First Report of Anthracnose of Shine Muscat Caused by *Colletotrichum fructicola* in Korea

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**ABSTRACT**

Anthracnose is one of the major problems for cultivating many crops, including vegetables, fruits, and trees. It is a continual threat for fruits grower worldwide. *Colletotrichum fructicola* was isolated from Shine Muscat berries showing typical anthracnose symptom in Korea. It was identified as *C. fructicola* based on morphology, pathological signs and concatenated sequences of internal transcribed spacer region of rDNA, glyceraldehyde-3-phosphate dehydrogenase, [\(\beta\)]-tubulin-2, chitin synthase-1, calmodulin, and the Apn2-Mat1-2 intergenic spacer and partial mating type (Mat1-2) gene. To the best of our knowledge, this is the first report first report of anthracnose of Shine Muscat caused by *C. fructicola* in Korea.

Shine Muscat (*Vitis vinifera* L.) is the grape cultivar bred by crossing Akitsu-21 (*Vitis labruscana* Baily × *V. vinifera*) and "Hakunan" (*V. vinifera*) [1]. Ever since its introduction to South Korea, it has become popular for its sweet seedless large grapes, and unique mango flavor. Grape growers have replaced their traditional Campbell cultivar with Shine Muscat and therefore the cultivation area for Shine Muscat has been growing rapidly in South Korea [2].

Anthracnose is a group of diseases caused by fungi especially *Colletotrichum* spp., that affects many plants, including vegetables, fruits, and trees. It is destructive to fruits, causing pre- and post-harvest decay. In August 2018, outbreak of anthracnose on Shine Muscat fruits occurred in a commercial orchard, Daegu, Korea and the disease incidence was approximately 25%. Symptoms were observed on the surface of fruits in the form of brown sunken lesion with concentric rings (Figure 1(A,B)). For the isolation of the causal agents, infected fruits showing typical anthracnose symptom brought to laboratory. Small sections containing healthy and necrotic tissues were cut from infected fruits, disinfested by immersing in 0.5% NaOCl for 3 min. The disinfested section of tissue rinsed in sterile water, dried by blotting, placed on water agar amended with 250 ppm streptomycin and incubated at 25°C in the dark. After 3 days of incubation, fungal hyphae were emerged from the margin of infected tissue. Five-millimeter diameter plug of newly emerging hyphae was transferred to potato dextrose agar (PDA) and incubated at 25°C in the dark. Pure cultures were obtained by following single spore isolation technique [3]. Primary conidial suspension was prepared in sterilized distilled water from 18 days old previously isolated colony. Concentration of conidial suspension was estimated using hemocytometer. Then 1 × 10^4 conidial suspension was prepared from the primary conidial suspension, sprayed over fresh PDA pates using sterilized wire loop and incubated at 25°C in the dark. After 3 days, a single germinating conidium was transferred to new PDA plates, incubated at 25°C in the dark and morphology of the colony was examined. Based on the similarity of the colony morphology, three isolates (ICKG2, ICKG4, and ICKG6) were selected for further morphological and molecular characterization. Twenty-day-old cultures were gray to white with immersing black perithecia in PDA (Figure 1(C,D)). For microscopic observation of teleomorph, 3–4 perithecia were picked up and mounted with lactophenol cotton blue on a glass slide. The sample was crushed by hitting gently over cover slide. Perithecia were brown and globose (Figure 1(E)). Ascospores were hyaline, slightly curved with round ends with the dimension of 13.5–25.6 μm × 4.0–6.1 μm (mean ± SD = 19.2 ± 2.9 × 5.0 ± 0.64 μm) (n = 25) (Figure 1(F)). Small mycelial mass was mounted on glass slide in...
sterilized distilled water for morphological analysis of morph. Abundant conidia were produced on short conidiophore. The conidia were hyaline, cylindrical, straight with rounded ends and 12.1–21.1 μm x 5.5–8.2 μm (mean ± SD = 159 ± 26 × 67 ± 9 μm) (n = 50) (Figure 1(G,H)). The glass slide containing conidia were put in a petri plate with moist tissue paper and incubated at 25°C in the dark for enhancing appressoria formation. Appressoria were brown, various in shape with smooth edge and 9.4–13.7 μm x 5.9–8.2 μm (mean ± SD = 10.8 ± 1.3 × 7.1 ± 0.7 μm) (n = 20) (Figure 1(G)). The morphological characteristics of present isolates were match with those of previous described Colletotrichum species within

Figure 1. Anthracnose of Shine Muscat caused by Colletotrichum fructicola. (A) Symptomatic Shine Muscat berries in infected orchard; (B) Sign and symptom appear on inoculated berries; (C, D) 20 days old fungal culture on PDA; (E) Perithecium; (F) Ascospore; (G) Conidiophore and appressoria (scale bar = 10 μm); (H) Conidia (scale bar = 10 μm).
C. gloeosporioides complex including C. fructicola [4–7]. Morphological characteristics of present isolates and close C. fructicola isolates are described in Table 1. Slight difference in morphological characteristics can be explained by growth media and type of host. Morphological characteristics of same species vary with the culture media and host type [4,8].

Molecular analysis could be the best way to support morphological analysis. For achieving this objective, HiGene Genomic DNA Prep Kit (Biofact, Daejeon, South Korea) was used to extract total genomic DNA from selected isolates following manufacturer instructions. Internal transcribed spacer (ITS) region of rDNA and partial gyceraldehyde-3-phosphate dehydrogenase (GAPDH), β-tubulin-2 (TUB2), chitin synthase-1 (CHS-1), calmodulin (CAL), and the Apn2-Mat1-2 intergenic spacer and partial mating type (ApMat) genes were amplified using primer sets ITS1F and ITS4 [9,10], GDF and GDR [11], BT2a and BT2b [12], CHS-79F and CHS-345R [13], CL1C and CL2C [4], and AM-F and AM-R [14], respectively. Resulting PCR products were purified, sequenced through Macrogen Inc. (Seoul, Korea). The obtained sequences of each gene region were subjected to sequence similarity analysis using Basic Local Alignment Search Tool (BLAST) and deposited in GenBank [15]. The accession number and BLAST analysis result of each gene of present isolates is shown in Table 2. BLAST search result revealed that present isolates were 99–100% molecularly similar with the C. fructicola isolates (Table 2).

For phylogenetic analysis, individual gene sequences of present isolates and fungal species within C. gloeosporioides complex were aligned with MEGA v. 6.0 using MUSCLE program and then concatenated with Mesquite v. 2.75 [16,17]. Two phylogenetic trees were constructed using maximum likelihood and maximum parsimony statistical method from concatenated sequences (ITS, GAPDH, TUB2, CHS-1, CAL, and ApMat) in MEGA 6 [16]. The phylogenetic analysis delineated the present isolates from shine mascal as C. fructicola (Figure 2). Koch’s postulates were performed by inoculating healthy V. vinifera fruits with conidial suspension (1 × 10⁶ conidia/ml) of ICKG2 isolate using wounding and unwounding methods. Seven-day-old culture of fungal isolate ICKG2 was used for preparing the conidial suspension. Fresh and healthy berries were collected from uninfected shine mascal orchard, surface sterilized with 0.5% NaOCl for 3 min followed by rinsing 2 times with sterile distilled water. The sterilized berries were dried by blotting and prepared for inoculation. For wounding method, berries were wounded with sterile pin to 1-mm depth and insulated with 10 μL droplet of conidial suspension. For unwounding method, berries were sprayed with conidial suspension. The experiment was replicated 15 times (one fruits per replicate).

Fruits received sterile water served as control. Treated fruits were incubated at 25°C in the dark in a box with moist paper tissue. Anthracnose symptoms appeared on fruits (Figure 1(B)), inoculated by wounding method after 7 days and disease incidence was (80%). Only 30% of inoculated fruits with unwounding method showed symptom after 14 days, whereas control fruits remained asymptotic. Same fungus (C. fructicola) was reisolated from inoculated fruits and identified following the methods described above. Based on morphology, molecular

### Table 1. Comparison of morphological characteristics between present isolate and previously reported Colletotrichum fructicola.

| Characteristics | Present isolate (ICK G4) | Colletotrichum fructicola* |
|-----------------|--------------------------|---------------------------|
| Ascospores      | Slightly curved with round ends | Fusiform-to-slightly curved with rounded ends |
| Color           | Hyaline                  | Hyaline                   |
| Size (μm)       | 13.5 ~ 25.6 × 4.0 ~ 6.1 | 13.6 ~ 24.0 × 2.6 ~ 6.2 |
| Conidia         | Cylindrical, straight with rounded ends | Cylindrical, straight with rounded ends |
| Color           | Hyaline                  | Hyaline                   |
| Size (μm)       | 12.1 ~ 21.1 × 5.5 ~ 8.2 | 13.1 ~ 19.8 × 3.1 ~ 7.0 |
| Appressoria     | Various in shape with smooth edge | Oval to fusoid with smooth edges |
| Color           | Brown                    | Brown                     |
| Size (μm)       | 9.4 ~ 13.7 × 5.9 ~ 8.2  | –                        |

*Description and illustration [6].

### Table 2. Molecular similarity of present isolates and previously reported Colletotrichum fructicola isolates.

| Present isolates | Gene (accession number) | Most similar Colletotrichum spp. isolates |
|------------------|-------------------------|------------------------------------------|
| ICKG2            | ITS (LC469121)          | C. fructicola isolate 22-21 (100%)       |
|                  | GAPDH (LC469127)        | C. fructicola isolate CPC 25976(100%)    |
|                  | TUB2 (LC469124)         | C. fructicola isolate HLNY-18 (99.76%)   |
|                  | CHS-1 (LC469130)        | C. fructicola isolate HJU-11 (100%)      |
|                  | CAL (LC469133)          | C. fructicola isolate HJU-11 (100%)      |
|                  | ApMat (LC469136)        | C. fructicola isolate CDTZG (100%)       |
| ICKG4            | ITS (LC469122)          | C. fructicola isolate 23-21 (99.47%)     |
|                  | GAPDH (LC469128)        | C. fructicola isolate CPC 25976(100%)    |
|                  | TUB2 (LC469125)         | C. fructicola isolate HLNY-18 (99.53%)   |
|                  | CHS-1 (LC469131)        | C. fructicola isolate HJU-11 (100%)      |
|                  | CAL (LC469134)          | C. fructicola isolate HJU-11 (99.86%)    |
|                  | ApMat (LC469137)        | C. fructicola isolate CDTZG (100%)       |
| ICKG6            | ITS (LC469123)          | C. fructicola isolate 23-21 (99.16%)     |
|                  | GAPDH (LC469129)        | C. fructicola isolate CPC 25976(100%)    |
|                  | TUB2 (LC469126)         | C. fructicola isolate HLNY-18 (100%)     |
|                  | CHS-1 (LC469132)        | C. fructicola isolate HJU-11 (100%)      |
|                  | CAL (LC469135)          | C. fructicola isolate HJU-11 (100%)      |
|                  | ApMat (LC469138)        | C. fructicola isolate CDTZG (100%)       |
characterization, and pathological signs, present isolates were identified as *C. fructicola*.

*C. fructicola* is a representative fungal species with in Musae clade of *C. gloeosporioides* complex [4]. It was isolated from coffee berry and described as a pathogenic fugal species by Prihastuti *et al.* for the first time [5]. Now, it is known as the dominant and aggressive fungal species responsible for anthracnose on many crops including apple, sandy pear (*Pyrus pyrifolia*), strawberry, and avocado [6,7,18,19]. To the best of our knowledge, there were no report on anthracnose of Shine Muscat caused by *C. fructicola* in Korea, this is the first such study. The outcome of this will help to choose the right control measure against anthracnose of Shine Muscat, the increasingly popular grape cultivar in South Korea.

**Disclosure statement**

No potential conflict of interest was reported by the authors.

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