Morphological and Molecular Identification of *Penicillium islandicum* Isolate KU101 from Stored Rice

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(Received on September 11, 2008; Accepted on November 6, 2008)

We have previously obtained a representative isolate KU101 of the predominant *Penicillium* species from rice under indoor storage conditions. In this study we attempted to characterize isolate KU101 using its morphological and molecular characteristics. When the micro- and macroscopic characteristics of isolate KU101 were compared with the *P. islandicum* reference isolate KCCM 34763, isolate KU101 was generally identical to those of isolate KCCM 34763, however, isolate KU101 grew faster and produced more orange to red pigments than isolate KCCM 34763. In a molecular-based identification, the nuclear sequence of the ITS1-5.8S-ITS2 region of isolate KU101 was most closely related to that of *P. islandicum*. Therefore, these results indicated that isolate KU101 from stored rice could be identified as *P. islandicum*, some isolates of which are known to produce mycotoxins.

**Keywords**: identification, *Penicillium islandicum*, stored rice

*Penicillium islandicum* Sopp. is one of the most destructive and harmful fungi affecting rice in storage. In Korea, imported rice infected with *P. islandicum* was detected in 1973 (Kim et al., 1973). Park et al. (2005) also described the existence of *P. islandicum* in 15 out of 45 Korean polished rice samples. In Japan, Sakai et al. (2005) reported that three out of 100 samples of domestic rice harvested in 2001 and 2002 were contaminated with *P. islandicum*, and among the contaminated samples, 82% of rice grain in a sample was infected. The contamination of paddy and milled rice by *P. islandicum* also occurred in Argentina and Paraguay (Tonon et al., 1997).

In our previous study (Oh et al., 2007), we examined the populations of fungi and bacteria in stored rice from rice processing complexes of the National Agricultural Cooperative Federation of 11 regions in Korea and found that all rice samples were contaminated with *Aspergillus* and *Penicillium* spp. including a species examined in this study. Furthermore, we examined rice under controlled condition to monitor the temporal changes of fungal and bacterial population and diversity (Oh et al., 2008). Consequently, we found that various *Penicillium* spp. existed in stored rice and that changes in the conditions tested had little impact on the types of species occurred. However, the population of one (representative isolate KU101) of the most frequently appearing *Penicillium* species dramatically increased up to about 40-70% of total fungi in brown rice as the relative humidity was increased. Thus, the objective of this study was to identify the predominant species of *Penicillium* occurring in the stored rice using morphological and molecular characteristics.

A representative isolate KU101 of the predominant *Penicillium* species was obtained from rice samples in one ton bags being stored indoor condition at Korea University, Seoul, Korea in 2006. The morphological identification of isolate KU101 was conducted using eight different types of media and compared to the *P. islandicum* reference isolate KCCM 34763 from the Korean Culture Center of Microorganisms (KCCM), Seoul, Korea. The media used (Frisvad and Samson, 2004; Pitt and Hocking, 1999) were: 1) czapek yeast extract agar (CYA), 2) czapek yeast autolysate with 5% NaCl (CYAS) agar, 3) czapek dox agar (CZ), 4) malt extract agar (MEA), 5) dichloran 18% glycerol agar (DG18), 6) 25% glycerol nitrate agar (G25N) to test the ability of isolate to grow at low a⁰, 7) yeast extract sucrose agar (YES), and 8) creatine sucrose agar (CREA) which shows acid or base production by fungi. All media were placed at 25°C for 7 days when comparing the growth rates at different temperatures using CYAs incubated at 15, 25, and 30°C. Colonies on each medium were compared for their diameters, overall colors, colors of conidia, reverse colors, and productions of sclerotia, exudates, and soluble pigments. Ehrlich reactions for detecting indole metabolites were conducted by the method of Lund (1995). Agar plugs (4 mm in diameter) from the center of a 7-day-old colony grown on CYA were cut and covered with discs (1 cm in diameter) of Whatman filter paper (No. 1) wetted with Ehrlich reagent (4-dimethylaminobenzaldehyde 2 g, 96%...
ethanol 85 ml, 37% hydrochloric acid 15 ml). The reaction was observed 5 and 10 min after treatments. The fungal growth (mm) on the media was determined with five plates (replications) and this experiment was repeated with similar results.

Microscopic features of isolates KU101 and KCCM 34763 on MEA, such as conidial heads, stipes, shapes of conidia, roughness of conidial walls, existence of ascospores or cleistothecia, lengths of phialides, and metulae, were determined using a microscope (× 1,000); these were compared with those of the *P. islandicum* described in the literature (Pitt, 2000). The branching patterns and appressedness of conidiophores were also observed to determine if they are mono-, bi-, or ter-verticillate and whether or not they are pressed close.

For molecular identification of isolate KU101, mycelial DNA was extracted from fungal cultures grown in potato dextrose broth at 28°C for 7 days. Extraction of DNA was conducted with the modified method of Boom et al. (1990). Polymerase chain reactions (PCR) were performed using oligonucleotide primer pairs, ITS1 (5'-TCCGTAGGTTGAA-CCTGCGG-3') and ITS4 (5'-TCTTCCGCTTATTGATA-TGC-3') (White et al., 1990), to amplify the nuclear ribosomal ITS1-5.8S-ITS2 region of *Penicillium* spp. The PCR contained the followings: 100 µl of each reaction mixture containing template 10 µl (10 ng/µl), forward primer (ITS1) 10 µl (20 ng/µl), reverse primer (ITS4) 10 µl (20 ng/µl), 2.5 mM dNTP 10 µl, 10x reaction buffer 10 µl, Taq 1 µl (5 unit/µl), and distilled water 49 µl, and the cycling conditions were 1) an initial denaturation of 4 min at 95°C, 2) 35 cycles of denaturation at 95°C for 30 s, annealing at 52°C for 1 min, and extension at 72°C for 1 min, followed by 3) a final extension at 72°C for 7 min. Following amplification, the PCR products were run on gels composed of 1.5% (w/v) agarose (Roche Diagnostics, Indianapolis, USA) in TAE buffer [0.045 M Tris, 0.089 M boric acid, 0.002 M EDTA (pH 8.4)] at 100 V for 120 to 150 min. The sequences of the ITS1-5.8S-ITS2 region of closely-related type strains of *Penicillium* species were obtained by Blast analysis against the GenBank DNA databases. Multiple sequence alignments were carried out using the ClustalX multiple sequence alignment program (European Molecular Biology Laboratory, Heidelberg, Germany). Phylogenetic and molecular evolutionary analyses were conducted using the Molecular Evolutionary Genetics Analysis (MEGA) version 3.1 program (The Biodesign Institute, Tempe, USA).

The 16S rDNA sequence of isolate KU101 has been deposited in GenBank under accession number FJ197220.

Colonies of isolate KU101 on CYA at 25 and 30°C were slightly sulcate, centrally rose, and velutinous to lightly floccose (Fig. 1A). Diameters of the colonies on CYA were, on average, 10.7 mm at 15°C, 23.9 mm at 25°C, and 30.2 mm at 30°C. Colors of mycelia on CYA were deep orange covering the central part of colony and grayish turquoise on the conidia especially in the margin of a colony. Conidio genesis was moderate and usually covered with orange to red-brown mycelia. Sometimes clear to pale yellow exudates were produced. Colonies on CYAS were, on average, 22.9 mm in diameter and similar to those on CYA except that their mycelia were deep orange. Colonies on CZ were, on average, 19.9 mm in diameter and slightly sulcate. Their mycelia were yellow, conidia grayish green, and conidio genesis was moderate. The reverse colors on CYA, CYAS, and CZ were strongly pigmented with an orangish brown to

![Fig. 1](image-url). Morphologies of (A) the *Penicillium islandicum* isolate KU101 and (B) the *P. islandicum* reference isolate KCCM 34763 on different media such as czapek yeast extract agar grown at 15°C (CYA15), CYA at 25°C (CYA), CYA at 30°C (CYA30), czapek yeast extract agar with 5% NaCl (CYAS), czapek dox agar (CZ), malt extract agar (MEA), dichloran 18% glycerol agar (DG18), 25% glycerol nitrate agar (G25N), yeast extract sucrose agar (YES), and creatine sucrose agar (CREA) at 25°C. O, obverse of the culture plate; R, reverse.