

Didymella acutilobae sp. nov. Causing Leaf Spot and Stem Rot in *Angelica acutiloba*

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ABSTRACT

During disease surveys of *Angelica acutiloba* plants in Korea, leaf spot symptoms were observed in a field in Andong in July 2019, and stem rot symptoms in vinyl greenhouses in Yangpyeong in April 2020. Incidence of leaf spot and stem rot of the plants ranged from 10 to 20% and 5 to 30%, respectively. Morphological and cultural characteristics of fungal isolates from the leaf spot and stem rot symptoms fitted into those of the genus *Phoma*. Molecular phylogenetic analyses of two single-spore isolates from the symptoms using concatenated sequences of LSU, ITS, TUB2, and RPB2 genes authenticated an independent cluster from other *Didymella* (anamorph: *Phoma*) species. Moreover, the isolates showed different morphological and cultural characteristics in comparison to closely related *Didymella* species. These discoveries confirmed the novelty of the isolates. Pathogenicity of the novel *Didymella* species isolates was substantiated on leaves and stems of *A. acutiloba* through artificial inoculation. Thus, this study reveals that *Didymella acutilobae* sp. nov. causes leaf spot and stem rot in *Angelica acutiloba*.

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1. Introduction

Angelica acutiloba (Siebold & Zucc.) Kitag. is a perennial plant belonging to the family Apiaceae. This plant is native to Japan and has been introduced to Korea and Manchuria [1]. It has been traditionally utilized as a medicinal herb with various pharmacological effects in Japan [2] and is cultivated as a vegetable in Korea.

We observed leaf spot symptoms with a yellow halo on *A. acutiloba* plants in a field in Andong, Gyeongbuk Province, Korea, during a disease survey in July 2019. We also observed stem rot symptoms on the plants in vinyl greenhouses in Yangpyeong, Gyeonggi Province, Korea, during a disease survey in April 2020. We obtained fungal isolates from the leaf spot and stem rot symptoms and examined morphological characteristics of the isolates. The morphological and cultural characteristics of the isolates fitted into those of the genus *Phoma* [3]. The isolates were very similar to each other in terms of morphological characteristics.

Morphological characteristics, host relationships, and cultural traits were investigated in the beginning stage for identification of *Phoma* spp. [3,4]. In addition, the genus *Didymella* was revealed as a teleomorph of the genus *Phoma* based on the phylogenetic

studies [4,5]. Many reported *Phoma* spp. have been re-identified into new genus or species gradually based on molecular phylogenetic studies [6–9].

It has been reported that *Didymella* sp. causes leaf spot of *A. acutiloba* in Japan [10]. In Korea, stem blight of *A. acutiloba* caused by *Phoma* sp. is recorded [11]. However, no significant study has not been proceeded to identify a causal agent of leaf spot and stem rot in the plant. This study was conducted to identify *Phoma* sp. isolates from leaf spot and stem rot symptoms of *A. acutiloba* in Korea using a multi-locus phylogenetic analysis with investigations of morphological and cultural characteristics. In addition, pathogenicity test of the isolates was carried out to reveal the causal agent of the leaf spot and stem rot in *A. acutiloba*.

2. Materials and methods

2.1. Disease survey and fungal isolation

We surveyed occurrence of diseases on *A. acutiloba* plants grown in a field in Andong, Gyeongbuk Province, Korea in July 2019 and three vinyl greenhouses in Yangpyeong, Gyeonggi Province, Korea in April 2020. One hundred leaves of the plants were observed for investigation of leaf spot occurrence in

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the field, and 100 plants for investigation of stem rot occurrence in the vinyl greenhouses. The investigation for occurrence of leaf spot and stem rot was conducted at three sites in the field or vinyl greenhouse. Lesion pieces (5–6 mm²) were cut from the leaves and stems of the diseased plants and surface-sterilized with 1% sodium hypochlorite solution for one minute. The sterilized pieces were placed on 2% water agar (WA) and incubated at 25 °C for 3–4 days. Then, growing mycelia from the pieces were transferred onto potato dextrose agar (PDA). The isolates were transferred onto oatmeal agar (OA), and single-spore isolates were obtained from 2-week-old OA cultures. One isolate from the leaf and stem, respectively, was selected among the single-spore isolates for identification and pathogenicity tests. A representative isolate was deposited in the Korean Agricultural Culture Collection (KACC), Wanju, Korea.

2.2. Investigation of cultural and morphological characteristics

Malt extracted agar (MEA), OA, and PDA were used to investigate cultural characteristics of the isolates (ANAC-1901 and ANAC-2001). The isolates were incubated on the media at 22 °C in the dark for seven days. Subsequently, another week of incubation was continued at 22 °C under alternating cycles of 13/11 h of near ultraviolet light and dark [6,7]. NaOH spot test [6] was performed on 1-week-old culture on MEA. Twenty pycnidia and 30 conidia from 2-week-old cultures on OA were observed to investigate their morphological characteristics under the light microscope (Nikon Eclipse Ci-L, Tokyo, Japan).

2.3. DNA extraction, PCR, and sequencing

Genomic DNA of the isolates was extracted using the protocol in a previous study [12], with slight modifications. Polymerase chain reaction (PCR) experiments were conducted to investigate partial large subunit nuclear ribosomal DNA (LSU), internal transcribed spacer regions 1 and 2 including 5.8S nrDNA (ITS), β -tubulin (TUB2), and RNA polymerase II second largest subunit (RPB2) gene regions using LR0R [13] and LR7 [14] for LSU, V9G [15] and ITS4 [16] for ITS, Btub2Fd and Btub4Rd [17] for TUB2, and RPB2-5f2 [18] and fRPB2-7cR [19] for RPB2. Conditions of PCR amplification for all the genes were followed as in the previous studies [7]. Takara Ex Taq (Takara Bio Inc., Kusatsu, Japan) was used to prepare PCR products of the two isolates following the manufacturer's instruction. Purification of the PCR products was conducted using the universal DNA purification kit

(Tiangen, Beijing, China), according to the manufacturer's instruction. The PCR products were sequenced using the same primers at Bionics Co., Ltd. (Seoul, Korea). The sequences were improved manually when necessary by SeqMan II (DNASTAR Inc., Madison, WI). The sequence data were deposited in GenBank.

2.4. Alignment and molecular phylogenetic analysis

The sequences of the isolates obtained from *A. acutiloba* and the pertinent sequences of *Didymella* spp. from the previous studies [7–9,20] (Table 1) were aligned by MUSCLE [21]. *Coniothyrium palmarum* (CBS 400.71) was served as an outgroup taxon. The multiple sequence alignments were conducted and improved when necessary using MEGA version 7 software [22]. The partition-homogeneity test for the sequences was carried out using PAUP version 4.0 software [23]. Then, neighbor-joining (NJ) analysis with the maximum composite likelihood model was performed with 1000 bootstrap (BS) replicates using MEGA version 7 software [22]. Bootstrap values equal to or greater than 50% were indicated at nodes. The evolutionary model for each gene was investigated using MrModeltest version 2.2 software [24]. The Bayesian analysis of the concatenated alignments was conducted by MrBayes version 3.2.4 software [25] based on the results of the model test. The calculation continued until the average standard deviation of split frequencies reached a value of equal to or less than 0.01. Generated trees were taken 25% burn-in process to calculate posterior probabilities (PPs). The probabilities equal to or greater than 0.9 were displayed at the nodes. The tree was visualized using FigTree version 1.4.4 software [26].

2.5. Pathogenicity test

One isolate from the leaf and stem of *A. acutiloba*, respectively, was used to verify its pathogenicity on leaves and stems of the host plant. Conidial suspension of each isolate harvested from 2-week-old cultures on OA was filtered through two layers of Miracloth (Sigma-Aldrich, St. Louis, MO) and suspended in sterile distilled water. *A. acutiloba* plants were grown in plastic pots (height: 14 cm; upper diameter: 15 cm; lower diameter: 10 cm) with commercial media in a vinyl greenhouse. A 20 ml conidial suspension ($1-2 \times 10^6$ conidia/ml) of each isolate was sprayed onto the leaves of 4-month-old *A. acutiloba* plants after shooting for inoculation test to leaves. A 10 ml conidial suspension ($1-2 \times 10^7$ conidia/ml) of each isolate was sprayed

Table 1. Isolates of *Didymella* spp. and *Coniothyrium palmarum* used for molecular phylogenetic analyses in this study.

Species	Strain/isolate ^a	Host/substrate	Locality	GenBank accession number ^b			
				LSU	ITS	TUB2	RPB2
<i>D. acutilobae</i>	ANAC-1901	<i>Angelica acutiloba</i>	Korea	OQ749982	OQ749980	OQ744070	OQ744072
	ANAC-2001	<i>Angelica acutiloba</i>	Korea	OQ749983	OQ749981	OQ744071	OQ744073
<i>D. brunneospora</i>	CBS 115.58	<i>Chrysanthemum roseum</i>	Germany	KT389723	KT389505	KT389802	KT389625
<i>D. chloroguttulata</i>	CGMCC 3.18351	Air	China	KY742211	KY742057	KY742299	KY742142
<i>D. combreti</i>	CBS 137982	<i>Combretum mossambicense</i>	Zambia	KJ869191	KJ869134	MT005626	MT018139
<i>D. dimorpha</i>	CBS 346.82	<i>Opuntia</i> sp.	Spain	GU238068	GU237835	MT018158	GU237606
<i>D. ellipsoidea</i>	CGMCC 3 18350	Air	China	KY742214	KY742060	KY742145	KY742302
<i>D. exigua</i>	CBS 183.55	<i>Rumex arifolius</i>	France	EU754155	GU237794	GU237525	EU874850
<i>D. gei</i>	CGMCC 3.20068	<i>Geum</i> sp.	China	MT229675	MT229698	MT249266	MT239095
<i>D. infuscatisspora</i>	CGMCC 3.18356	<i>Chrysanthemum indicum</i>	China	KY742221	KY742067	KY742309	KY742152
<i>D. ligulariae</i>	CGMCC 3.20070	<i>Ligularia sibirica</i>	China	MT229676	MT229699	MT249267	MT239096
<i>D. microchlamydospora</i>	CBS 105.95	<i>Eucalyptus</i> sp.	U.K.	GU238104	FJ427028	FJ427138	KP330424
<i>D. pteridis</i>	CBS 379.96	<i>Pteris</i> sp.	The Netherlands	KT389722	KT389504	KT389801	KT389624
<i>D. segeticola</i>	CGMCC 3.17489	<i>Cirsium segetum</i>	China	KP330455	KP330443	KP330399	KP330414
<i>D. subrosea</i>	CBS 733.79	<i>Abies alba</i>	France	MN943747	MN973540	MT005643	MT018174
<i>D. suiyangensis</i>	CGMCC 3.18352	Air	China	KY742243	KY742089	KY742330	KY742168
<i>Coniothyrium palmarum</i>	CBS 400.71	<i>Chamaerops humilis</i>	Italy	EU754153	AY720708	KT389792	KT389592

^aCBS: Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands; CGMCC: China General Microbiological Culture Collection, Beijing, China.

^bLSU: 28S large subunit of the nrRNA gene; ITS: internal transcribed spacer regions 1 and 2 including 5.8S nrDNA gene; TUB2: β -tubulin; RPB2: RNA polymerase II second largest subunit.

onto the stems at the soil surface level of the *A. acutiloba* plants for inoculation test to stems. The inoculated plants were placed in plastic boxes (63 × 44 × 47 cm) under 100% relative humidity at room temperature (24–26 °C). Control plants were sprayed with the same quantity of sterile distilled water and placed under the same conditions as the inoculated plants. After five days, the inoculated plants were taken out from the boxes and kept indoors. The pathogenicity of the isolates was assessed based on the induced symptoms 10–12 days after inoculation. The inoculation test was performed in triplicate.

3. Results

3.1. Molecular phylogeny

According to the model test, the best fitting model for sequences of LSU, ITS, TUB2, and RPB2 were detected as HKY + I, HKY + I + G, GTR + I, and GTR + G, respectively. The partition-homogeneity test (p value = 0.94) confirmed that the phylogenetic trees of each gene have a common underlying structure, which enable us to concatenate the alignments of each gene. The concatenated sequences of the isolates and relevant *Didymella* spp. contained a total 2130 characters (794, 453, 285, and 599 characters of LSU, ITS, TUB2, and RPB2, respectively). Two analyses indicated no significant differences in tree topology (data not shown). Hence, only the NJ tree with BS values was shown with PPs at nodes (BS/PP). The representative isolates, ANAC-1901 and ANAC-2001 obtained from leaf spot and stem rot symptoms, respectively, were determined as members of the genus *Didymella* (Figure 1). The isolates were positioned in a group and verified as the same species because of no difference in their sequences. In addition, the isolates were not

categorized as any species within the genus. Furthermore, a single clade consisting of the isolates was constructed with a high BS value and PP that was clearly separated from other closely related species such as *Didymella bellidis* (Neerg.) Qian Chen & L. Cai (anamorph: *Phoma bellidis* Neerg.) [7,27], *Didymella segeticola* (Q. Chen) Q. Chen, Crous & L. Cai (anamorph: *Phoma segeticola*) [8,28], and *Didymella suiyangensis* Qian Chen, Crous & L. Cai [8]. GenBank accession numbers of the isolates, ANAC-1901 and ANAC-2001 are OQ749982–OQ749983, OQ749980–OQ749981, OQ744070–OQ744071, and OQ744072–OQ744073 for LSU, ITS, TUB2, and RPB2, respectively.

3.2. Taxonomy

Didymella acutilobae G.B. Lee and W.G. Kim, sp. nov. (Figure 2)

Mycobank no.: MB 849458.

Etymology: Named derived from specific epithet of the host plant, *Angelica acutiloba*.

Holotype: Isolated from stem of *A. acutiloba*, Yangpyeong, Gyeonggi Province, Korea (37° 27' 51" N and 127° 33' 16.999" E), April 2020, W.G. Kim, ex-holotype culture (KACC 410302).

Cultural and morphological characteristics: Diameter of 1-week-old cultures on MEA, OA, and PDA was 54–61 mm (av. 58 mm), 53–54 mm (av. 54 mm), and 53–55 mm (av. 53 mm), respectively. The culture on MEA showed brown to black colony with white mycelium and light black concentric rings (Figure 2(A)). The culture on OA showed brown to dark olivaceous colony with white mycelium (Figure 2(B)). The culture on PDA showed white to light brown colony with white mycelium and whitish brown concentric rings (Figure 2(C)). NaOH spot test on MEA was negative.

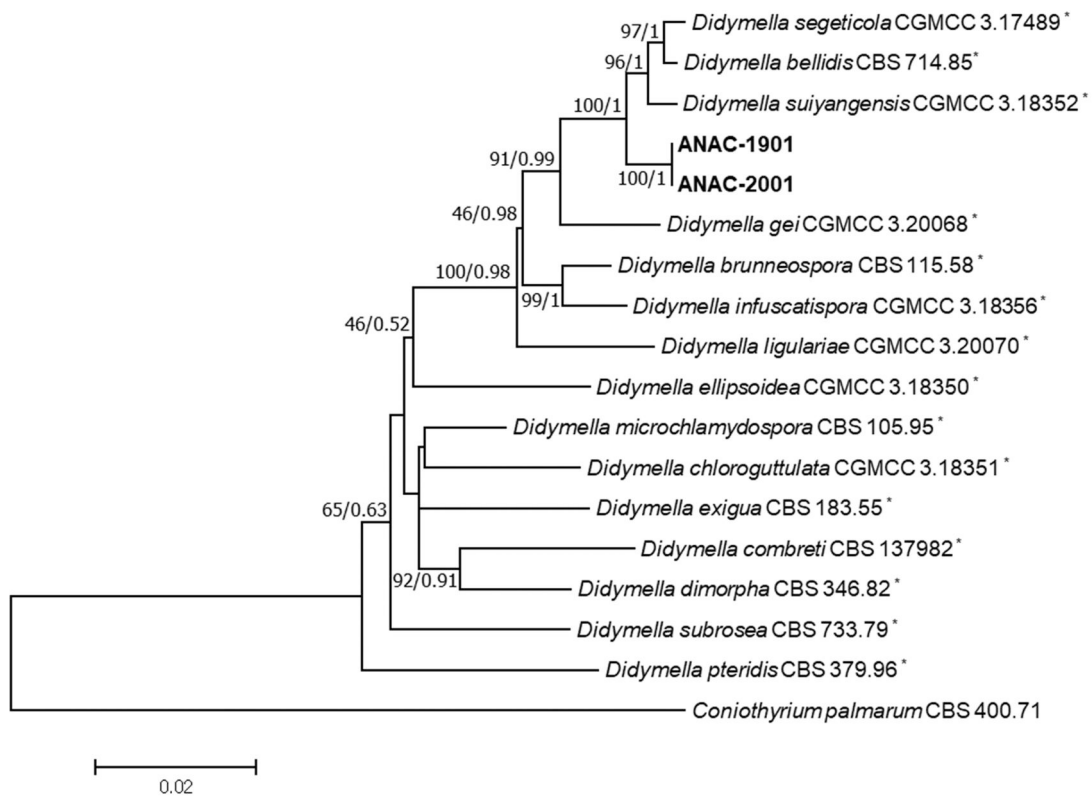


Figure 1. A phylogenetic tree generated from the neighbor-joining analysis with maximum composite likelihood model based on a concatenated alignment of partial large subunit nuclear ribosomal DNA, internal transcribed spacer regions 1 and 2 including 5.8S nrDNA, β -tubulin, and RNA polymerase II second largest subunit sequences of two isolates (ANAC-1901 and ANAC-2001) of *Didymella acutilobae* sp. nov. and relevant *Didymella* spp. Bootstrap values (BS) and posterior probabilities (PP) are given at nodes (BS/PP). The bar represents the number of nucleotide substitutions per site. The phylogenetic tree was rooted to *Coniothyrium palmarum* (CBS 400.71). *The reference strains.

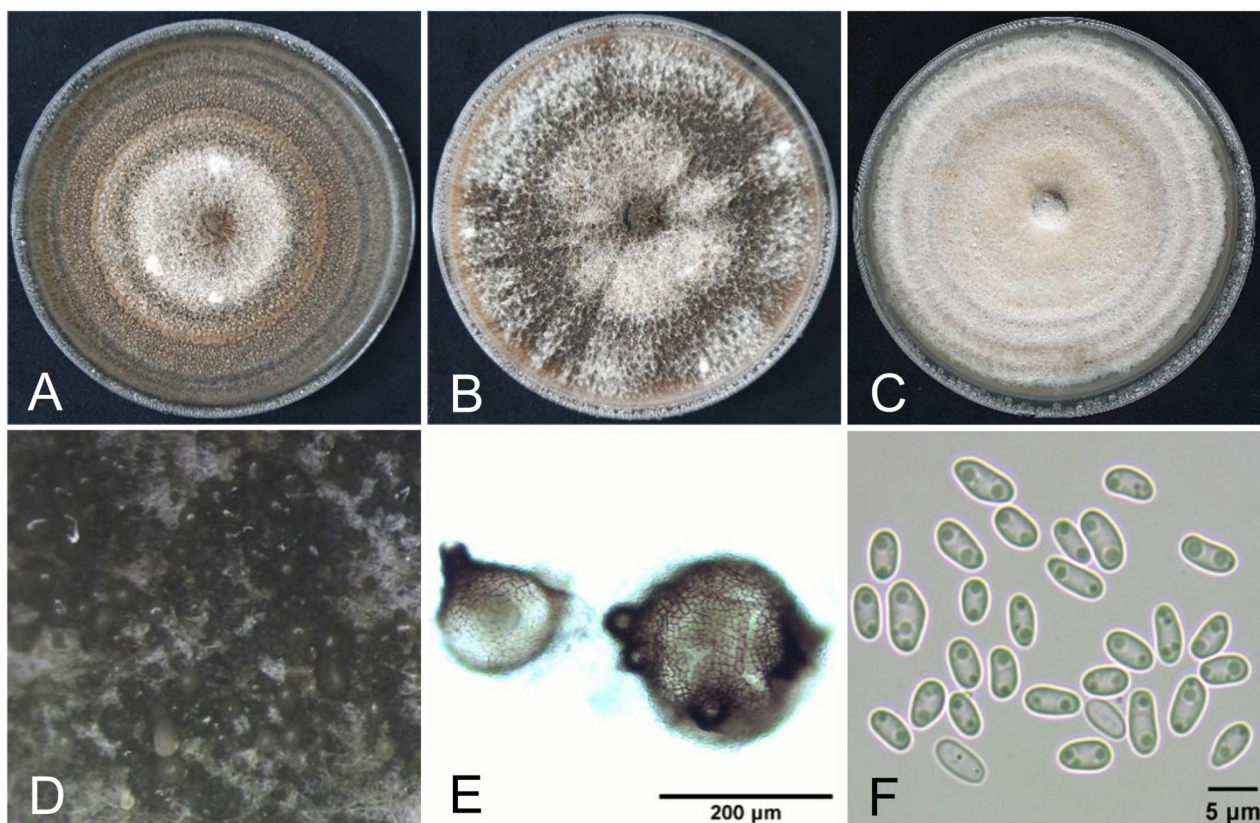


Figure 2. Cultural and morphological features of *Didymella acutilobae* sp. nov. (A) Two-week-old colonies on malt extract agar; (B) oatmeal agar; (C) potato dextrose agar. (D, E) Pycnidia produced in oatmeal agar. (F) Conidia.

Teleomorph was not observed in the cultures. Pycnidia usually half submerged in the agar or on the surface (Figure 2(D)), 70–240 µm in diameter, solitary or confluent, globose, brown to black, with 1–5 ostioles, non-papillate or papillate (Figure 2(E)). Conidia 2.9–6.5 × 1.6–3.0 µm (av. 4.7 × 2.3 µm), ellipsoidal or slightly curved, aseptate with usually two bipolar guttules (Figure 2(F)). Conidial matrix white. Chlamydo spores absent. Summarized descriptions on the cultural and morphological characteristics of *D. acutilobae* and closely related *Didymella* spp. are shown in Table 2. *D. acutilobae* was somewhat dissimilar to the closely related *Didymella* spp. in the cultural and morphological characteristics.

3.3. Disease incidence and pathogenicity

During the disease surveys in Korea, leaf spot symptoms were found on *A. acutiloba* plants in the investigated field in Andong, Gyeongbuk Province in July 2019, and stem rot symptoms in the investigated vinyl greenhouses in Yangpyeong, Gyeonggi Province in April 2020. The leaf spot symptoms were circular to elliptical, brown to dark brown with a yellow halo, and 2–5 mm in diameter in the early stage (Figure 3(A)). In the late stage, they enlarged to irregular shape of 10–20 mm in diameter. The stem rot symptoms showed wilt and blight due to rot of the lower part of the stem (Figure 3(B)).

Incidence of leaf spot and stem rot of the plants during the disease surveys ranged from 10 to 20% and 5 to 30%, respectively.

The isolates (ANAC-1901 and ANAC-2001) from leaf and stem rot lesions of *A. acutiloba* induced leaf spot and stem rot symptoms in the inoculated plants of *A. acutiloba* (Figure 3(C,D)), but no symptoms were observed on the leaves and stems of the control plants (Figure 3(E,F)). The induced symptoms were similar to those observed on the plants of *A. acutiloba* during the disease surveys. Re-isolation of the isolates from the induced leaf spot and stem rot lesions was confirmed based on morphological characteristics.

4. Discussion

According to the phylogenetic analyses, two isolates of *D. acutilobae* were located in the genus *Didymella* with positioning in as an independent group from other closely related species such as *D. bellidis*, *D. segeticola*, and *D. suiyangensis*. Compared to morphology of *D. bellidis*, *D. segeticola*, and *D. suiyangensis* [7,8,27,28], *D. acutilobae* produced much smaller conidia than those of *D. segeticola* and *D. suiyangensis*. *D. acutilobae* developed much bigger pycnidia than *D. segeticola* and showed more ostioles of pycnidia than *D. segeticola* and *D. suiyangensis*. The growth rates of *D. acutilobae* on MEA,

Table 2. Summarized descriptions on the cultural and morphological characteristics of *Didymella acutilobae* and closely related *Didymella* species.

<i>Didymella</i> spp.	Morphological characteristics ^a		Colony on media ^b and result of NaOH spot tests	Reference
	Pycnidia	Conidia		
<i>D. acutilobae</i>	70–240 µm in diameter. Solitary or confluent, globose, brown to black, with 1–5 ostioles, non-papillate or papillate.	2.9–6.5 × 1.6–3.0 µm. Ellipsoidal or slightly curved aseptate with usually 2 bipolar guttules. Conidial matrix white. Chlamydo spores absent.	MEA: brown to black with light concentric rings; 54–61 mm. OA: brown to dark olivaceous; 53–54 mm. PDA: white to light brown with concentric rings; 53–55 mm. NaOH spot test: negative.	Present study
<i>D. bellidis</i> (CBS 714.85)	50–260 µm in diameter. Globose to irregular shape, glabrous, honey to black, with 1–5 ostioles, non-papillate or slightly papillate.	3.8–6.4 × 1.8–2.6 µm. Ellipsoidal, aseptate, with two polar guttules. Conidial matrix salmon to saffron. Chlamydo spores absent.	MEA: olivaceous to grey; 76–77 mm. OA: white to colorless, but salmon color in center; 68 mm. NaOH spot test: positive.	[7,27]
<i>D. segeticola</i> (CGMCC 3.17489)	90–105 × 75–95 µm. Subglobose, glabrous, pyriform and irregular shape in later, 1–2 ostioles, on an elongated neck.	4.5–7 × 2.5–4 µm. Ellipsoidal to ovoid or cylindrical, aseptate with 1–6 polar guttules, conidial matrix crème-white.	MEA: white and green in center; 64–66 mm. OA: white to grey; 56–65.5 mm. PDA: white to grey; 52–59 mm. NaOH spot test: negative.	[8,28]
<i>D. suiyangensis</i> (CGMCC 3.18352)	90–240 × 55–180 µm. Globose to irregular shape, covered by some hyphal outgrowths, brown, 1 ostiole, slightly papillate or papillate.	3.5–7 × 2–3 µm. Ellipsoidal to oblong, smooth, aseptate with indistinct guttules. Conidial matrix cream.	MEA: grey to olivaceous; 59–64 mm. OA: white to buff; 52–55 mm. PDA: White to greyish brown; 57–61 mm. NaOH spot test: positive.	[8]

^aTwenty pycnidia and 30 conidia of each isolate were examined in the present study.

^bDiameter of colony on MEA, OA, and PDA was measured after incubation at 22 °C for one week. Other colony features were investigated after incubation at 22 °C for two weeks. MEA: malt extracted agar; PDA: potato dextrose agar; OA: oatmeal agar.

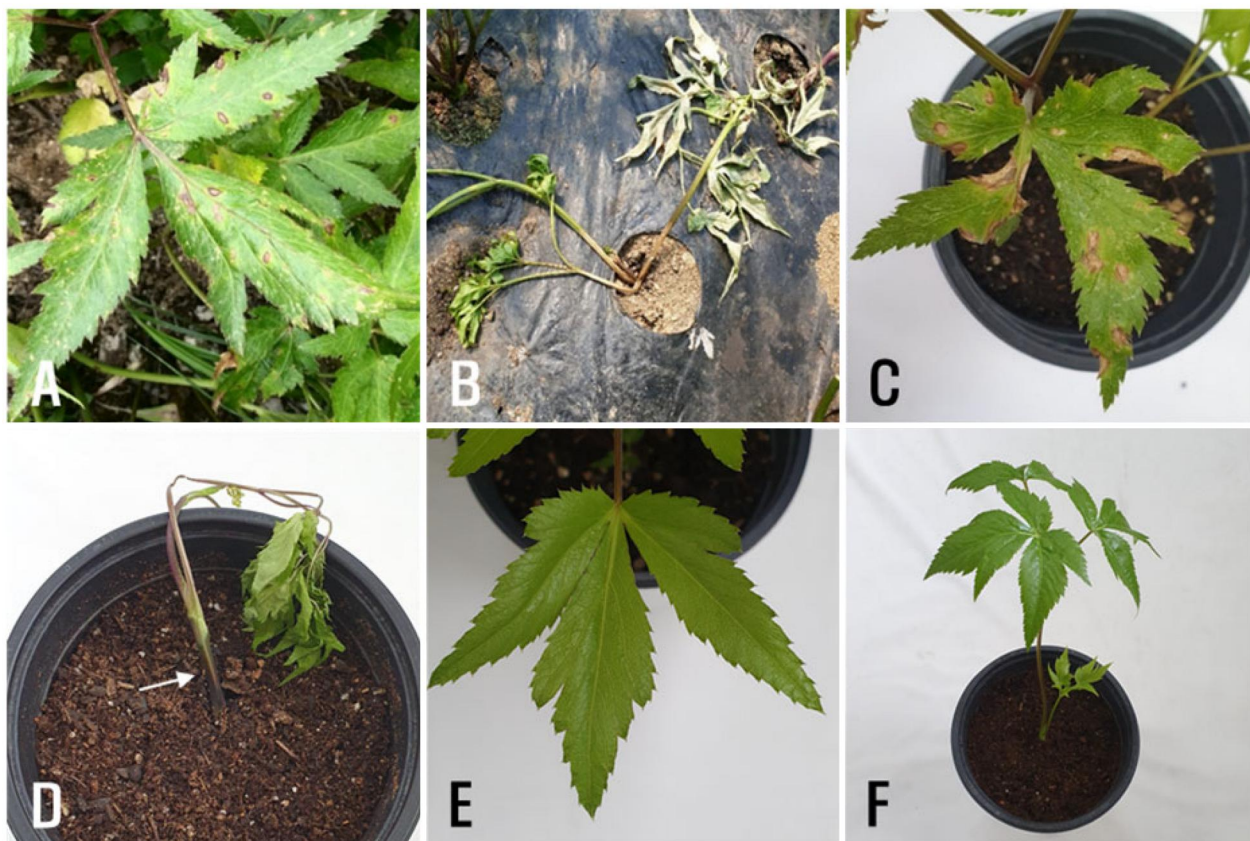


Figure 3. Leaf spot and stem rot symptoms of *Angelica acutiloba* plants. (A, B) Symptoms on the leaves and stems observed in the investigated field and vinyl greenhouse, respectively. Induced symptoms on the leaf (C) and the stem (D) by artificial inoculation with the isolates of *Didymella acutilobae* sp. nov. in pathogenicity tests. The white arrow (D) indicates a stem rot lesion formed on the stem. (E, F) Non-inoculated control plants.

OA, and PDA were generally much slower than those of *D. bellidis*, *D. segeticola*, and *D. suiyangensis*. In NaOH spot test, only *D. acutilobae* and *D. segeticola* exhibited negative reaction among closely related *Didymella* spp. Based on the phylogenetic tree analyses and investigations of the morphological characteristics, it is suggested that the isolates signify a novel species in the genus *Didymella*. Hence, we propose *Didymella acutilobae* sp. nov. as a fungal pathogen causing leaf spot and stem rot in *Angelica acutiloba*.

Disclosure statement

No potential conflict of interest was reported by the authors.

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