First Report of Two Colletotrichum Species Associated with Bitter Rot on Apple Fruit in Korea – C. fructicola and C. siamense

Myung Soo Park*, Byung-Ryun Kim**, In-Hee Park** and Soo-Sang Hahn*

*School of Biological Sciences and Institute of Microbiology, Seoul National University, Seoul, Korea; **Environmentally Friendly Agriculture Division, Chungnam Agricultural Research and Extension Services, Yesan, Korea

ABSTRACT
Bitter rot caused by the fungal genus Colletotrichum is a well-known, common disease of apple and causes significant yield loss. In 2013, six fungal strains were isolated from Fuji apple fruits exhibiting symptoms of bitter rot from Andong, Korea. These strains were identified as Colletotrichum fructicola and C. siamense based on morphological characteristics and multilocus sequence analysis of the internal transcribed spacer rDNA, actin, calmodulin, chitin synthase, and glyceraldehyde-3-phosphate dehydrogenase. Pathogenicity tests confirmed the involvement of C. fructicola and C. siamense in the development of disease symptoms on apple fruits. This is the first report of C. fructicola and C. siamense causing bitter rot on apple fruit in Korea.

1. Introduction
Bitter rot caused by fungi in the genus Colletotrichum is one of the most common diseases of apple fruit and causes significant yield loss of apple crop worldwide [1]. Colletotrichum gloeosporioides and C. acutatum are the two most common species causing bitter rot on apples [2]. It is difficult to identify Colletotrichum to species based only on morphology, due to the morphological similarity at different growth conditions [3]. Recently, Colletotrichum was studied based on morphology and multilocus phylogenetic analysis, grouping the 189 recognized species into 11 species complexes [4]. Additional studies described several new species from different hosts in the C. acutatum and C. gloeosporioides complexes [5,6]. Of the described species, five have been reported to cause bitter rot of apple fruit: C. floriniae and C. nymphaeae (C. acutatum species complex) and C. siamense, C. theobromicola, and C. fructicola (C. gloeosporioides species complex) [1].

Apple (Malus pumila Mill.) is grown worldwide. In Korea, the total area of apple cultivation is approximately 31,620 ha and is gradually increasing (KOSTAT, Statistics Korea, http://kostat.go.kr/portal/eng/). To date, about 40 pathogens have known associations with apple in Korea, including anthracnose, white rot, Alternaria leaf spot, Marssonina blotch, and bacterial shoot blight [7]. Two species of Colletotrichum (C. gloeosporioides and C. acutatum) were reported to be associated with apple anthracnose disease in Korea [7]. Bitter rot is a serious disease of apple fruit in Korea and causes significant yield loss [8]. C. gloeosporioides and C. acutatum were previously identified as causative agents bitter rot of apple in Korea [7,9]. These studies were based on morphological features and sequence analysis of a single gene (ITS or β-tubulin) – data that has now been shown to be unreliable (morphological plasticity, low resolution of DNA) for species identification. The aim of the present study was to identify the causative agent of bitter rot on apple fruit from Andong, Korea. We identify six unknown fungal strains associated with bitter rot based on morphological features and multilocus sequence analyses of the internal transcribed spacer (ITS) rDNA, actin (ACT), calmodulin (CAL), chitin synthase (CHS-1), and glyceraldehyde-3-phosphate dehydrogenase (GAPDH), and provide verification with pathogenicity tests.

2. Materials and methods
2.1. Sampling and isolation
Fungal isolates were collected from Fuji apple fruit with symptoms of bitter rot collected from orchards in Andong, Korea during 2013. The fruit tissues were surface-sterilized by dipping them in 0.1%...
Genomic DNA was extracted directly from mycelia of fungal strains using a modified cetyltrimethylammonium bromide (CTAB) extraction protocol [10]. PCR amplifications of ITS, ACT, CAL, CHS-1, and GAPDH were performed in a C100 Thermal Cycler (Bio-Rad, Hercules, CA, USA) using previously described methods [6]. DNA sequencing was performed at Macrogen (Seoul, Korea), using an ABI PRISM 3730XL Analyzer (Life Technologies, Gaithersburg, MD, USA). Sequences were proofread and edited using MEGA ver. 5.0 [11] and aligned with reference sequences of type strains downloaded from GenBank using the default settings of MAFFT v7 [12]. Maximum likelihood phylogenetic analyses were performed using RAxML [13] for an ITS only dataset for initial identification, followed by the concatenated data set (ITS, ACT, CAL, CHS-1, and GAPDH) for species identification. Analyses were run with the GTR + G model of evolution and 1000 bootstrap replicates. All sequences were deposited in GenBank (Table 1).

2.3. Culture and morphological characteristics

To observe macroscopic culture characteristics, the strains were cultured on PDA and incubated at 25 °C for one week. A mycelium plug (5 mm diameter) was taken from the actively growing edge of the colony and incubated on PDA at 25 °C for seven days under dark conditions. To observe microscopic characters, mounts of strains were made in 3% KOH (lactic acid). Measurements were performed using a light microscope (Nikon Eclipse 80i) for at least 30 conidia.

Table 1. Information and GenBank accession numbers of Colletotrichum strains isolated this study.

| Species            | Collection no. | Location          | Pathogenicity | ITS             | ACT             | CAL             | CHS-1            | GAPDH          |
|--------------------|----------------|-------------------|---------------|-----------------|-----------------|-----------------|-----------------|----------------|
| C. fructicola      | CNARE13111     | Andong, Gyengangbuk-do | +             | MG807419        | MG831722        | MG831728        | MG831734        | MG831740        |
| CNARE13132         | Andong, Gyengangbuk-do | +             | MG807420        | MG831723        | MG831729        | MG831735        | MG831741        |
| C. siamense        | CNARE13100     | Andong, Gyengangbuk-do | +/+           | MG807421        | MG831724        | MG831730        | MG831736        | MG831742        |
| CNARE13105         | Andong, Gyengangbuk-do | ++            | MG807422       | MG831725        | MG831731        | MG831737        | MG831743        |
| CNARE13126         | Andong, Gyengangbuk-do | +++           | MG807423       | MG831726        | MG831732        | MG831738        | MG831744        |
| CNARE13152         | Andong, Gyengangbuk-do | +++           | MG807424       | MG831727        | MG831733        | MG831739        | MG831745        |

*: no infection and symptom; ±: 0.1–6 mm; ++: 7–12 mm; and +++: 12 mm.

3. Results

3.1. Phylogenetic analyses

Based on the ITS sequence analysis and morphological features, six strains isolated from apples with bitter rot symptoms were identified as members of the C. gloeosporioides species complex (data not shown). Next, for the analysis of the concatenated dataset (ITS, ACT, CAL, CHS-1, and GAPDH) to identify the unknown strains to species level, we increased sampling of GenBank sequences of type strain in the C. gloeosporioides species complex. The corresponding lengths of the aligned fragments of six strains for the five loci were as follows: 582 bp for ITS; 275 bp for ACT; 708 bp for CAL; 286 bp for CHS-1; 283 bp for GAPDH. Two strains (CNARE13111 and CNARE13132) formed a monophyletic group with C. fructicola CBS 130416 (type strain from Coffea arabica) and CBS125397 (from Tetragastris panamensis) (100% bootstrap value),
Table 2. Comparison of the morphological characteristics of the strain used in this study with those of previously reported C. fructicola and C. siamense.

| Characteristic       | C. fructicola CNARE13132 | C. fructicola* | C. siamense CNARE13126 | C. siamense* |
|----------------------|--------------------------|---------------|-------------------------|--------------|
| Colony morphology    | Cottony, gray to dark gray at the center | Cottony, gray to dark gray at the center | Cottony, aerial mycelium grayish white, orange conidial masses at the center | Cottony, aerial mycelium white, orange conidial masses at the center |
| Conidia size (µm)    | 11.0–14.0 × 4.5–5.5 | 11.5–17.5 × 3–5.5 | 11.0–14.5 × 4.0–5.0 | 12.0–15.5 × 4.0–5.5 |
| Conidia shape        | Cylindrical, both ends rounded | Cylindrical, both ends rounded | Cylindrical, both ends bluntly rounded | Cylindrical, both ends bluntly rounded |
| Colony radius (mm)   | 71–77 | 74–79 | 65–72 | 79 |

*Described by Lee et al. [9].

3.2. Taxonomy

**Colletotrichum fructicola** Prihastuti, L. Cai & K.D. Hyde., Fungal Divers. 39: 89–109 (2009) (Table 2; Figure 2(b)).

Colonies on PDA initially white mycelia that became grey to dark grey at the center with age on the front side and greyish green on the reverse side. Colonies grew 10.1–11.0 mm/day at 25 ± 1°C under dark and were 70.7–77.0 mm diameter after seven days. Aerial mycelium dense, cottony pale grey, without visible conidial masses. Acervuli absent in culture. Setae absent. Conidia common in mycelium, one-celled, smooth-walled, hyaline, straight, aseptate, cylindrical with both ends rounded, sometimes oblong, sizes ranging 11.0–14.0 × 4.5–5.5 μm. Chlamydospores were not observed.

Remarks: Colletotrichum fructicola is morphologically similar to Colletotrichum kahawae, but it can be distinguished from C. kahawae by small conidia and rapid growth on PDA [3].

**Colletotrichum siamense** Prihastuti, L. Cai & K.D. Hyde., Fungal Divers. 39: 89–109 (2009) (Table 2; Figure 2(b)).

Colonies on PDA initially greyish white mycelia that became pale brownish to pinkish with age on the front side and pinkish on the reverse side. Colonies grew 9.3–10.3 mm/day at 25 ± 1°C under dark and were 65.1–72.1 mm diameter after seven days. Aerial mycelium dense, cottony, greyish white, with visible conidial masses at inoculum. Setae absent. Conidia common in mycelium, one-celled, smooth-walled, hyaline, cylindrical with both ends bluntly rounded, sometimes oblong, sizes ranging 11.0–14.5 × 4.0–5.0 μm.

Remarks: Colletotrichum siamense is morphologically similar to C. acutatum, but it can be distinguished from C. acutatum by cylindrical conidia with both ends bluntly rounded [3].

3.3. Pathogenicity

Despite differences in virulence, all strains produced characteristic signs of bitter rot on apple fruit, such as a sunken zone with concentric rings of conidial masses. C. siamense (CNARE13100, CNARE13105, CNARE13126, and CNARE13152) produced larger...
lesions compared to *C. fructicola* (CNARE13111 and CNARE13132). Of the six strains isolated in this study, strain CNARE13126 of *C. siamense* showed the strongest virulence (Table 2; Figure 2).

4. Discussion

Apple fruits from Andong, Korea collected in 2013 showed characteristic signs of bitter rot—a sunken zone with concentric rings of conidial masses. Based on morphological features, multilocus sequence analyses, and pathogenicity tests, we identified *C. fructicola* and *C. siamense* to be the causative agents of bitter rot of apple fruit in Korea. To the best of our knowledge, this is the first report of *C. fructicola* and *C. siamense* in Korea causing bitter rot on apple fruit. To the best of our knowledge, this is the first report of *C. fructicola* and *C. siamense* in Korea causing bitter rot on apple fruit.

*C. fructicola* and *C. siamense*, members of the *C. gloeosporioides* species complex, were first described as a pathogen of *C. arabica* berries [15]. These species have wide host range including apple and are associated with bitter rot worldwide [6].

Many new *Colletotrichum* species from various hosts were described using morphological features and multilocus phylogenetic analysis of ITS, ACT, CAL, CHS-1, and GAPDH [1,5,6,16]. *C. gloeosporioides* is known as one of the most important pathogens that infect a range plant species. However 25 isolates of *C. gloeosporioides* from tropical fruits were re-identified as *C. asianum*, *C. fructicola*, *C. horii*, *C. kahawae*, and *C. gloeosporioides* based on morphological characteristics and multilocus phylogenetic analysis of ITS, ACT, CAL, CHS-1, and GAPDH [16]. Seven *Colletotrichum* species are known as causal species of bitter rot of apple (*C. gloeosporioides*, *C. acutatum* *C. fioriniae*, *C. fructicola*, *C. nymphaeae*, *C. siamense*, and *C. theobromicola*) [1,2]. *C. gloeosporioides* and *C. acutatum* were previously reported to cause bitter rot of apples in Korea (Korea disease list). These two species were not found in this study; we may have missed these species due to a low number of isolates in our study, or strains in previous studies may have been misidentified. Further study of bitter rot of apple fruit in Korea is needed to clarify this situation.

Bitter rot in Korea is caused by at least two *Colletotrichum* species, *C. fructicola* and *C. siamense*, with the latter being the more aggressive species. Studies of bitter rot and other plant pathogens should combine morphology multigene phylogenetic analysis for more reliable identification. We expect such an approach will uncover more fungal species associated with plant pathogens. In addition to more studies identifying causative agents of plant pathogens, studies of species distribution can provide valuable information for better management of bitter rot on apple in Korea.

Figure 2. *Colletotrichum fructicola* CNARE13132 (a) and *C. siamense* CNARE13126 (b). (a) Colony morphology on potato dextrose agar (PDA) after 7-day culture at 25°C (front); (b) Colony morphology on PDA after 7-day cultures at 25°C (reverse); (c) Conidia Bar scale, 10 μm (micrometer); (d) Symptoms induced by artificial inoculation.
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