RESEARCH NOTE

Erysiphe convolvuli, a Powdery Mildew Occurring on *Convolvulus arvensis* in Korea

In-Young Choi^{1,†}, Young-Eun Na^{2,†}, Lamiya Abasova¹, Joon-Ho Choi¹, and Hyeon-Dong Shin^{1,3,*} ¹Department of Agricultural Biology, Jeonbuk National University, Jeonju 54896, Korea ²National Institute of Agricultural Science, Rural Development Administration, Wanju 55365, Korea ³Division of Environmental Science and Ecological Engineering, Korea University, Seoul 02841, Korea

[†]The first two authors contributed equally to this study. Corresponding author: hdshin@korea.ac.kr

ABSTRACT

Anamorph of a powdery mildew was collected from *Convolvulus arvensis* (Convolvulaceae) in 2021 in Korea. Based on the morphological examination and molecular phylogenetic analyses of the internal transcribed spacer and large subunit rDNA gene, the powdery mildew was identified as *Erysiphe convolvuli*. This powdery mildew species is commonly found in Europe and North America, but rarely in East Asia. The holomorph of this powdery mildew was first recorded on *Calystegia hederacea* in Korea. Conclusively, this is the first report of *E. convolvuli* on *C. arvensis* and the second record of this powdery mildew species in Korea.

Keywords: Anamorph, Calystegia hederacea, Chasmothecia, Field bindweed, Invasive plant

Convolvulus arvensis L., commonly known as field bindweed, is a perennial plant in the family Convolvulaceae. The native origin of this plant is Eurasia, however, in recent centuries, its distribution has expanded to most temperate-subtropical regions. In East Asia, it was introduced into Japan in the 1900s and spread to all areas in the 1940s, while its introduction into Korea was recorded in 1980 along the western coast of the Korean peninsula. Nowadays *C. arvensis* is invasive in all areas of South Korea [1].

In the course of our routine forays to collect phytopathogenic fungi, a previously unknown powdery mildew was found on this plant in Seoul in the summer of 2021 (Fig. 1A). Initially, the infected plants displayed white powdery colonies due to the net-like mycelial mats with an abundance of conidiophores and conidia on both sides of the leaves and stems (Fig. 1B). As the disease progressed, the powdery patches grew larger and denser, causing leaf senescence and early defoliation. No chasmothecia were found until the death of the infected leaves in December 2021.

Detailed morphological characteristics of the powdery mildew were observed using an Olympus BX50 microscope (Olympus, Tokyo, Japan). Photomicrographs were taken with a Zeiss AX10 microscope equipped with an AxioCam MRc5 camera (Carl Zeiss, Oberkochen, Germany). The hyphae were superficial, straight to wavy, branched, and 4-7 μ m wide. Hyphal appressoria were multi-lobed, single or in pairs, and 3-7 μ m wide. Conidiophores are arising from superficial hyphae, solitary, positioned between two hyphal septa usually not central, upright, 54-102×7-9 μ m, and composed of 3-4 cells with



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under the terms of the Creative Commons Attribution Non-Commercial License (http: //creativecommons.org/licenses/by-nc/4.0/) which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited. straight foot cells (Fig. 2A and B). Conidia were formed singly on conidiophores, cylindrical to ellipsoidcylindrical, $34-54 \times 14-18 \mu m$, vacuolate, and devoid of conspicuous fibrosin bodies (Fig. 2C). Germ tubes were produced on the perihilar position of the conidia and were variable in shape (Figs. 2D-I). These characteristics were consistent with the description of *Erysiphe convolvuli* DC. [2,3].

To confirm the morphology-based identification, two specimens (KUS-F32396 and F32411, Korea University Herbarium, Seoul, Korea) were used for molecular-phylogenetic analysis. Genomic DNA was extracted from the mycelium using MagListoTM 5M plant genomic DNA extraction kits (BIONEER, Daejeon, Korea) according to the manufacturer's instructions. The internal transcribed spacer (ITS1 and ITS2) and large subunit (LSU) gene of rDNA were amplified and sequenced with an application of ITS1-F/PM6 and PM3/TW14 primers, respectively [4]. Newly obtained sequences in this study were evaluated using a BLASTN search and deposited in GenBank (Accession Nos. OM033508 and OM033509 for ITS; OM033511 and OM033512 for LSU). The results for ITS and the LSU were 100% identical with sequences from MN203981, MT377709, LC328325 of *Erysiphe convolvuli*. Phylogenetic analysis was conducted in PAUP* 4.0.b using an alignment consisting of 18 sequences, of which 16 were retrieved from GenBank [5]. *Golovinomyces vincae* (AB769444) was used as an outgroup taxon. The robustness of a maximum parsimony tree was evaluated using bootstrap (BS) analysis with the application of 1000 replicates. Sequences from our specimens were grouped with sequences of *Erysiphe convolvuli* in a distinct clade, which is supported by the highest BS value (Fig. 3).



Fig. 1. Powdery mildew caused by *Erysiphe convolvuli* on *Convolvulus arvensis*, which is climbing over hedge-grown *Euonymus japonicus* shrubs on the roadside (A). Close-up view of *E. convolvuli* infection on *C. arvensis* growing over *E. japonicus* (right arrow). Note that young *C. arvensis* leaves are newly infected (left arrow) and uninfected (upper arrow) (B).



Fig. 2. *Erysiphe convolvuli* found on *Convolvulus arvensis*. A-B: Conidiophores. Note the septa (arrows) in the hypha. C: Conidia. D-I: Conidia in germination. Note that germ tubes are short or long, straight or curved, and may or may not have terminal appressoria.



Fig. 3. Maximum parsimony tree based on the combined internal transcribed spacer (ITS)+ large subunit (LSU) sequences of *Erysiphe convolvuli* and other *Erysiphe* spp. retrieved from GenBank. The isolates obtained in this study are shown in bold. Bootstrap values (>80%) are indicated on related branches.

Erysiphe convolvuli has been found on *C. arvensis* plants within its indigenous area, including northern Africa (Canary Islands, Egypt, Libya, and Morocco), western Asia (Afghanistan, western part of China, India, Iran, Iraq, Israel, Lebanon, Pakistan, Saudi Arabia, Azerbaijan, and Turkey), and nearly all of Europe [6]. This powdery mildew species has also been recorded on *C. arvensis* plants in its invasive area, including China (Beijing) and the United States [7]. Interestingly, there have been no reports of this powdery mildew in Japan [8], suggesting the absence of *E. convolvuli* in Japan, where five introduced species of *Convolvulus* plants have been found (http://www.ylist.info). Likewise, *E. convolvuli* was recently found on *Calystegia hederacea* in Korea [3]. This is the first report of powdery mildew caused by *E. convolvuli* on *C. arvensis* in Korea. This finding could help understand recent invasions and the spread of this powdery mildew.

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