The bZIP Transcription Factor MoAP1 Mediates the Oxidative Stress Response and Is Critical for Pathogenicity of the Rice Blast Fungus Magnaporthe oryzae

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Abstract
Saccharomyces cerevisiae Yap1 protein is an AP1-like transcription factor involved in the regulation of the oxidative stress response. An ortholog of Yap1, MoAP1, was recently identified from the rice blast fungus Magnaporthe oryzae genome. We found that MoAP1 is highly expressed in conidia and during invasive hyphal growth. The Moap1 mutant was sensitive to H2O2, similar to S. cerevisiae yap1 mutants, and MoAP1 complemented Yap1 function in resistance to H2O2, albeit partially. The Moap1 mutant also exhibited various defects in aerial hyphal growth, mycelial branching, conidia formation, the production of extracellular peroxidases and laccases, and melanin pigmentation. Consequently, the Moap1 mutant was unable to infect the host plant. The MoAP1-eGFP fusion protein is localized inside the nucleus upon exposure to H2O2, suggesting that MoAP1 also functions as a redox sensor. Moreover, through RNA sequence analysis, many MoAP1-regulated genes were identified, including several novel ones that were also involved in pathogenicity. Disruption of respective MoGG_01662 (MoAAT) and MoGG_02531 (encoding hypothetical proteins) genes did not result in any detectable changes in conidial germination and appressorium formation but reduced pathogenicity, whereas the mutant strains of MoGG_01230 (MoSSADH) and MoGG_15157 (MoACT) showed marked reductions in aerial hyphal growth, mycelial branching, and loss of conidiation as well as pathogenicity, similar to the Moap1 mutant. Taken together, our studies identify MoAP1 as a positive transcription factor that regulates transcription of MoGG_01662, MoGG_02531, MoGG_01230, and MoGG_15157 that are important in the growth, development, and pathogenicity of M. oryzae.

Introduction
Organisms such as plants have evolved to develop many efficient defense systems against pathogenic microbes. Among them, reactive oxygen species (ROS), primarily superoxide and H2O2, produced by plasma membrane-localized NADPH oxidases [1], are regarded as one of the fastest defense reactions against pathogen attack [2]. During the plant defense response, ROS was used by apoplastic peroxidases on the cell wall to synthesize lignin and other phenolic polymers that prevent pathogen invasion into the host [3]. There are reports that ROS produced at the site of an attempted invasion may also function as a secondary messenger in the induction of various plant defense-related genes and is essential for the pathogen-associated molecular pattern (PAMP)-triggered immunity (PTI) response in plants [4,5,6,7]. Additionally, due to the toxicity, ROS accumulated at the site of pathogen invasion can directly kill the pathogen [8,9]. Conversely, plant pathogens have also developed many strategies, including enzymatic and non-enzymatic ones, to detoxify ROS and successfully invade their hosts [10,11,12,13]. The relative sensitivity of the fungal pathogen to ROS may also depend on the effectiveness of its own ROS detoxification or tolerance machinery. In fungal pathogens, transcription factor-mediated host-derived ROS detoxification through regulation of gene expression is important in plant-microbe interactions [14,15,16]. Detoxifying enzymes, either preformed or inducible, including superoxide dismutase, catalases, and peroxidases, are thought to contribute to the tolerance of ROS in pathogenic fungi [15,17,18,19].

Saccharomyces cerevisiae transcription factor Yap1 functions as one of the most important determinants of the yeast’s response to the oxidative stress and Yap1 is responsible for transcriptional activation of various genes involved in ROS detoxification [20,21,22,23,24]. Loss of Yap1 function resulted in increased sensitivity to external stresses. A comparison of AP1 transcription factors from eukaryotic organisms revealed a conserved N-terminal basic leucine zipper

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Author Summary

*Magnaporthe oryzae* is a causal agent of rice blast disease and an important model for understanding of fungal development and pathogenicity. To examine the molecular mechanisms involved in conidium formation and appressorium development of *M. oryzae*, we identified the transcriptional factor MoAP1 as a regulator of the oxidative stress response. Our results indicated that MoAP1 is a stage-specific regulator for conidium formation, morphology, aerial hyphal growth, and also growth in planta. Additionally, we identified four novel genes whose functions were linked to MoAP1 and pathogenicity. Disruption of *MGG_01662* (encoding aminobutyrate aminotransferase, MoAat) and *MGG_02531* (hypothetical protein) caused minor phenotypic changes but attenuated virulence, and disruption of *MGG_01230* (encoding succinic semialdehyde dehydrogenase, MoSsadh) and *MGG_15157* (encoding acetyltransferase, MoAct) resulted in drastic reductions in the growth of aerial hyphae and hyphal branching as well as loss of conidiation and pathogenicity. Our studies extend the current understanding of AP1 functions in fungi and reveal that the MoAP1-mediated regulatory network is associated with the pathogenicity of *M. oryzae*.

(bZIP) DNA-binding domain, consisting of a leucine zipper that mediates dimerization [25] and an adjacent basic region that specifically interacts with DNA sequences [24]. At the C-terminus, the cysteine-rich domains (c-CRD) are highly conserved [26] and play a key role in Yap1-mediated resistance to the oxidative stress and, together with an n-CRD, for the appropriate subcellular localization of the Yap1 protein [27,28]. Additionally, mutation of cysteine residues in Yap1 resulted in increased sensitivity to a variety of oxidizing compounds and drugs [29]. To date, Yap1 homologs were identified in several fungal pathogens [14,15,30,31,32], which of oxidizing compounds and drugs [29]. To date, Yap1 homologs were identified in several fungal pathogens [14,15,30,31,32], which were identified in several fungal pathogens [14,15,30,31,32], which play a key role in Yap1-mediated resistance to the oxidative stress and, together with an n-CRD, for the appropriate subcellular localization of the Yap1 protein [27,28]. Additionally, mutation of cysteine residues in Yap1 resulted in increased sensitivity to a variety of oxidizing compounds and drugs [29]. To date, Yap1 homologs were identified in several fungal pathogens [14,15,30,31,32], which share the function in stress tolerance but differ in pathogenicity. Yap1-mediated ROS detoxification was an essential virulence determinant in *Usitago maydis* [15], *Alternaria alternata* [14], and *Candida albicans* [31], but it had no role in virulence of *Cochliobolus heterostrophus* and *Aspergillus fumigatus* [30,32].

*Magnaporthe oryzae* is a pathogen of both economical and scientific importance [33,34]. Like most other fungal pathogens, conidia of *M. oryzae* play a central role in the disease cycle. When attached on the host surface, conidia begin to germinate and develop appressorium from the end of the germ tubes [35]. The mature appressorium generates enormous turgor pressure (8 MPa) to help penetrate the plant cuticle and enter the plant cells [36]. After penetration, infection hypha spread through the rice leaf cells and typical necrotic lesions develop on the surface of rice leaves. Eventually, aerial conidioles differentiate from hyphae on the lesion and newly formed conidia are released to serve as secondary inocula for new infections. In the past two decades, efforts have been made to study the conidiation process, formation of appressorium, and host plant responses to infection. Studies have suggested that *M. oryzae* infectious hyphae is biotrophic and it secretes effectors, such as biotrophy-associated secreted (BAS) proteins that can alter host cellular defense processes [37,38]. The availability of genome sequences for both *M. oryzae* and rice host provided a new platform to identify pathogenicity-related genes and to understand molecular pathogenesis at the genome level [39,40].

In this study, we identified MoAP1 as a homolog of the bZIP transcription factor AP1. We also identified four other proteins as the downstream targets of MoAP1, which were also involved in conidiation and pathogenicity.

Results

Identification of MoAP1 from *M. oryzae*

Using *S. cerevisiae* Yap1 sequence as trace, we identified the locus MGG_12814 ([XP_001408783](http://www.broad.mit.edu/annotation/genome/magnaporthe_grisea/Home.html)) [40]. MGG_12814 was predicted to encode a 576-amino acid (aa) protein that shares substantial similarity (19–50%) to a number of fungal AP1 proteins. We thus named the protein MoAP1. Analysis of MoAP1 showed several conserved domains including a bZIP DNA-binding domain, a nuclear localization domain near the N-terminus (aa 150–214), and a cysteine-rich domain at the C-terminus (aa 492–551). Alignments of MoAP1 with other Yap1-like proteins revealed high similarity in the bZIP and c-CRD domains (see Figure S1A and S1B). The alignments of the bZIP domains also revealed the most conserved residues Q161, N162, A165, Q166, A168, F169, and R170 in the basic region (see Figure S1A). The c-CRD domain is also rich in cysteine (C506, C530, and C539) and serine (S540) residues, and a putative nuclear export sequence (NES) within the c-CRD (aa 526–535) is a possible binding site for the Crm1p-like exporter (see Figure S1B) [41]. The phylogenetic relationship of MoAP1 to other AP1 proteins revealed that AP1-like proteins in filamentous fungi apparently separated from those of unicellular yeasts, with *M. oryzae* MoAP1 most similar to *Gibberella zeae* GzAP1 ([XP_388976](http://www.ncbi.nlm.nih.gov/nuccore/78251633)) and *Fusarium oxysporum* FoAP1 ([XP_388976](http://www.ncbi.nlm.nih.gov/nuccore/78251633)) (see Figure S1C).

MoAP1 complemented the growth defect of a yeast yap1 mutant under oxidative conditions

To determine the function of MoAP1, we tested whether MoAP1 complements a *yap1* mutant of *S. cerevisiae*. An expression vector pYES2 containing the full-length MoAP1 gene was transformed into the *yap1* mutant. As a control, the *yap1* mutant was also transformed with an empty pYES2 vector. When plated on glucose- (suppression) or galactose-containing (induction) medium in the absence of oxidizing agents, no growth defect was observed (see Figure S2). However, when 0.3 mM hydrogen peroxide (H$_2$O$_2$) was added, only the wild-type strain grew on glucose-containing medium. On medium containing both 0.3 mM H$_2$O$_2$ and galactose, the *yap1* pYES:MoAP1 formed colonies similar to the wild type, while the *yap1* mutant carrying the empty pYES2 vector was significantly inhibited (see Figure S2). This indicated that the ability of the *S. cerevisiae yap1* mutant to cope with H$_2$O$_2$ stress could be complemented by the introduction of *MoAP1*. However, judging by growth, the complementation appeared to be partial (see Figure S2).

MoAP1 is highly expressed in conidia and in invasive hyphae

To explore the function of MoAP1, we examined its expression in *M. oryzae* by quantitative real-time polymerase chain reaction (qRT-PCR, see Figure S3A). The abundance of *MoAP1* transcripts during vegetative growth in liquid CM was relatively lower than in conidia, when compared with the expression of the control actin gene (*MGG_03982.5* (avg. dCt = 6.0). In the invasive growth in planta, *MoAP1* mRNA accumulation decreased almost two-fold at 8 hrs and 11-fold at 24 hrs post-inoculation (hpi) on rice leaves, compared with that of conidia at 0 hpi. However, the transcript abundance began to increase after 24 hpi, reaching almost the same level as that of conidia at 72 hpi (see Figure S3A). Thus, we concluded that the *MoAP1* was highly activated during conidiation and infection.
MoAP1 is necessary for aerial hyphal growth but not mycelial radial growth

To understand the role of MoAP1 in growth, we generated a Moap1 mutant strain using the Moap1 disruption allele linked to a hygromycin resistance marker gene (see Figure S3B). Linear DNA fragments amplified from the Moap1 disruption allele were used to transform the protoplasts of the wild type strain Guy11. Putative transformants were screened on hygromycin media and verified by PCR amplification (see Figure S3C). The mutants were also confirmed by RT-PCR (see Figure S3D) and Southern blotting analysis (see Figure S3E and S3F). Two deletion mutants, Moap1-9 and Moap1-20, were selected for further analysis. To complement the mutant strain, the genomic DNA sequence of MoAP1 containing a 2-kb promoter region was reintroduced into the Moap1 mutant and verified using PCR amplification (see Figure S3C and S3D).

On CM medium for 5 days at 28°C, the Moap1 mutants showed no apparent defect in radial growth, but presented as a flat colony due to the reduced aerial hyphal growth (Figure 1A and 1B) with altered pigmentation (Figure 1D). We further incubated Guy11, Moap1 mutants, and the complemented strain in liquid CM for 48 hrs and found that the mutant strains formed compact mycelia masses, in contrast to the sparse ones formed by the wild-type and complemented strains (Figure 1C). We also compared hyphal branching patterns of the Moap1 mutants with the control strains using Calcofluor white (CFW) staining and found that the hyphal branching was severely reduced in the Moap1 mutant (see Figure S4A). Together, these results indicated that MoAP1 is essential for proper growth and hyphal branching.

MoAP1 disruption resulted in abnormal conidium morphology and reduction in conidia formation

Conidia, which are borne on specialized stalks called conidiophores, play an important role in the disease cycle of rice blast [42]. Given that the Moap1 mutant showed reduced hyphal growth, we investigated the role of MoAP1 in conidia formation. Conidiation of the wild-type strain (Guy11), Moap1 mutants (Moap1-9 and Moap1-20), and the complement strain (Moap1/AP1) was determined in 10 day old RDC cultures [43]. The most striking finding was that conidiation was dramatically reduced by approximately 30 to 40-fold in Moap1 deletion mutants (Moap1-9, 30-fold; Moap1-20, 40-fold), compared with the wild-type and complement strains.
Of the few conidia that formed in Moap1-9 and Moap1-20, most exhibited abnormal, elongated, and spindly morphology (Figure 2C and 2D). Combined with the expression profiles of MoAP1 by qRT-PCR that showed a much higher level of MoAP1 expression in conidia (see Figure S3A), we concluded that MoAP1 plays an important role in conidial formation.

**MoAP1 disruption leads to hypersensitivity to the oxidative stress**

*S. cerevisiae* Yap1 was involved in the oxidative stress response [21,22]. To investigate whether MoAP1 exhibits the same function, the wide-type, Moap1 mutants, and the complemented (Moap1/AP1) strains were exposed to H$_2$O$_2$. The mycelia growth of the Moap1 mutants was apparently affected (Figure 3A and 3B). Exposure to 2.5 and 5 mM H$_2$O$_2$, respectively, led to an average 19% (2.5 mM) or 22% (5 mM) greater growth inhibition rate than the wild-type strain (Figure 3B). The involvement of the MoAP1 gene in H$_2$O$_2$ resistance was confirmed by genetic complementation in which the complemented strain (Moap1/AP1) was as resistant to H$_2$O$_2$ as the wild-type strain (Figure 3A and 3B).

**MoAP1 disruption led to excess oxidative bursts during conidiation and germination**

In fission yeast *Schizosaccharomyces pombe*, the transcription factor Pap1, together with another transcription factor SpAtf1, regulates the genes involved in ROS homeostasis and the response to the extrinsic oxidative stress [44,45]. Intracellular ROS is known to have multiple functions in fungal pathogenicity [14,15,30,32,46]. In *M. oryzae*, intracellular ROS is the key to its virulence in rice seedlings, and the NADPH oxidase mutants lost virulence on the susceptible rice cultivar CO-39 because of their obstructed ROS production [46]. Thus, visualization of ROS accumulation was performed to investigate ROS metabolism during conidiation and germination in Moap1 mutants. We first investigated the production of ROS using dihydrorhodamine 123, which exhibits green fluorescence during reduction by superoxide radicals. Using this technique, it appeared that the Moap1 conidium accumulated higher amounts of superoxide than the wild-type (Figure 4A). Such increased accumulation of superoxide was also detected in the Moap1 mutants during conidial germination. Green fluorescence was typically more intense in the germ tubes and mature appressoria of the Moap1 mutants than the wild-type strain (Figure 4A). To further confirm enhanced accumulation of ROS in the Moap1 mutants, another kind of reactive oxygen species detection probe, nitroblue tetrazolium (NBT), which forms a dark-blue water-insoluble formazan precipitate on reduction by superoxide radicals, was used. Using this procedure, we found that the Moap1 mutants accumulated higher amounts of superoxide, with more intense formazan precipitates in the germ tubes and mature appressoria (Figure 4B). In contrast, the appressoria and infection hyphae of the wild type strain had less formazan precipitates than the Moap1 mutants (Figure 4B). Thus, both staining results indicated that MoAP1 disruption leads to excess oxidative bursts in conidiation and germination.

![Figure 2. MoAP1 disruption leads to abnormal conidial morphology and reduced conidiation. (A) Development of conidia on conidiophores is affected by Moap1 deletion. Strains grown on RDC medium for 7 days were examined by light microscopy. Bars equal 100 μm. (B) Statistical analysis of conidia production. The conidia produced by the wild type strain (Guy11), the mutants and complemented strains grown on RDC medium for 10 days were collected, counted, and analyzed by Duncan analysis (p<0.01). Asterisks indicate significant differences among Guy11, the Moap1 mutants and complemented strains. Error bar represents standard deviation. (C) Conidium morphology. Conidia were harvested from RDC medium, diluted to 1.0×10^5 spore/ml, and observed by light microscopy. Bars equal 10 μm. (D) Conidia size comparison. The conidia sizes were determined as width by length from 150 conidia of each strain. Asterisks indicate that the difference is statistically significant. Error bars represent standard deviations. doi:10.1371/journal.ppat.1001302.g002](https://www.plospathogens.org_article_image/10.1371/journal.ppat.1001302.g002)
Subcellular localization of MoAP1 is modulated by H$_2$O$_2$

AP1 proteins are translocated from the cytoplasm to the nucleus following the oxidative stress [14,15,27,30]. To analyze if this is also true for MoAP1, we generated a C-terminal MoAP1::eGFP fusion gene under the drive of the TrpC promoter and introduced the fusion gene into the wild-type strain. Conidia of the MoAP1::eGFP containing transformants were harvested and observed under a fluorescence microscope. In the absence of the oxidative stress, the MoAP1::eGFP fusion protein was distributed throughout the cell and was apparently excluded from nuclei (see Figure S5). In contrast, fluorescence was concentrated in a single spot after exposure to 2 mM H$_2$O$_2$ for 2 hrs, as shown by the colocalization of the eGFP and DAPI fluorescence signals (see Figure S5), indicating the nuclear localization of MoAP1::eGFP in response to the oxidative stress.

MoAP1 is required for invasive hyphae growth and pathogenicity

To determine whether MoAP1 was involved in pathogenicity, conidial suspensions (1 x 10$^5$ conidia/ml) of both the Moap1 mutant...
and wild-type strains were sprayed onto 4-week-old rice seedlings *(Oryza sativa cv CO-39)*. At 5 days after inoculation, symptoms had fully emerged on rice leaves inoculated with the wide-type strain, but no lesion on the *Moap1* mutant inoculated rice leaves (see Figure 8). When observation was made at 7 days post infection, there were still no typical lesions developed on *Moap1* mutant infected leaves. Only small necrotic-like dark brown spots were observed occasionally (Figure 5A). The loss of pathogenicity was complemented by reintroducing the *MoAP1* gene into the *Moap1* mutant (Figure 5A). Similarly, pathogenicity test with conidia suspension or mycelial plug on barley leaves displayed very similar result (see Figure S6A and S6A).

To investigate possible reasons for the lost pathogenicity, an onion epidermis penetration assay was performed. At 48 hours post-inoculation, both the wild-type and the *Moap1* mutant strains could penetrate the onion epidermis cell, but the invasive hyphae of the wild type strain freely expanded into the onion epidermis cells, in contrast to the restricted growth of the *Moap1* strain (Figure 5B). The deficiency of hyphae expansion could be complemented by reintroduction of the *MoAP1* gene (Figure 5B). Further assay using the rice leaf sheath generated the similar results (Figure 5C). Moreover, DAB staining showed the accumulation of ROS at the infection site of the *Moap1* mutant, but not the wild type and complemented strains (Figure 5D). These results indicated that *MoAP1* is required for invasive hyphae growth and the defect in invasive hyphae might be responsible, at least in part, for the loss of pathogenicity.

**MoAP1** disruption attenuates the activity of extracellular peroxidases and laccases

In *M. oryzae*, defects in cell wall composition can influence appressorium formation and impair successful infection of rice host [47,48,49,50]. Congo Red (CR), which binds to cell wall component β-1,4-glucan [51], is commonly used to detect cell wall integrity. To determine whether MoAP1 has a role in cell wall integrity, CR was added to CM medium (200 μg/ml). No difference was found in the growth of mycelium between the *Moap1* mutant (15% inhibition) and the wild-type strain (16% inhibition). However, the degradation halo of CR by the *Moap1* mutant was not as apparent as the wild type (Figure 6A). This indicated a deficiency of the CR-degrading activity in the *Moap1* mutant. An enzyme activity assay using culture filtrates further indicated that the *Moap1* mutant nearly lost all of the peroxidase activity (Figure 6D). Moreover, we determined the activity of additional extracellular enzyme laccases in either solid or liquid CM medium, and found that, in each case, the decreased laccase activity was observed in the *Moap1* mutant, with less oxidized dark purple stain around its colony and lower levels of laccase activity in the culture filtrate, compared with the wild-type strain (Figure 6B and 6C). These data suggest that *MoAP1* disruption resulted in a decreased peroxidase and laccase activities.

**Addition of copper sulfate to the Moap1 mutant restores laccase activity and complements pigmentation**

We observed altered pigmentation of the *Moap1* mutants on CM medium, with a yellowish-brown color to the mutant strains compared with a dark pigmentation of the wild-type strain (Figure 1D). It is well known that copper sulfate can stimulate the biosynthesis of melanin by inducing the laccase activity [52,53]. When copper sulfate was added to CM medium at 1 mM, which did not affect mycelial growth of either the *Moap1* mutants or the wild-type strain, the laccase activity was restored to the *Moap1* mutants, as indicated by dark pigmentation (see Figure S7). Combined with the accumulation of reactive oxygen species in conidia and mycelia, we hypothesized that the *Moap1* mutant was likely in a hyperoxidative state, which may, in turn, lead to reduced oxidative cross-linking of melanin, and which could be compensated for by an increase in the laccase activity.

**Differential expression of pathogenicity-associated genes revealed genes linked to MoAP1**

To understand reasons for phenotypic changes and the loss of pathogenicity in the *Moap1* mutants, we generated serial analysis of gene expression (SAGE) libraries for the wild-type strain (Guy11, 4,924,107 tags) and the *Moap1* mutant (4,690,301 tags) using mycelia grown in liquid CM medium. To confirm gene expression patterns derived from the SAGE libraries, 10 down-regulated genes in the *Moap1* mutant were randomly selected and validated by qRT-PCR. The results showed that each gene expression pattern was consistent with that in the SAGE data (Figure 7A and Table S1), despite the discrepancy of the fold-change being higher in qRT-PCR than in SAGE data.

To identify genes that were subjected to regulation by MoAP1, we compared the gene expression profiles between the wild-type
strain and the *Moap1* mutant. In total, 497 genes were up-regulated and 682 genes were down-regulated (Figure 7B). These genes were functionally grouped into GO categories based on manual curation, as described in Materials and Methods. We found that 69 genes related to redox-homeostasis were altered in expression and 5/6 of those genes, either with the signal peptide or not, were down-regulated in the *Moap1* mutant (Figure 7C). We also noted a significant decrease in the expression of genes involved in transcriptional regulation, protein degradation, lipid metabolism, secondary metabolism, cellular transportation, and cell development (Figure 7C).

Since the *Moap1* mutants were non-pathogenic due to the confinement of its invasive hyphae within the originally penetrated onion skin cells and rice leaf sheath cells, we examined the putative target genes of MoAP1 based on their previously defined roles and identified seven genes that were previously characterized to regulate virulence (Table 1). Because of the severely decreased conidiation of the *Moap1* mutants, we screened the SAGE database to identify genes that were involved in conidiation. A previously defined gene *MoCOS1* (*MGG_03977*) involved in conidiophore stalk formation [54] and a novel vitamin B6 synthesis amidotransferase-encoding gene (*MGG_05981*) [55] were found to be also significantly down-regulated in the *Moap1* mutant (see Table 1 and Figure S8A). Additionally, three laccase-encoding genes (*MGG_13464*, *MGG_11608*, and *MGG_09139*) were also found to be severely down-regulated (see Table 1 and Figure S8B).

MoAP1 regulates the expression of four genes involved in pathogenicity

Among MoAP1 regulated genes, we selected 10 whose expression was severely down regulated, and of which 7 contained the putative AP1 binding sites (Figure 8, Table 2 and Table S1), and characterized their functions by generating individual gene disruption mutants. Among the deletion mutants, we compared the phenotypes in mycelial growth, conidiation, appressorium formation, and pathogenicity on rice, and found that disruption of the transcription factor MoOefC (*MGG_10422*), the hypothetical protein-encoding gene (*MGG_13654*), the laccase protein-encoding gene (*MGG_13464*), the

![Figure 5. Pathogenicity of the *Moap1* mutant strain.](image-url)
Mossadh and MoAct are required for appressorium-like structure-mediated penetration

Generally, *M. oryzae* infects rice aerial organs, such as leaves and stems, through appressoria, which develops from conidia. However, hyphae can also invade rice roots [56] and wounded leaf tissue [57]. Previous studies suggested that the hyphal tips can also form an appressorium-like structure to break the rice leaf cuticle and cause disease [54,58]. As both the *Mossadh* and *MoAct* mutants lost pathogenicity on rice leaves, we hypothesized that the lost pathogenicity might be due to the lack of the ability to breach the rice leaf cuticle. Given that successful lesion development requires the development of appressoria, we examined appressorium formation on the onion skin using microscopy and found that both the wild-type and the mutants developed appressorium-like structures at the hyphal tips (Figure 9A and 9C). However, the appressorium-like structures formed by the *Mossadh* (Figure 9E) and *Moact* mutants (Figure 9F) were significantly fewer than that formed by the wild-type strain (34%). Furthermore, those appressorium-like structures could not penetrate the onion skin cells, compared with the easy penetration of the cells by the wild-type strain (Figure 9B and 9D). The penetration assay using the rice leaf sheath generated similar results (Figure 9G). These results indicate that both *MoSsadh* and *MoAct* are required for appressorium-like structure formation, as well as penetration.

*Mossadh* and *MoAct* are equally required for invasive hyphal growth

The results above indicated that *Mossadh* and *MoAct* mutants lost the ability to penetrate the plant cell. If these results are the main reason for the lost pathogenicity, given the conditions, abraded rice leaves should restore pathogenicity to the mutants. To examine this, we inoculated wounded rice leaves with agar plugs containing mycelial tips to evaluate pathogenicity. Both the *Mossadh* and *MoAct* mutants were unable to cause symptoms, while the wild-type strain produced visible diffuse lesions on rice leaves 5 days after inoculation (Figure 10A and 10B). The similar result was found when the observation was made at 7 days post infection (dpi) (see Figure S9B).

To further investigate the loss of pathogenicity, we injected the mycelial fragments into rice leaves using a syringe. The wild-type strain caused typical necrotic symptoms along the injection site and formed developmental lesions at 5 dpi (Figure 10C). However, the *Mossadh* mutant caused no necrotic symptoms, while the *MoAct* mutant caused severely restricted necrosis (Figure 10C). These results suggested that the loss of pathogenicity in *Mossadh* and *MoAct* mutants is not only due to failure in penetration, but also to the lost ability of forming invasive hyphae. Similar results were observed at 7 dpi (see Figure S9A).

*Mossadh* and *MoAct* mutants are both hypersensitive to H$_2$O$_2$

The stress-tolerance mechanisms of plant pathogens play an important role in virulence [14,15,59,60]. To assess whether the putative MoAP1 targets MoSsadh and MoAct play an active role in the tolerance to exogenous H$_2$O$_2$, the mutant strains were inoculated on H$_2$O$_2$-containing CM medium. The assay results showed that both type of mutants were more sensitive to H$_2$O$_2$ than the wild-type strain (Figure 11A and 11C). The mycelial growth of the *Mossadh* mutant was severely inhibited on CM medium containing 5 mM H$_2$O$_2$, with 20% (*Mossadh-4*) and 17% (*Mossadh-6*) greater inhibition rates than the wild-type strain (Figure 11B). Meanwhile, the *MoAct* mutants displayed higher sensitivity to H$_2$O$_2$ than either the *Moap1* mutants or the *Mossadh* mutants, with an average 10% greater mycelial growth inhibition rate than the wild-type strain at 2 mM H$_2$O$_2$ and 61% at 5 mM H$_2$O$_2$. The sensitivity of the mutants to H$_2$O$_2$ was complemented by reintroduction of the respective wild-type genes (Figure 11D).

*MoSsadh* and *MoAct* mutants displayed similar morphological phenotypes to *Moap1* mutants

To further investigate the roles of *MoSsadh* and *MoAct* on hyphal growth, we compared their growth on CM and RDC
medium. The *Mossadh* and *Moact* mutants exhibited reduced growth (see Figure S10), and the aerial hyphae was also sparser and thinner, compared with Guy11 (Figure 12A). When observed by light microscopy, no conidia were found (Figure 12C). The colonies of the two mutants were also smooth appearing and displayed 5-10 radial folds in each colony, similar to the *Moap1* mutants (Figure 8 and 12A). To further understand possible reasons for this, we grew the *Moap1* mutants in liquid CM medium and found that it could form compact mycelia mat, in contrast to the sparse one formed by the wild-type and the complemented strain (Figure 12B). Meanwhile, the hyphal branching patterns of Guy11, the *Mossadh* mutants, the *Moact* mutants, and the complemented strains were examined using CFW staining, which revealed that hyphal branching was severely reduced in the *Mossadh* and *Moact* mutants (see Figure S4B and S4C). Moreover, the colonies of the *Mossadh* and *Moact* mutants were less pigmented than the wild-type and complemented strains (see Figure S10A and S10B). Combined with the results of the *Moap1* mutants, these results further indicated that MoAP1 controls the morphological phenotypes of *M. oryzae* by regulating genes including *MoSSADH* and *MoACT*.

Disruption of *MoSSADH* and *MoACT* attenuated peroxidase and laccase activities

To functionally analyze possible reasons for attenuated virulence of both the *Mossadh* and *Moact* mutants in rice, we examined the cell wall integrity, which is regarded as the most important factor affecting appressorium formation in *M. oryzae* [47,48,49,50]. By the addition of CR to CM medium at 200 μg/ml, the mycelial growth of both the *Mossadh* mutants (average 17% inhibition rate) and *Moact* mutants (average 16% inhibition rate) was similar to that of the wild-type strain (average 16% inhibition rate). However, no degradation halo of CR on either the *Mossadh* or *Moact* mutant plates was present in comparison to the wild type.

Figure 7. Differential gene expression analysis on transcriptomes of the *Moap1* mutant and Guy11 strains. (A) Real time RT-PCR validated the RNA-SEQ results. Real time RT-PCR was carried out to confirm the RNA-SEQ results through random selection of genes that were down-regulated in the *Moap1* mutant. Laccase genes (I), extracellular peroxidase gene (II), and genes involved in the redox homeostasis (III) were down-regulated in the *Moap1* mutant. IV and V indicated MGG_01230 (*Mossadh*) and MGG_10315 (*MoMpg1*) that were down-regulated. (B) Numbers of altered genes expression in *Moap1* mutants. Gene expression profiles were analyzed and 682 genes were down-regulated while 497 genes were upregulated in *Moap1* mutants in comparison to the wild type strain (Guy11). Genes whose expression were up or down as indicated by expression profiling were chosen based on the log2 ratio (*Moap1*/Guy11) values that were either 1.5- fold more or - less. DR and UR denote down- and up-regulation. (C) Functional grouping of genes up- or down-regulated in *Moap1* mutants. The up-regulated (in purple) and the down-regulated (in cyan) genes were divided into 23 groups according to their putative functions as described in Materials and Methods.

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and complemented strains, which displayed a visible degradation halo of CR (see Figure S11A). To test for secreted peroxidases, culture filtrates of the *Mossadh* mutant, *Moact* mutant, wild-type Guy1, and complemented strains were collected and assayed for secretion of extracellular peroxidases. The results showed that both the *Mossadh* and *Moact* mutants were deficient in peroxidase activities (see Figure S11C and S11E). For the altered pigmentation in the *Mossadh* and *Moact* mutants, we again compared the laccase activity, hypothesizing that the polyphenol oxidase function may be impaired in both these mutants. The oxidation of the laccase substrate 2,2'-azino-di-3-ethylbenzthiazoline-6-sulfonate (ABTS) showed that the laccase activity was reduced in both the *Mossadh* and *Moact* mutants. The oxidized dark-purple stain around the colonies of the *Mossadh* and *Moact* mutants was less than that of Guy1 (see Figure S11B). Similarly, the laccase activity in the filtrates of the *Mossadh* and *Moact* mutants was also reduced (see Figure S11D and S11F). These data suggested that the deletion of either *Mossadh* or *Moact* equally results in decreased activity of peroxidases and laccases in *M. oryzae*, similar to the *Moap1* mutants.

### Discussion

In this study, we have characterized *M. oryzae* MoAP1 as a homolog of fungal AP1 protein, such as *S. cerevisiae* Yap1 and Schiz. pombe Pap1. Similar to other members of the AP1 family [12,29], MoAP1 contains a bZIP domain at the N-terminal, a nuclear localization motif, and c-CRD and/or n-CRD domains that are vital for cellular localization of Yap1 and resistance to oxidative damages [20]. In both filamentous and unicellular fungi, the nuclear localization of AP1 under the oxidative stress is a crucial step for the function of transcription regulation [27], and intramolecular disulfide bridge formation by two cysteine residues from the c-CRD and n-CRD regions is thought to be necessary for its nuclear translocation [41,61]. It has also been established that the AP1 proteins are translocated in the nucleus in response to the oxidative stress [20,62]. We have found that MoAP1 could partially complement the H$_2$O$_2$-sensitive phenotype of a yeast *yap1* mutant, *Moap1* deletion mutants were sensitive to H$_2$O$_2$, and MoAP1 localized in the nucleus when exposed to H$_2$O$_2$, all suggesting that MoAP1 is vital for resistance to the oxidative damage in *M. oryzae*.

To survive, fungi have evolved sophisticated mechanisms for adapting to stresses from intracellular or extracellular sources. During developmental processes, a fungus encounters various stresses, including the toxic by-products of its metabolism and the oxidative stress generated through aerobic respiration [63,64]. It has also been documented that the cellular environment within host plants also serves as one of the major sources of stress to invading fungal pathogens [65,66]. To evade the stress, fungi need specialized adaptation mechanisms. It is known that the AP1 transcription factors are one of the major regulators to activate genes in responding to the exogenous oxidative stress. In *Schiz. pombe*, two parallel H$_2$O$_2$-responsive pathways exist. The transcription factor Pap1, together with the MAP kinase Sty1 pathway, regulate gene expression in response to the oxidative stress [67]. In *S. cerevisiae* and *Candida albicans*, deletion of *YAP1* leads to reduced tolerance to the oxidative stress and/or attenuated virulence [21,22,62]. Studies of *U. maydis* also showed that Yap1 plays a critical role in the H$_2$O$_2$ detoxification system and in the infection of maize plants [15]. A similar role was also found for *A. alternata* AaAP1 [14]. In *C. heterostrophus* and *A. fumigatus*, the AP1 proteins ChAP1 and AfAP1 mediate H$_2$O$_2$ response but are not required for pathogenicity [14]. This indicates that the AP1 proteins are conserved regulators of the oxidative stress, but their roles in virulence are diverged. The *M. oryzae* AP1 protein appears to be close to *U. maydis* AP1 and AaAP1 in that it is required for both H$_2$O$_2$ resistance and pathogenicity. As a step to further advance our knowledge of AP1 proteins, we utilized the transcription analysis by the SAGE approach to identify genes involved in redox homeostasis whose expression is also down-regulated due to *Moap1* mutation. Based on our results, we hypothesize that MoAP1 is a key factor in regulating genes involved in the detoxification of H$_2$O$_2$, and that the severely restricted infectious hyphal growth of the *Moap1* mutants may be due to the reduced tolerance of the oxidative stress, which is responsible for the loss of pathogenicity.

### Table 1. SAGE data for genes encoding known pathogenicity factors or pathogenicity-associated functions.

| Gene_ID   | Description                                | ELSD$^a$ (Moap1/WT) | p-Value  | Reference | Putative MoAP1 binding sites $^b$ |
|-----------|--------------------------------------------|---------------------|----------|-----------|----------------------------------|
| MGG_10315 | Hydrophobin (Mgp1)                         | −9.72               | 0        | [89]      | Yes                              |
| MGG_07536 | Metalloprotease/Zn amino-peptidase          | −2                  | 5.36E-05 | [55]      | No                               |
| MGG_09312 | Transcription factor Zn2Cys6               | −1.781              | 4.46E-45 | [55]      | No                               |
| MGG_00111 | P-type ATPase (Pde1)                       | −1.58               | 0.00043  | [102]     | Yes                              |
| MGG_05344 | Snodpro1 homolog (Msp1)                    | −1.39               | 0        | [103]     | No                               |
| MGG_06424 | PEX3-peroxisome                           | −1                  | 0.01     | [55]      | No                               |
| MGG_09898 | Adenylate cyclase (Mac)                    | −1                  | 0.027    | [104]     | No                               |
| MGG_03977 | Putative Zinc-Finger Protein               | −4.52               | 3.29E-48 | [54]      | No                               |
| MGG_05981 | vitamin B6 synthesis amidotransferase      | −2.70               | 2.74E-243| [55]      | Yes                              |
| MGG_13464 | laccase                                    | −7.61               | 0        | Unpublished| Yes                              |
| MGG_11608 | laccase-2                                  | −3.40               | 4.20E-159| Unpublished| No                               |
| MGG_09139 | laccase-1                                  | −2.32               | 1.44E-13 | Unpublished| Yes                              |

$^a$ELSD indicated expression level in SAGE database.

$^b$Yes indicating at least one putative MoAP1 binding site was identified in the promoter region of the gene while ‘No’ stand for non putative MoAP1 binding site was identified in the promoter region.

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Figure 8. Phenotype observation and pathogenicity assay of targeted gene deletion mutants. Ten down-regulated genes in the Moap1 mutant were selected for gene deletion and the phenotype of each mutant was compared and displayed. Gene ID numbers were sourced from www.broad.mit.edu/annotation/genome/magnaporthe_grisea/. a Log2 ratio (Moap1/Guy11) value stands for gene expression fold differences in the Moap1 mutants. b Co represents conidiation and c AP represents appressorium. d The diseased leaves were harvested at 5 dpi, and the results at 7 dpi displayed in Figure S12. e The letter Y stands for yes and N stands for no.

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Table 2. AP1 binding motif in the promoter region of the down-regulated genes.

| Gene_ID     | Description                                      | ELSD (a) (Moap1/WT) | p-Value | Putative MoAP1 binding sequence (b) |
|-------------|--------------------------------------------------|---------------------|---------|------------------------------------|
| MGG_12814.6 | Conserved hypothetical protein (MoAP1)          | 1e-75               |         | GCTTACTTC                          |
| MGG_01230.6 | Succinate-semialdehyde dehydrogenase (MoSuad)   | -4.11               | 0.0     | GTACTAA, TGAGTA, AGACTAA           |
| MGG_15157.6 | Acetyltransferase (MoAct)                        | -4.91               | 3e-20   | ATACTAA, TTACTCA                   |
| MGG_13464.6 | Laccase                                          | -7.61               |         | CAAGTCAGC                          |
| MGG_01682.6 | 4-aminobutyrate aminotransferase (MoAd)         | -5.17               |         | ACTGACTAG, TTAGTAAAG, TTACTCC      |
| MGG_02378.6 | Glutamate decarboxylase -1 (Mooye)              | -4.45               |         | TTAGTAAGT                          |
| MGG_10422.6 | C6 transcription factor OefC (Mooye)            | -3.81               |         | TTGAGTAAGT                         |
| MGG_13654.6 | Hypothetical protein (Mooye)                    | -9.59               |         |                                 |
| MGG_03817.6 | Metalloprotease 1 (Mooye)                       | -6.88               | 4e-26   | AGAGTAA                            |

(a)ELSD indicated expression level in SAGE database.
(b)Sequences in the table indicating the putative MoAP1 binding site that identified in the promoter region of the gene.

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Figure 9. The appressorium-like structure is unable to penetrate the onion epidermis or rice leaf sheath cells. (A and C) The appressorium-like structure develops at the hyphal tips. The appressorium-like structure was induced by placing the mycelia blocks on the onion epidermis cell for 24 hrs. (Scale bars equal 20 μm). (B and D) Onion epidermis cell penetration assay. The hyphae blocks were inoculated on the onion skin cells for 48 hrs. Penetration was observed using DIC microscopy. (Scale bars equal 20 μm). (E and F) Statistical analysis of appressorium-like structure development. A total of 300 hyphal tips were examined and counted. Dark column indicates Guy11, white column indicates the Mossadh or Moact mutant, and gray column indicates the Mossadh/SSADH or Moact/ACT complement strain. Error bar represents standard deviations and asterisks indicate that the differences were significant. (G) Rice leaf sheath penetration assay. Appressorium like structure formed by the Mossadh mutant and Moact mutant are incapable of penetration into the rice leaf sheath cells 48 hrs post injection (Scale bars = 20 μm).

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Figure 10. Pathogenicity test of Mossadh and Moact mutant strains on the wounded rice leaves. (A) Pathogenicity test of Mossadh mutants on the wounded rice leaves. The mycelia blocks of the wild type strain Guy11, Mossadh mutants, and the complemented strain were inoculated on the wounded rice leaves and then cultured under moist conditions with 28°C for 5 days. The wounded rice leaves with the CM agar plugs on was used as negative control. This experiment was performed three times with 10 pieces of rice leaves for each strain. Similar results were obtained in each test and this picture showed the representative result. (B) Pathogenicity test of Moact mutants on the wounded rice leaves. The mycelia blocks of Guy11, Moact mutants and the complemented strain were inoculated on the wounded rice leaves and observations made as above. (C) Pathogenicity test of the mutant strain by injection of hyphae fragments. The hyphae fragments of Guy11, the mutant strains and respective complemented strains were prepared as described in Materials and Methods. From left to right, leaves injected with water, wild type strain, Mossadh mutant, and Moact mutant. The arrowheads in black indicate injection sites with necrosis, while the arrowheads in white indicate injection sites without necrosis.

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ROS production serves many functions in eukaryotic cells, including those in cellular defense [13,69,69]. Its role in host resistance and pathogen invasion is highly dependent on the types of plant pathogen and host interaction [9,70,71,72]. In plants, the generation of ROS is regarded as one of the first responses to fungal invasion [9]. For pathogens and entophytes, the production of ROS also has an important role and the disruption of intracellular ROS homeostasis can result in functional defects. In the endophytic fungus *Epichloë festucae*, ROS acts as a negative regulator to prohibit excessive fungal proliferation and, thus, allow the fungus to maintain a mutalistic relationship with its host plant [73]. Furthermore, the deletion of *NOX* encoding NADPH oxidase or its regulatory subunit RacC leads to defects in ROS production, but the mutant remains highly pathogenic [73,74]. However, the deletion of the *NOX*-like gene in the ergot fungus *Claviceps purpurea* yields mutants with reduced conidial formation and pathogenicity [75]. Moreover, the NADPH oxidases of both *M. oryzae* and the gray mold fungus *Botrytis cinerea* are key for the generation of intracellular ROS and their deletions make them lose pathogenicity on host plants [46,76]. Conversely, deletion of the *AbTMPL* gene encoding a transmembrane protein in *A. brassicicola* led to hypersensitivity to the oxidative stress and excess oxidative bursts during conidiation and plant invasion, and ultimately impaired the virulence on green cabbage [77]. Thus, ROS generation seems to have different effects during fungi and plant interactions, and the regulation of ROS levels is essential for fungal development as well as virulence [46,69,72,73,76,77,78]. In this study, we highlighted the significance of intracellular ROS homeostasis in relation to fungal development. Because *MoAP1* was highly expressed in conidiation and the *Moap1* mutant exhibited abnormal conidiogenesis, excess ROS accumulation in conidia, and loss of pathogenicity, we concluded that *MoAP1* is involved in important mechanisms for balancing ROS levels during conidiation, and the disruption of intracellular ROS homeostasis is responsible for the loss of virulence.

In *M. oryzae*, deletion of the catalase-encoding gene (*MoCATB*) caused reduced pigmentation, which is similar to the *Moap1* mutants because of its hyperoxidant state [49]. In this study, we also observed that the activity of the secreted laccase was reduced in *Moap1* mutants, and that the addition of copper sulfate stimulated the laccase activity and restored melanin biosynthesis to the *Moap1* mutants. This indicated that the decreased laccase activity and the excess ROS levels resulted in less pigmentation in the *Moap1* mutants.

In fungi, secreted peroxidases are regarded as an important component in helping pathogens to detoxify host-derived ROS during plant-microbe interactions [15,16,60]. Recently, it was shown that the decreased expression of peroxidase genes likely resulted in a reduced ability to scavenge host-derived ROS and, thus, an attenuation of virulence [15,16,60]. We detected a deficiency of secreted peroxidase activity in the *Moap1* mutant by comparing CR discoloration and assaying the peroxidase activity in culture filtrates. In the comparison of the SAGE data, we identified several extracellular peroxidase-encoding genes displaying significant reductions in transcription (see Table S1), which suggested that these may result in the decreased peroxidase activity in the *Moap1* mutants. Furthermore, the activity of the laccase, which is involved in pathogenicity of certain fungi [79], was severely reduced in the *Moap1* mutants. Moreover, we also identified that the laccase activity was similarly decreased in the *Mossadh* and *Moact* mutants. Such results prompted us to question whether the decreased laccase activity in the *Mossadh* and *Moact* mutants was associated with that displayed in the *Moap1* mutant. Expression of two laccase genes (MGG_13464 and MGG_11660)

**Figure 11. ** *Mossadh* and *Moact* are hypersensitive to H$_2$O$_2$. (A) Phenotype of the *Mossadh* mutants under the oxidative stress. The wild type strain, *Mossadh* mutants and the complemented strain were inoculated on CM agar medium with or without 5 mM H$_2$O$_2$ and cultured at 28°C for 5 days. (B) Statistical analysis of mycelia growth rate with or without H$_2$O$_2$. The analysis was similar to Figure 3B. Error bars represent the standard deviations, and asterisks represent significant differences among Guy11, *Mossadh* mutants and the complemented strain (p<0.01). (C) Phenotype of the *Moact* mutant strains under the oxidative stress. Guy11, *Moact* mutants, and the complemented strain were inoculated on CM agar medium with or without 2 to 5 mM H$_2$O$_2$ and cultured at 28°C for 3 days. (D) Statistical analysis of mycelia growth. Error bars represent the standard deviations, and asterisks indicate that the differences among Guy11, *Moact* mutants and the complemented strain were statistically significant (p<0.01).

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was indeed down-regulated in the Moap1 mutants (see Figure S8B). Based on these results, we postulated that the decreased expression of MoSSADH and MoACT is responsible for the reduced laccase activity in the Moap1 mutants.

The identification of MoAP1-regulated genes by SAGE analysis revealed similar functional categories of genes that are down-regulated in the Moap1 mutant. Among these genes were some well-studied ones directly involved in pathogenicity and novel ones such as MGG_01230 and MGG_15157, which are likely to be involved in H2O2 tolerance. Disruption of either gene showed the complete loss of pathogenicity. The individual mutants also displayed similar morphogenesis defects, including less aerial hyphae but more compact mycelial growth, hypersensitivity to H2O2, decreased or loss of extracellular laccases and peroxidases activity, and reduced hyphal branching. These phenotypes are consistent with those observed in the Moap1 mutants, indicating that MoAPI has a direct role in regulating genes such as MoSSADH and MoACT to control growth and morphologies, as well as tolerance to H2O2 and ultimately pathogenicity. Likewise, disruption of MoAPI function may also affect other aspects such as protein translation and degradation, and secondary metabolism as evidenced by the SAGE data (see Table S1) that lead to the defect in pathogenicity.

Like most fungal pathogens, conidiogenesis and appressorium development are key steps in the colonization of host plants by M. oryzae. After attaching to the host surface, the conidia began to produce germ tubes and then developed a specialized infection structure, an appressorium with 8 MPa turgor pressure at the tip of the tube, to help the fungus penetrate host cell barriers [36]. The above processes are controlled by a precise developmental program in response to stimuli from the host and environment. In this program, conidiogenesis is a complex process that involves a cascade of morphological events. In M. oryzae, MoAPI disruption did not affect the developmental stages, such as hyphal growth, appressorium formation and penetration, but severely affected the ability to produce conidia and infectious hyphae growth. This observation was consistent with the results of qRT-PCR, which suggests that MoAPI is a stage-specific regulator during conidiation and infection. In an effort to fully understand the developmental defects of the Moap1 mutants, we screened the SAGE data and found that the zinc finger transcription factor MoCOS1 [54], a determinant of conidiophore formation, was also severely down-regulated in the Moap1 mutants (see Figure S8A), indicating the possibility that it is responsible for decreased conidiation. A detailed analysis of the MoAPI downstream genes revealed that the mutation of either MoSSADH or MoACT gene caused complete loss of the ability to generate conidia. Taken together, these findings suggest that the transcriptional factor MoAPI controls conidiation via a complex mechanism, involving the regulation of MoCOS1, MoSSADH, and MoACT expression.
Certainly, we still could not rule out the possibility that some unknown gene(s) regulated by MoAP1 are also responsible for the conidiation in M. oryzae.

Because both the Mossadh and Moact mutants lost the ability to produce conidia, the pathogenicity assay was performed through inoculation of the hyphal tip plug that showed the complete loss of pathogenicity for the two mutants. It is known that appressorium-mediated penetration is required for full virulence of M. oryzae [80,81,82,83,84]. Recent studies of MoCos1 and other transcription factors revealed that the hyphal tips of the MoCos1 mutant could develop appressorium-like structures on the host surface and cause infection [54,59]. It was presumed that these transcription factors might play a role in an unknown mechanism in mycelia-mediated infection. To fully understand the loss of pathogenicity, we compared the formation of the appressorium at the hyphal tips between the wild-type strain and the two mutants, and found that all of them could form such appressorium-like structures. This finding is similar to the results of Kim et al. [58]. However, such hyphae-driven appressorium by the Mossadh and Moact deletion mutants were severely limited in their ability to penetrate into onion skin cells and rice leaf sheath cells and cause disease symptoms on leaves. This indicates that the hyphae-driven appressorium-mediated penetration may be functionally similar to penetration by normal appressorium. In the Mossadh and Moact mutants, the loss of pathogenicity might be due to the loss in the penetration ability.

The γ-aminobutyrate (GABA) shunt is a metabolic pathway that bypasses two successive steps of the tricarboxylic acid (TCA) cycle and is present in many organisms [85,86,87]. In plants, the activity of this pathway is predominantly associated with the response to abiotic and biotic stresses [88]. Recent studies have considered that a mutation of AtSSADH in Arabidopsis thaliana leads to growth abnormalities, hypersensitivity to the environmental stress, and ROS accumulation on the trichomes [87]. However, the cellular function of succinic semialdehyde dehydrogenase is still uncharacterized in fungal pathogens. In M. oryzae, our findings revealed that the Mossadh mutant displayed retarded mycelial growth, hypersensitivity to oxidative stress, and dramatically reduced aerial hyphae. Further, the mutant completely lost the ability to cause infections. We initially suspected that the loss of pathogenicity in the Mossadh mutant might be due to the inability to penetrate the host cell, but after injection of mycelial fragments in the rice leaf cell, the Mossadh mutant still could cause necrosis around the injection site. Together with the fact that stress-tolerance mechanisms of plant pathogens play an important role in virulence [14,15,16,60] and the sensitivity of the Mossadh mutants to H2O2, we postulated that M. oryzae MoSsadh might be responsible for both the penetration and invasive growth in planta, and the defects in stress-tolerance may be a result of the restriction of invasive hyphae growth in planta, causing the complete loss of pathogenicity.

**Materials and Methods**

**Fungal strains and growth conditions**

*M. oryzae* strain Guy11 was used as wild type throughout this work. Both Guy11 and its derivative mutants were cultured on complete medium (CM) [89] for 3–15 days at 28°C to assess the growth and colony characteristics. Fungal mycelia were harvested from liquid CM and used for genomic DNA and RNA extractions. To observe the vegetative growth under the oxidative stress condition, H2O2 (Aldrich, 323381, 3 wt. %) was mixed in solid CM, and diameters of fungal colonies were measured after 3 to 5 days as indicated. For the activation of the laccase activity in the Moap1 mutants, copper sulphate was amended in the CM medium for 1 mM at final concentration. Cell wall integrity assay was performed by growing strains in Congo Red (CR, Aldrich, 860956) amended CM medium (200 μg/ml) for 5 days. For conidial collection, strains were normally maintained on corn meal (RDC) medium [43] at 28°C for 10 days and then transferred to constant fluorescent light condition to promote conidiation for another 3–5 days. Conidia were obtained by rubbing mycelia with water followed by filtration through Miracloth (Calbiochem, San Diego, USA). For mycelial growth assay, strains were inoculated on the liquid complete medium for 48 hrs and then transferred to the Petri dish for photograph. To observe conidiophore development and conidiation, strains were inoculated on RDC for 5 days and mycelia were rubbed with a glass rod before transferring to the constant fluorescent light condition to promote conidiation for another 2 days. S. cerevisiae strains were grown in SD medium supplemented with appropriate amino acids and with glucose (3% (w/v)) or galactose (2% (w/v)). All growth assays were repeated for three times, with three replicates each time.

**Characterization of gene disruption mutants**

Vegetative growth of Moap1 mutants was measured on complete agar medium for 5 days, while the vegetative growth of Mossadh and Moact mutants were measured on the same medium for 4 days. All the experiments were performed with triple replicates in three independent experiments. The ability to produce conidia was measured by counting the numbers of conidia from 10-day old RDC plates as described previously [43]. Conidia were collected by rubbing the plate with 5 ml of sterilized distilled water. Conidia were counted using a hemacytometer under a microscope and their germination and appressorium formation were measured on a hydrophobic coverslip. Conidial suspensions of 30 μl (1×10^6 spores/ml) were dropped onto a coverslip and placed in a moistened box at 28°C. After 8 hrs of incubation, the percentage of conidia germinating and germinated conidia-forming appressoria was determined by microscopic examination of at least 100 conidia. This test was done at least three times, each with three replicates.

**Yeast yap1 mutant complementation**

*S. cerevisiae* BY4741ΔYML007w (yap1) and the strain from which it was derived, BY4741 (MATa his3Δ1 leu2Δ0 met2Δ0 ura3Δ0) were obtained from Euroscarf. The full-length of *M. oryzae* Moap1 cDNA (1.7 kb) was amplified using primers pairs FL2700/F and FL2701/R. The PCR products, digested with EcoRI and SpH1, were cloned into pYES2 (Invitrogen) and transformed into BY4741ΔYML007w. Colonies were selected on SD medium lacking uracil, and the wild type yeast strains BY4741 as well as the *yap1* deletion mutant BY4741ΔYML007w transformed with empty pYES2 vector were used as a control. Transformed yeast cells were grown on SD medium without uracil containing either glucose (3% (w/v)) or 2% galactose respectively. Five-microliter drops from serial dilutions from cultures with an OD600 of 0.5 were spotted on plates with and without 0.3 mM H2O2 and grown for 3 days at 30°C.

**Targeted gene disruption and complementation of MoAP1 and MoAP1 target genes**

For constructing the Moap1 gene replacement construct, a 1.0-kb upstream flanking sequence fragment and 0.9-kb downstream flanking sequence was amplified from *M. oryzae* genomic DNA by PCR using primer pairs FL1992(F)/FL1993(R) and FL1994(F)/
FL1995(R), respectively (Table S2). The two flanking sequences were joined together by overlap PCR with primer pair FL1992(F) and FL1995(R), and the amplified ~2 kb fragments were purified and cloned into a pMD19-T vector (Takara, Co, China) to generate plasmid pMD19-Moap1. The Hygromycin resistance gene cassette [90] was prepared by PCR amplification with primer pair FL1111(F)/FL1112(R) (Table S2) and inserted in the plasmid pMDT-Moap1 at the PmeI enzyme site to generate the final disruption construct pMD-Moap1-HPI. The 3.4-kb fragment was amplified with FL1992(F) and FL1995(R) primers and transformed into Guy11 protoplasts. The protoplast-mediatated transformation of M. oryzae Guy11 was carried out as described [88]. Transfomants were selected on solid CM agar medium supplemented with 300 µg/ml hygromycin B. To identify the gene-deleted mutants, Hygromycin B resistant transformants were screened using primers FL2382(F)/FL2383(R) (Table S2). The mutants were further verified by Southern hybridization. To generate the complemented construct of the Moap1 mutant, a 4.4 kb DNA fragment including the putative promoter and the coding sequence was amplified and inserted into the plasmid pCB1352, according to Zhang et al. [88]. Disruption of Moap1 target genes was performed similarly with primer pairs listed in the Table S2.

Nucleic acid manipulation and Southern blotting

DNA extraction was performed as described by Talbot and associates [88], while gel electrophoresis, restriction enzyme digestion, ligation, and Southern blot hybridization were performed using standard procedures [91]. DNA hybridization probes were random primer labeled with digoxigenin-11-dUTP using DIG-High prime according to the manufacturer’s instructions for digoxigenin high-prime DNA labeling and the detection starter kit (Roche Applied Science, Penzberg, Germany). Total RNA was isolated from frozen fungal mycelia using the RNA extraction kit (Macherey-Nagel, Bethlehem, PA, USA) following the manufacturer’s instructions. To measure the relative abundance of gene transcripts, RNAs were extracted from mycelia grown in CM liquid medium for 2 days at 28°C in a 150-rpm orbital shaker. To measure the relative abundance of Moap1 transcripts during diverse fungal developmental stages, the total RNA samples were extracted from mycelia grown in CM liquid medium, conidia and plants inoculated with the conidia of Guy11 (1×109 spores ml⁻¹) for 8, 24, 48 and 72 hrs, respectively, by the method described above. The primer sets used to detect transcripts of Moap1 and its related genes from M. oryzae are listed in Table S2.

Quantitative RT–PCR, RT–PCR, and gene expression analysis

For RT-PCR and quantitative real time RT-PCR (qRT-PCR), 5 µg of total RNA were reverse transcribed into first-strand cDNA using the oligo(dT) primer and M-MLV Reverse Transcriptase (Invitrogen). Confirmation of deletions and reintroduction of Moap1, MoSAHDD and MoACTF genes were made with primer pairs FL2382(F)/FL2383(R), FL6745(F)/FL6746(R), and FL6749(F)/FL6750(R) (Table S2). 32 cycles of RT-PCR were run on a Bio-Rad PTC2000 Peltier Thermal Cycler. The stable expression actin gene (MGCC_03982.5) amplified by primer pairs FL474(F)/FL475(R) (Table S2) was used as internal control. qRT-PCR reactions were performed following previously established procedures [16]. To compare the relative abundance of target gene transcripts, the average threshold cycle (Ct) was normalized to that of actin gene for each of the treated samples as 2^{-ΔΔCt}, where ΔΔCt = (Ct_target gene–Ct_actin gene) test condition−(Ct_target gene–Ct_actin gene) CM [16]. qRT-PCR was performed with three independent pools of tissues in three sets of experimental replicates and primers pairs used in this section were listed in Table S2.

Plant infection assays

Onion epidermis penetration assays were performed using the method as previously described [43]. Plant infection assays were performed on four-week old susceptible rice seedlings (O. sativa) CO-39 or seven-day old barley seedlings (Four arris) by spraying 4 ml of the conidial suspensions with a sprayer. Inoculated plants were placed in a moist chamber at 28°C for first 24 hrs in darkness, and then transferred back to another moist chamber with a photoperiod of 12 hrs under fluorescent lights. The disease severity was assessed at 5 or 7 days after inoculation. Approximately six centimeter long diseased rice blades were photographed to evaluate the virulence of the mutants. For determining the pathogenicity of the mutants without conidia, mycelia tip plugs of the wild type strain Guy11, Mossadh and Moact mutants were inoculated on the healthy or wounded rice leaves or barley leaves for 5 or 7 days and keps in the same condition as described above. For the infiltration infection assay, 0.1 gram mycelia of the tested strains was broken into pieces using a glass rod and 50 µl of each suspension were injected into the leaves of 4-week-old rice plants and cultured for 5 or 7 days under the condition as described above. These experiments were all replicated three times.

ROS and superoxide detection

Intracellular ROS levels of M. oryzae were monitored during the infection related structure formation using the oxidant-sensitive probe dihydrodorhodamine-123 (Molecular Probes, Carlsbad, CA) and nitroblue tetrazolium (NBT) as previously described [16]. For dihydrodorhodamine-123 staining, drops of conidial suspension (30 µl) were placed on the coverslips and cultured for up to 24 hrs. At each interval, the water surrounding the conidia was removed carefully and replaced with final concentration of 50 µM dihydrodorhodamine-123 (Merck, Whitehouse Station, NJ) at 28°C for 2 hrs, then rinsed twice with phosphate-buffered saline and viewed under a fluorescence microscope (Olympus IX71) equipped with a digital camera by short exposure to UV light. NBT staining was performed as described [92]. Superoxide production during conidial germination and infection related structure formation was viewed by microscopy.

Measurement of the extracellular enzyme activities

Laccase activity on solid medium was measured as described [92] with little modification. A 5×5 mm hyphal tip plug was inoculated on CM medium supplemented with 0.2 mM 2, 2'-azino-di-3-ethylbenzathiazoline-6-sulfonate (ABTS, Sigma) for 3 days. The assay for the activation of the laccase activity was performed by addition of 1 mM copper sulphate to the CM medium containing 0.2 mM ABTS and cultured under 28°C for 4 days. For detection of peroxidase secretion, a 5×5 mm hyphal tip plug was placed on CM medium containing 200 µg/ml Congo Red for 5 days. The measurement of peroxidase and laccase activities in culture filtrates was performed as described [88].

CFW and DAPI staining

Calcofluor staining using Fluorescent Brightener 28 (10 µg/ml, Sigma-Aldrich) for the microscopy of mycelial branches was performed as described [93]. Both the mutants and the wild type were inoculated on the coverslips that contain a thin layer of agar
medium and cultured for 48 hrs. Mycelial tip plugs were removed and stained with 10 μg/ml CFW for 10 min in darkness, rinsed twice with PBS and viewed under a fluorescence microscope (Olympus IX71). For the localization of MoAP1, conidia of Gny11 transformed with plasmid pCB1532::TrpC::Moap1::GFP was treated with 2 mM H2O2, stained with DAPI (50 μg/ml, Sigma) for 10 min, and visualized under a microscope (Olympus IX71). GFP fluorescence was detected using a 450 to 490-nm excitation filter and a 520-nm barrier filter and the DAPI fluorescence was detected under UV light using a 360 to 400 nm excitation filter.

Bioinformatics analysis

The full sequence of MoAP1 was downloaded from the M. oryzae database (www.broadinstitute.org/annotation/genome/nanoglyphus_gnome). Yap1 homology sequences from different organisms were obtained from GenBank (www.ncbi.nlm.nih.gov/BLAST) using the BLAST algorithm [94]. Sequence alignments were performed using the Clustal_W program [95] and the phylogenetic tree was viewed using Mega3.0/Beta program [96]. Orthologs were identified between M. oryzae predicted proteins and proteins in the GO database [97] via searching reciprocal best hits with the following cut-off: e-value, 1.0e-3, and identity, 20%. Results from local alignment using BLAST and prediction of signal peptides from SignalP 3.0 software [98] and a manual literature review were used to make final assignments to GO functional categories. Primers used in this study were designed by using Primer3 Input (version 0.4.0) and commercially synthesized (envigen Co., Shanghai, China). To predict API binding sites, yeast API binding motif sequences (MTTACGTAAK, TTAGTMAGC and TTASTMA) [99,100,101] were used to search in the 1000 bp-upstream sequences set of the up- and down-regulated genes from SAGE, and no more than one mismatch was allowed.

Supporting Information

**Figure S1** Comparison of API protein conserved domains and a dendrogram of fungal API proteins. (A) Comparison of the conserved bZIP domains of fungal API proteins arranged by Clustal W program. Identical amino acids residues are shaded. (B) Alignment of the conserved c-CRD domain of fungal API proteins. Asterisks indicate the conserved cysteine residues. (C) A dendrogram of fungal API proteins. The phylogenetic tree was created with Mega3.0 beta by the established parameter in the program. GenBank accession numbers are as follows: M. oryzae MoAP1 (EDK00544), F. oxysporum FoAPI (XP_388976), C. heterostrophus ChAPI (AAP64131), P. triticicrepitis PtAPI (XP_001931984), A. oryzae AorAPI (BAE92562), S. cerevisiae ScYAP1 (CAA41536), U. maydis UmayYAP1 (XP_758338) and C. albicans Cap1P (EAK94712).

**Figure S2** MoAPI complements the H2O2 sensitivity of a S. cerevisiae Δ yap1 mutant. The growth of S. cerevisiae BY4741::pYES2, BY4741DYML007w+pYES2, BY4741DYML007w+pYES2::Moap1 was tested on SD plates with glucose (top left panel), galactose (top right panel), SD plates with glucose supplemented with 0.3 mM H2O2 (bottom left panel), and with glucose supplemented with 0.3 mM H2O2 (bottom right panel). Found at: doi:10.1371/journal.ppat.1001302.s001 (1.75 MB TIF)

**Figure S3** The MoAPI phase specific expression, targeted gene replacement and complementation. (A) The phase specific expression of MoAPI. The expression of MoAPI was measured by quantitative real-time RT-PCR with cDNA from samplings for infectious growth, vegetative growth, and conidia. The relative abundance of Moap1 transcripts during infectious growth (from ungerminated conidia to in planta fungal cells 72 hpi) was normalized by comparing with vegetative growth in liquid CM (Relative transcript level = 1). Each sample was harvested from 10 plants and three independent experiments, each with three replicates, were performed. Significant differences are presented in the figure (P < 0.01), and the error bar represents the standard deviation. (B) MoAPI targeted gene replacement. A 1.89-kb fragment of the Moap1 coding region was replaced with a 1.4-kb fragment containing the hygromycin B resistance cassette to create the Moap1 mutant. The DNA fragment at the inner space of MoAPI deletion region was used as the probe to validate the Moap1 deletion transfectants by PCR amplification and Southern hybridization analysis (scale bar = 1 kb). (C) Genomic PCR was used to validate the deletion of Moap1 gene and reintroduction of Moap1 coding region to complement the mutant strain. (D) Semi-quantitative RT-PCR was carried out to confirm the deletion and reintroduction of MoAPI gene. Complete inactivation of Moap1 transcription in the deletion mutants was verified by reverse transcription (RT)-PCR using cDNA of the wild type strain, the Moap1 mutants, and the complemented strain. (E and F) Southern hybridization analysis was used to validate the deletion of the MoAPI gene and the addition of a single copy integration of the HPH gene. The arrowhead in E (left) showed a single band hybridized by the HPH probe in the mutant. No band was present in the wild type strain.

Found at: doi:10.1371/journal.ppat.1001302.s003 (1.24 MB TIF)

**Figure S4** Hyphal branching reduction in the Moap1, Massadh, and Moact mutants. (A) Branching patterns of mycelia on agar media containing coverslips 48 hrs after incubation. Frequent branching occurs at the mycelia of wild type while no or a few hyphal branches were observed in the Moap1 mutants. Calcofluor white staining is used to indicate the position of the mycelia. Bar = 50 μm. (B) Branching patterns of mycelia on agar media containing coverslips 48 hours after incubation. Frequent branching occurs at the mycelia of wild type while no or a few hyphal branches were observed in the Massadh mutants. Calcofluor white staining indicates the position of the mycelia. Bar = 50 μm. (C) Branching patterns of mycelia on agar media containing coverslips 48 hrs after incubation. Frequent branching occurs at the mycelia of wild type while no or a few hyphal branches were observed in the Moact mutants. Calcofluor white staining is used as the indicator for the position of the mycelia. Bar = 50 μm.

Found at: doi:10.1371/journal.ppat.1001302.s004 (2.19 MB TIF)

**Figure S5** Subcellular localization of MoAPI in the presence of H2O2. For green fluorescence observation, both conidia of wild type strain Gny11 (WT) and WT transformed with pCB1531::TrpC::Moap1::GFP were treated with DAPI and then observed under an Olympus microscope with a specific filter set as described in the materials and methods. For subcellular localization, conidia of WT transformed with pCB1531::TrpC::Moap1::GFP was treated with or without 2 mM H2O2 and then observed as described above.

Found at: doi:10.1371/journal.ppat.1001302.s005 (0.70 MB TIF)

**Figure S6** Pathogenicity test of Moap1, Massadh and Moact mutant strains on the barley leaves. (A) Pathogenicity test of Moap1 mutant on barley leaves. 4 ml conidial suspension (1×106 conidia/ml) of each strain was sprayed on seven-day-old barley seedlings (Four airs) and cultured as described in Figure 5A and the results were observed at 7 dpi. The barley leaves spraying of gelatin was used as negative control. (B and C). Pathogenicity test of Moap1, Massadh and Moact mutant on barley leaves. Mycelia blocks of the
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