A Short-chain Dehydrogenase/reductase Gene is Required for Infection-related Development and Pathogenicity in *Magnaporthe oryzae*

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The phytopathogenic fungus *Magnaporthe oryzae* is a major limiting factor in rice production. To understand the genetic basis of *M. oryzae* pathogenic development, we previously analyzed a library of T-DNA insertional mutants of *M. oryzae*, and identified ATMT0879A1 as one of the pathogenicity-defective mutants. Molecular analyses and database searches revealed that a single T-DNA insertion in ATMT0879A1 resulted in functional interference with an annotated gene, MGG00056, which encodes a short-chain dehydrogenase/reductase (SDR). The mutant and annotated gene were designated as *Mosdr1* and *MoSDR1*, respectively. Like other SDR family members, *MoSDR1* possesses both a cofactor-binding motif and a catalytic site. The expression pattern of *MoSDR1* suggests that the gene is associated with pathogenicity and plays an important role in *M. oryzae* development. To understand the roles of *MoSDR1*, the deletion mutant Δ*Mosdr1* for the gene was obtained via homology-dependent gene replacement. As expected, Δ*Mosdr1* was nonpathogenic; moreover, the mutant displayed pleiotropic defects in conidiation, conidial germination, appressorium formation, penetration, and growth inside host tissues. These results suggest that *MoSDR1* functions as a key metabolic enzyme in the regulation of development and pathogenicity in *M. oryzae*.

**Keywords**: appressorium, conidiation, rice blast, short-chain dehydrogenase/reductase (SDR)

*Magnaporthe oryzae* is an ascomyceteous phytopathogen that causes rice blast, one of the most destructive diseases found in all rice-growing countries (Ou, 1985). The disease continues to be a major constraint in rice production, and it is exacerbated by such factors as global population growth. Development of the disease requires appropriate fungal development and pathogenicity, which are controlled by complicated cellular processes. Disease occurs when the conidium (asexual spore) attaches to suitable host tissues. After conidial attachment, a germ tube emerges at the apex that differentiates into a specialized infectious structure, called an appressorium, upon its perception of host factors. Enormous internal turgor pressure (>8 MPa) is generated inside the appressorium, which enables the fungus to penetrate host cells through mechanical disruption of the host’s epidermal barriers (Howard et al., 1991). The fungus then develops invasive hyphae in the host cell membrane, colonizes whole cells, and escapes via conidiation.

*M. oryzae* has emerged as an important model organism for studying the molecular basis of fungal development and pathogenicity due to its genetic amenability, genomic sequence data, and the severe economic impact of rice blast (Dean et al., 2005; Valent, 1990). Several studies have shown that conserved core signaling pathways are associated with the transduction of signals into cellular responses for appressorium-mediated disease development. These include cyclic adenosine monophosphate (cAMP) (Choi and Dean, 1997; Mitchell and Dean, 1995), mitogen-activated protein (MAP) kinase (Jeon et al., 2008; Xu and Hamer, 1996; Xu et al., 1997), and Ca²⁺-dependent signaling pathways (Choi et al., 2009a; Choi et al., 2009b). Whereas the effects of conserved signaling pathways on *M. oryzae* development and pathogenicity are well documented, little is known of the other determinants and pathways that direct infection-related development and pathogenicity. Although successful disease development by *M. oryzae* involves intercellular signaling as well as metabolic and regulatory networks that are highly integrated, most studies have focused on signaling pathways without considering the interplay that exists between them.

Members of the short-chain dehydrogenase/reductase (SDR) family are present in all forms of life, from simple organisms to higher eukaryotes (Jörnvall et al., 1999), indicating their versatility and fundamental importance in metabolic processes (Persson et al., 2009). Most dehydrogenases (about 25%) belong to the SDR family (Kallberg and Persson, 2006). As the first characterized member of the SDR family was *Drosophila* alcohol dehydrogenase, this family used to be called “insect-type” or “short-chain”
alcohol dehydrogenase (Villatorrya et al., 1989). SDRs are also known as NAD(P)H-dependent oxidoreductases, which are distinct from the medium-chain dehydrogenase/reductase (MDR) and aldo-keto reductase (AKR) superfamilies (Jönnvall et al., 1995). There are two conserved motifs in the SDR family. The first motif at the N-terminus defines the cofactor-binding site (TGxxxxGxG). This region consists of alternating β-sheets and α-helices (β-α-β), homologous to Rossmann-fold elements with a typical α/β-folding pattern (Oppermann et al., 2001; Rossmann and Argos, 1978). The second region houses specific residues for the catalytic site (YxxxxK), which is not present in the MDR or AKR superfamily (Jönnvall et al., 1995).

SDRs play diverse and important roles in cellular differentiation and signaling. SDRs are involved in retinoid and steroid hormone biosynthesis in mammals (Oppermann et al., 2001) and sex determination in maize by controlling cell death (DeLong et al., 1993). Other SDRs have been found to convert xanthoxin to abscisic acid (ABA)-aldehyde in ABA biosynthesis (Cheng et al., 2002), which is critical for plant growth, development, and stress responses. Members of the SDR family such as trihydroxynaphthalene reductase (3HNR) are key enzymes in fungal melanin biosynthesis (Thompson et al., 1997), which is required for pathogenicity in *M. oryzae*. Also, an SDR has been shown to be involved in the biosynthesis of fumonisin toxin in *Gibberella moniliformis* (Butchko et al., 2003).

To better understand *M. oryzae* development and pathogenicity, we previously generated a genome-wide mutant library via *Agrobacterium tumefaciens*-mediated transformation (ATMT) and screened mutants to find genes required for the disease development (Jeon et al., 2007). In this study, we characterized one of pathogenicity defective T-DNA insertional mutants, ATMT0879A1, that is renamed MoSDR1. Deletion mutants of MoSDR1 were generated through homology-dependent gene replacement to confirm the involvement of MoSDR1 in pathogenicity. Analyses of the ΔMosdr1 mutants revealed that MoSDR1 is essential for conidiation, conidial germination, and appressorium development.

### Materials and Methods

**Fungal strains and culture conditions.** *M. oryzae* strain KJ201, which was obtained from the Center for Fungal Genetic Resources (CFGR; http://cfgr.snu.ac.kr), was used as the wild-type strain. The wild-type strain and transformants generated in this study were cultured on either oatmeal agar medium (OMA; 5% oatmeal (w/v) and 1.5% agar) or V8 juice agar medium (4% V8 juice and 1.5% agar) at 25°C under constant fluorescent light to promote conidiation (Lee and Lee, 1998). Genomic DNA and total RNA were extracted from four-day-old mycelia cultured in complete liquid medium (0.6% yeast extract, 0.6% tryptone, and 1% sucrose) as described previously (Chi et al., 2009a).

**Measurement of mycelial growth, conidiation, conidial germination, and appressorium formation.** Mycelial growth was measured in six-well plates containing V8 agar (SPL Lifesciences Inc., Gyeonggi-Do, Korea) five days after inoculation with a mycelia agar plug (4 mm in diameter). The degree of conidiation was measured from ten-day-old oatmeal agar cultures in six-well plates. Conidia were collected with 5 ml of distilled water and counted using a hemacytometer under a light microscope. Counting was done in three independent experiments with triplicates. Conidial germination and appressorium formation were measured on the hydrophobic side of a coverslip or GelBond film (FM BioProducts, Rockland, ME, USA) as described previously (Lee and Dean, 1993). Conidia were harvested from six-day-old oatmeal agar cultures with sterile distilled water and adjusted to a concentration of 10⁶ conidia/ml. Drops (40 μl of the conidial suspension) were placed on the hydrophobic side of the coverslips and incubated in a box containing pre-moistened wipes at 25°C for 8 h. The percentages of conidial germination and appressorium formation from germinated conidia were determined by counting more than 100 conidia in at least four independent experiments with triplicate samples under a microscopy.

**Pathogenicity assays.** Conidia were harvested from ten- to twelve-day-old cultures on oatmeal agar plates with sterilized distilled water. A conidial suspension (1×10⁶ conidia/ml) containing 250 ppm Tween 20 was used for the spray inoculation of seedlings of a susceptible rice cultivar (*Oryza sativa* cv. Nakdong). The inoculated plants were placed in a moisture chamber at 25°C for 24 h in the dark, and then transferred to a growth chamber with one photoperiod of 16/8-h (light/dark). Disease severity was assessed seven days after inoculation based on the diseased leaf area (DLA) and lesion types, as described previously (Koh, 1986). For the infiltration assay, 100 μl of a conidial suspension (5×10⁶ conidia/ml) was injected using a syringe without a needle onto the leaves of three-week-old rice plants. Disease symptoms were observed at seven days after inoculation. For the sheath infection assay, a spore suspension (10⁵/ml in 0.25% gelatin) was placed under the leaf sheaths of five-week-old rice plants. The incubated plants were covered with a plastic bag to maintain high humidity and laid down horizontally inside. At 48 h after inoculation, the sheaths were trimmed to produce epidermal layers above the midvein and used for microscopic obser-
vation (Chi et al., 2009b; Koga et al., 2004). The assays for pathogenicity were repeated three times using triplicate samples.

**Nucleic acid manipulation and polymerase chain reaction (PCR).** Genomic DNA was isolated as described previously (Kim et al., 2005). Restriction enzyme digestion, agarose gel fractionation, cloning, and DNA gel blotyping were performed using standard methods (Sambrook et al., 1989). Briefly, fungal genomic DNA was digested with ScaI, separated on a 0.7% agarose gel, and transferred onto a Hybond N+ membrane (Amersham Pharmacia Biotech, Little Chalfont, UK). The probes used for DNA hybridization were labeled with $^{32}\text{P}$-dCTP using the Rediprime II Random Prime Labeling System (Amersham Pharmacia Biotech, Piscataway, NJ, USA) according to the manufacturer’s instructions. The hybridization membrane was exposed to a phoshorimager (BAS-2040, Fuji Photo Film, Tokyo, Japan) and visualized using a phosphorimager software. Total RNA was isolated from the mycelia using an Easy-Spin RNA extraction kit (iNtRON Biotechnology, Seoul, Korea). For reverse transcription-PCR (RT-PCR), 3 μg of total RNA were reverse-transcribed into first-strand cDNA using oligo (dT) primers and the SuperScript First-Strand Synthesis System (Invitrogen Life Technologies, Carlsbad, CA, USA) according to the manufacturer’s instructions. RT-PCR was performed on a Perkin-Elmer 9600 DNA thermal cycler in a 20-μl reaction mixture containing 1 μl of cDNA, 1 μl of deoxyribonucleoside triphosphates (2.5 mM dNTP mix), 2 μl of 10× PCR buffer, 100 nM each primer, and 1 U of Taq polymerase. Real-time quantitative RT-PCR (qRT-PCR) was performed as described previously (Kim et al., 2005) using the AB7500 Real-Time PCR system (Applied Biosystems, Foster City, CA, USA). The conditions included 3 min at 95°C (1 cycle) followed by 15 s at 95°C, 30 s at 60°C, and 30 s at 72°C (40 cycles). Each qRT-PCR mixture (final volume 10 μl) contained 5 μl of Power SYBR® Green PCR Master Mix (Applied Biosystems), 3 μl of forward and reverse primers (100 nM each), and 2 μl of cDNA (12.5 ng/μl). The oligonucleotide sequences (SDR1-RT for RT-PCR and SDR1-Q for qRT-PCR) are listed in Table 1. To compare the relative abundance of target gene transcripts, the average threshold cycle (Ct) was normalized to that of $\beta$-tubulin for the samples as follows: $2^{-\triangle \text{Ct}}$, where $-\Delta \text{Ct} = (C_{\text{target gene}} - C_{\beta\text{-tubulin}})$. The fold-change for a target gene during fungal development, as compared to its growth in liquid complete medium, was calculated as $2^{\Delta \text{Ct}}$, where $-\Delta \text{Ct} = (C_{\text{target gene}} - C_{\beta\text{-tubulin}})_{\text{liquid complete medium}} - (C_{\text{target gene}} - C_{\beta\text{-tubulin}})_{\text{liquid complete medium}}$ (Livak and Schmittgen, 2001). qRT-PCR was performed twice using three independent tissue pools. The flanking region of the T-DNA was obtained by thermal asymmetric interlaced-PCR (TAIL-PCR) as described previously (Liu and Whittier, 1995).

**Targeted disruption of MoSDR1.** The targeted gene disruption vector for MoSDR1 was constructed by double-joint PCR as described previously (Gritz and Davis, 1983; Yu et al., 2004). Fragments of the 5' (1,263 bp) and 3'-flanking regions (1,264 bp) of the gene were amplified using the primers UF/UR and DF/DR, respectively (Table 1). A 2.1-kb hygromycin phosphotransferase gene (HPH) cassette was amplified using primers HPHF and HPHR (Table 1) from pBCATPH (Gritz and Davies, 1983). The replacement construct was amplified with primers UF/DR using the fused product as the template. Protoplasts from strain KJ201 were transformed directly with the purified deletion construct after amplification. Hygromycin-resistant transformants were selected on TB3 (0.3% yeast extract, 0.3% casamino acid, 1% glucose, and 20% sucrose) medium supplemented with 200 μg/ml hygromycin B (Calbiochem, San Diego, CA, USA) and screened by PCR. Putative two knock-out mutants were confirmed by DNA blotyping and RT-PCR. Since the mutants were phenotypically indistinguishable, a representative mutant named as ΔMosdr1 was used in this study.

**FUN-1 staining.** Conidial viability was examined by staining with the fluorescent dye FUN-1 (Molecular Probes, Eugene, OR, USA). A conidial suspension (1×10$^6$ conidia/ml in 1% glucose, 10 mM Na-HEPES, and 20 μM FUN-1, pH 7.2) was incubated at 30°C in the dark for 60 min, then observed under optics equipped with epifluorescence (Millard et al., 1997). Stained conidia were viewed and photographed with excitation at 450-490 nm and emission at 520 nm under an Axioscan Universal microscope (Carl Zeiss Microscope Division, Oberkochen, Germany) equipped with a fluorescein filter set.

**Results**

**MoSDR1-DNA is a pathogenicity-defective T-DNA mutant of M. oryzae.** We previously established a T-DNA insertional library via ATMT, which generated 21,070 hygromycin-resistant mutants (Jeon et al., 2007). ATMT0879A1, a pathogenicity-defective mutant, was identified using a high-throughput screening system. As shown in Fig. 1A, ATMT0879A1, which is referred as MoSDR1-DNA, was nearly nonpathogenic; it produced a few tiny lesions that failed to develop further, whereas the wild-type strain caused typical symptoms of rice blast on the susceptible rice cultivar Nakdong. Assessment of pathogenicity according to the method of Koh et al. (1987) revealed that MoSDR1-DNA significantly reduced the disease severity compared to wild-
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Southern blot analysis revealed that integration of a single copy of the T-DNA occurred in MoSDR1TDNA (data not shown). The right flanking sequence of the T-DNA was determined by TAIL-PCR as described previously (Choi et al., 2007; Liu and Whittier, 1995). The results indicated that the T-DNA was inserted into the promoter region of an annotated gene, MGG00056 in supercontig 6.21 of the genome sequence. These data allowed us to create a schematic diagram of the integration of the T-DNA in ATMT0879A1 (Fig. 2A). RT-PCR revealed that the expression level of MGG00056 was reduced by two-fold in MoSDR1TDNA, whereas the expression level of MGG11682 upstream of MGG00056 was unaffected compared to that in the wild-type strain (data not shown). These data suggest that the pathogenicity defect of MoSDR1TDNA results from functional interference with the annotated gene MGG00056.

**Fig. 1.** Pathogenicity test of the *M. oryzae* mutants MoSDR1TDNA and Δ Mosdr1. (A) Disease development in rice. Seedlings of a susceptible rice cultivar (cv. Nakdong) were inoculated with conidia (1×10^6/ml) of the indicated strains. Photos of the inoculated leaves were taken seven days after spray inoculation. (B) Disease severity was measured based on the diseased leaf area and lesion types. The values given are the means ± SD from three replicates inoculated by each strain. Three rice plants were used for each replicate.

**Fig. 2.** Schematic diagram of the T-DNA insertion in ATMT0879A1 and the targeted deletion of MoSDR1. (A) Schematic diagram of the T-DNA insertion in ATMT0879A1. The insertion is located -271 bp from the start codon in MGG00056. (B) Schematic diagram showing the targeted deletion of MoSDR1. The MoSDR1 replacement vector contained the homologous sequences flanking hph for homology-dependent replacement. S and closed arrows indicate the SacI restriction site and primer positions, respectively. (C) Genomic DNA was isolated from wild-type (lane 1), Δ Mosdr1 (lane 2), and an ectopic transformant (lane 3), digested with SacI, and probed with the 3′-flanking region. (D) MoSDR1 expression in the wild-type and Δ Mosdr1. Total RNA was isolated from the mycelia of wild-type (lane 1) and Δ Mosdr1 cells (lane 2), then subjected to RT-PCR.

**MGG00056 encodes a member of the SDR family.** A comparison of the cDNA sequence of MGG00056 with the genomic sequence confirmed that the gene consists of two exons and one intron and encodes a protein consisting of 286 amino acids with a predicted molecular mass of 29 kDa. MGG00056 was found to encode an SDR from a series of database searches, including BLASTP and PSORT (http://db.psort.org); it is hereafter referred to as MoSDR1. MoSDR1 showed significant similarity to predicted proteins from filamentous fungi, including 73 and 72% identity with proteins from *Aspergillus oryzae* and *Penicillium chrysogenum*, respectively (Fig. 3). MoSDR1 possesses an atypical NAD/NADH-binding motif (TGxxxxGxG instead of TGxxxxGxG) and a typical catalytic site (YxxS/T-A/K), similar to SDR family members found in prokaryotic and eukaryotic organisms. The variability detected in the binding motif of MoSDR1 has been reported in other homologs.
The SDR family includes classical and extended types that differ in terms of their lengths and cofactor-binding motifs (Kallberg et al., 2002). Targeted deletion of MoSDR1 reveals its role in disease development. To determine the function of MoSDR1, the coding region of MoSDR1 was replaced with a hygromycin B expression cassette via a homology-dependent replacement strategy (Fig. 2B). Purified MoSDR1 deletion constructs after amplification were used to transform fungal protoplasts. Hygromycin-resistant transformants were selected on a regeneration medium containing 200 μg/ml hygromycin B and screened by a PCR using the primers DR/HPHF (Table 1). An MoSDR1 deletion mutant, ΔMosdr1, was confirmed by Southern blot analysis and RT-PCR (Fig. 2C and D). We first compared the pathogenicity of ΔMosdr1 with that of MoSDR1 T-DNA. As expected, ΔMosdr1 showed remarkably reduced pathogenicity compared to wild-type, similar to MoSDR1 T-DNA (Fig. 1). This result indicates that MoSDR1 is required for M. oryzae pathogenicity.

MoSDR1 is required for conidial development. Quantitative RT-PCR revealed that MoSDR1 mRNAs were highly expressed in conidia (>3-fold), appressoria (>2-fold), and in planta (>3-fold), relative to those in mycelia. To characterize the roles of MoSDR1 during M. oryzae development, we performed functional analyses using a ΔMosdr1 mutant.
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Development, phenotypic characterization of the strains was carried out. Measurements of mycelial growth showed that ΔMosdr1 was slightly retarded compared to wild-type on complete medium, similar to MoSDR1T-DNA (Table 2). The morphologies of the colonies of the two strains were indistinguishable (data not shown). Quantitative measurement of the conidia indicated that conidial production was significantly reduced in MoSDR1T-DNA. As expected, the deletion of MoSDR1 resulted in a significant reduction in conidiation as shown in MoSDR1T-DNA. These results indicate that MoSDR1 plays a role in asexual (conidial) reproduction.

MoSDR1 is indispensable for preinfection-related development and pathogenicity. The germination of MoSDR1T-DNA conidia was significantly reduced, whereas about 98% of wild-type conidia germinated (Table 2). Therefore, we tested whether the deletion of MoSDR1 affects this stage of preinfection development. A large proportion (>80%) of ΔMosdr1 conidia failed to germinate on a hydrophobic surface (Table 2). After conidial germination, the wild-type strain formed dome-shaped appressoria at the ends of the germ tubes after 6 h of incubation on a hydrophobic surface. More than 95% of the wild-type conidia formed appressoria (Table 2); however, appressorium development in both ΔMosdr1 and MoSDR1T-DNA was significantly reduced.

To determine whether the ΔMosdr1 mutant exhibited invasive growth in plant cells, a conidial suspension was infiltrated into rice leaves by injection using a syringe. The wild-type strain produced blast lesions, whereas ΔMosdr1 failed to cause the disease. These data indicate that the mutant lost the ability to grow inside plant cells (Fig. 4A).

To test the role of MoSDR1 in penetration, rice leaf sheath cells were inoculated with a conidial suspension. The wild-type strain penetrated the epidermal cells of the leaves and grew invasively (Fig. 4B). In contrast, the ΔMosdr1 mutant was unable to penetrate the plant surface, showing hyphal growth on all surfaces. These results suggest that the functioning of MoSDR1 is required for appressorium-mediated penetration and further growth inside the host.

**MoSDR1 is required for conidial viability.** To determine whether the inability of the ΔMosdr1 conidia to germinate was associated with conidial viability, we stained the conidia with the dye FUN-1. Briefly, conidia were freshly harvested in a solution (2% glucose and 10 mM HEPES, pH 7.2) from seven-day-old oatmeal agar plates, and incubated at 30°C for 60 min in dark. Most of the wild-type conidia (>97%) exhibited orange fluorescence, indicating that they were metabolically active; in contrast, most of the

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**Table 2. Comparison of phenotypic characteristics among the strains**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Mycelial growth (mm)</th>
<th>Conidiation (10⁵/ml)</th>
<th>Conidial germination (%)</th>
<th>Appressorium formation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>3.1 ± 0.1</td>
<td>17.4 ± 1.1</td>
<td>98.0 ± 1.0</td>
<td>96.4 ± 1.5</td>
</tr>
<tr>
<td>MoSDR1T-DNA</td>
<td>2.7 ± 0.2</td>
<td>0.6 ± 0.1</td>
<td>15.4 ± 0.5</td>
<td>3.2 ± 0.8</td>
</tr>
<tr>
<td>ΔMosdr1</td>
<td>2.6 ± 0.1</td>
<td>0.7 ± 0.2</td>
<td>17.4 ± 0.8</td>
<td>4.3 ± 1.5</td>
</tr>
</tbody>
</table>

*a* Mycelial growth was measured at five days post-inoculation on V8 agar medium. The data are presented as the means ± SD of three independent experiments with triplicates.

*b* Conidiation was measured by counting the number of conidia collected from ten-day-old oatmeal agar cultures in six-well plates.

The percentage of conidial germination on a hydrophobic surface was measured under a light microscope using conidia harvested from six-day-old oatmeal agar plates.

The percentage of appressorium formation from germinated conidia on a hydrophobic surface was measured using conidia harvested from six-day-old oatmeal agar plates.

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**Fig. 4. In planta assays for fungal growth and penetration.** (A) Infiltration assay for invasive growth inside plant cells. Conidial suspensions (5×10⁴ conidia/ml) were infiltrated into rice leaves. Photographs were taken seven days after inoculation. (B) Sheath inoculation assay. Forty microliters of a conidial suspension (3×10⁴/ml) of the strains were inoculated onto rice sheath cells for 48 h. The arrowhead indicates appressorium (App).
The Melanization of MoSDR1

MoSDR1 is a member of the SDR family of proteins, which are known to be involved in the biosynthesis of melanin, a pigment that is crucial for fungal pathogenicity. In particular, the MoSDR1 gene encodes a new member of the SDR family. The targeted deletion of MoSDR1 resulted in a significant reduction in conidial reproduction whereas its mycelial growth was not affected, indicating the requirement of MoSDR1 in conidiation. This result correlates with the observation that the expression of MoSDR1 was dramatically induced during conidiation. This implies that the event of conidiation is mediated via the transcriptional regulation of MoSDR1 gene, rather than direct regulation of MoSDR1 activity. Consistent with this idea was the observation that phenotypic defects in the MoSDR1ΔT-DNA were quite similar to those in the ΔMosdr1. Development of conidia, as propagules, is a key step to ensure fungal survival and infectability. In general, fungi terminate conidial development through desiccation after all the requirements are prepared in conidia. Although conidial morphology of MoSDR1 deletion mutant was indistinguishable from the wild-type conidia, MoSDR1 conidia failed to germinate. The germination defect of ΔMosdr1 conidia may not be caused by their inability to sense environmental factors such as nutrients and moisture, but by a metabolic abnormality in the process of conidial development. In a histochemical study, we showed this to be the case that the ΔMosdr1 conidia lost viability. In addition, appressorium development was completely abolished on germinated conidia of the ΔMosdr1.

These results suggest that MoSDR1-mediated metabolic regulation is indispensable not only for pre-infection stages, but for pathogenic development.

MoSDR1 showed a high similarity to filamentous fungal proteins. It is tempting to speculate that the MoSDR1 and its homologous proteins have a common enzymatic mechanism among filamentous fungi. SDRs utilize a wide range of substrates, including steroids, alcohols, sugars, and aromatic compounds. The substrate of MoSDR1, which may be rich in conidia based on the phenotypes of ΔMosdr1, has not been identified yet. Further studies are needed to elucidate the metabolic role of MoSDR1 in conidial development.
Acknowledgments

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