelia and more hyphae accompanied by cytoplasmic exosmosis. The control fruits (Fig. 3d) did not show any symptom. The fungi were re-isolated from the lesions of inoculated wax apple fruits and the re-identified (GUCC 9719.3 and GUCC 9719.4) sequencing four gene regions.

![Figure 3. Symptoms developing on Syzygium samarangense fruits inoculated with Lasiodiplodia syzygii. a. Symptom at 3rd day; b. Symptom at 5th day; c. Symptom at 7th day; d. Control.](image-url)
Discussion

This study revealed a new species of *Lasiodiplodia*, *L. syzygi* from rotting fruits of *Syzygium samarangense*. Phylogenetic analyses, based on ITS, LSU, TEF1-a and *tub2*, showed that it is phylogenetically closer to *L. rubropurpurea*. Comparisons of DNA base-pair differences in the four loci, as well as morphological differences, confirmed the novelty of this species. The fungus was proved to be pathogenic and, therefore, it is the causal agent of the post-harvest water-soaked brown lesions on wax apple.

Wax apple (*Syzygium samarangense*) is known to be affected by many fungal pathogens that often cause economic losses. These include *Colletotrichum gloeosporioides* (Udayanga et al. 2013) and *Lasiodiplodia theobromae* which was the causal agent of black spot disease (Che et al. 2015), *Pestalotiopsis* spp. and *Phytophthora* spp. The fruit disease of the current study did not show any typical symptoms of black spot caused by *L. theobromae*. Furthermore, the pink or orange spore masses, typical of anthracnose caused by *C. gloeosporioides* or epidermal to superficial, acervular conidiomata reported by Maharachchikumbura et al. (2013) for *Pestalotiopsis*, were not seen in the current study. The fruit rot caused by *Phytophthora* spp. spread more rapidly (only 2 or 3 days up to a whole fruit) and results in a sour taste on fruits. However, the *L. syzygii* needed about seven days to completely rot the fruit and did not cause any sour taste in the fruits. Thus, the study reports a new disease on wax apple.

*Lasiodiplodia* resides in Botryosphaeriaceae, *Botryosphaeriales* (Hongsanan et al. 2020) and comprises several species known to cause important or potentially important diseases on woody hosts, mostly in the tropics or sub-tropics (Phillips et al. 2019). Very few species of this family appear to be host-specific (Dissanayake et al. 2016). In south-western China and adjoining areas, agriculture and forestry play an important role in the local economy, which might facilitate the spread of this wax apple disease. Thus, research needs to focus on the occurrence of this newly-discovered pathogen in other economically-important plants and in other locations, as well as how to manage it by biological or chemical control approaches. It is also remarkable to find a new disease on such an important commercial fruit indicating that there are numerous new taxa to be discovered in Thailand (Hyde et al. 2018) and Botryosphaeriaceae (Hyde et al. 2020).

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