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Systematic Analysis of Zn₂Cys₆ Transcription Factors Required for Development and Pathogenicity by High-Throughput Gene Knockout in the Rice Blast Fungus

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Abstract

Because of great challenges and workload in deleting genes on a large scale, the functions of most genes in pathogenic fungi are still unclear. In this study, we developed a high-throughput gene knockout system using a novel yeast-*Escherichia-Agrobacterium* shuttle vector, pKO1B, in the rice blast fungus *Magnaporthe oryzae*. Using this method, we deleted 104 fungal-specific Zn₂Cys₆ transcription factor (TF) genes in *M. oryzae*. We then analyzed the phenotypes of these mutants with regard to growth, asexual and infection-related development, pathogenesis, and 9 abiotic stresses. The resulting data provide new insights into how this rice pathogen of global significance regulates important traits in the infection cycle through Zn₂Cys₆TF genes. A large variation in biological functions of Zn₂Cys₆TF genes was observed under the conditions tested. Sixty-one of 104 Zn₂Cys₆ TF genes were found to be required for fungal development. In-depth analysis of TF genes revealed that TF genes involved in pathogenicity frequently tend to function in multiple development stages, and disclosed many highly conserved but unidentified functional TF genes of importance in the fungal kingdom. We further found that the virulence-required TF genes *GPF1* and *CNF2* have similar regulation mechanisms in the gene expression involved in pathogenicity. These experimental validations clearly demonstrated the value of a high-throughput gene knockout system in understanding the biological functions of genes on a genome scale in fungi, and provided a solid foundation for elucidating the gene expression network that regulates the development and pathogenicity of *M. oryzae*.

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Data Availability: The authors confirm that all data underlying the findings are fully available without restriction. All relevant data are within the paper and its Supporting Information files. And RNA-sequencing data are also available at NCBI's Gene Expression Omnibus (GEO accession number GSE57146).

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Introduction

Magnaporthe oryzae is the best-studied phytopathogenic fungus, which was voted first in the top 10 list of fungal plant pathogens by an international community of molecular plant pathologists [1]. The importance of this filamentous ascomycete fungus is not only owing to the fact that the rice blast disease caused by the fungus is the most destructive disease of rice throughout the world, which typically leads to 20-30% losses and even complete loss in grain production during regional epidemics [1,2], but also to its being a primary model in the study of hostfungal pathogen interactions [3]. The rice blast fungus has a complicated life cycle including hyphal growth, conidiogenesis, conidial germination, appressorium formation and plant infection, which provides substantial biological information in eukaryotic development and pathogenesis. The rice blast fungus is highly amenable to molecular genetic manipulation, and the functions of numerous genes are identified by gene knockout or ectopic insertion [4-6].

Transcription factors (TFs) are proteins that bind to specific DNA sequences, thereby controlling the flow of genetic information from DNA to mRNA. In the rice blast fungus, more than 522 putative TF proteins have been identified from 12,991 M. oryzae proteins (www.ftfd.snu.ac.kr; www.broadinstitute.org). Therefore, nearly 4.02% of genes in the genome code for TFs, which makes this family the single largest family of *M. oryzae* proteins. Recent functional analyses of single or several TF genes revealed their critical biological roles in fungal development, pathogenesis and response to the environment, for instance, in hyphal growth (MNH6, MSTU1, MoCRZ1 and MoSWI6) [7-10], conidiogenesis (COM1, CON7, COS1, MNH6, and MoHOX2/HTF1) [7,11–16], conidial germination (TRA1) [17], appressorium formation (MoLDB1, MoSOM1 and MoCDTF1) [18,19], plant infection (COM1, MNH6, MIG1, MST12/MoHOX8, and MoSfl1) [7,11,16,20-22], and response to oxidative stress(-MoATF1 and MoAP1) [23,24] or light (MgWC-1) [25]. However, the biological functions of most TFs have not been revealed, mainly because it is difficult to delete genes on a large scale and

Author Summary

Magnaporthe oryzae is not only the fungus causing the rice blast disease, which leads to 20-30% losses in rice production, but also a primary model pathosystem for understanding host-pathogen interactions. However, there is no high-throughput gene knockout system constructed, and little is known about most of the genes in this fungus. We developed a high-throughput gene knockout system, and using this system, we obtained null mutants of 104 fungal-specific Zn₂Cys₆ transcription factor (TF) genes by screening 8741 primary transformants in M. oryzae. We analyzed the functions of these TF genes in development, pathogenesis, and stress responses under 9 conditions. We found that 61 Zn₂Cys₆ TF genes play indispensable and diversified roles in fungal development and pathogenicity. CNF1 is the first reported TF gene that strongly and negatively regulates asexual development in the rice blast fungus, and CCA1, CNF1, CNF2, CONx1, GPF1, GTA1, MoCOD1 and PCF1 are required for pathogenicity. We further found via RNA-seq that GPF1 and CNF2 have similar mechanisms in gene expression regulation related to pathogenicity. The resulting data provide new insights into how Zn₂Cys₆ TF genes regulate important traits during the infection cycle of this rice blast pathogen.

because a high-throughput gene knockout method has not been previously established in M. oryzae.

Several loss-of-function techniques, such as homologous recombination [4], insertional mutagenesis [6] and RNA interference [26], have been used to investigate gene functions in fungi. Although insertional mutagenesis and RNA interference have been effectively used to inactivate gene functions, they all have limitations that prevent them from becoming high-throughput gene knockout techniques, including 1) inability to cover the genome and the heavy workload to clone genes in the insertional mutagenesis method [6], and 2) the silencing of potential unintended targets by the RNA interference method [26]. Gene knockout through DNA homologous recombination is a primary technique used to inactivate gene functions. More than 650 putative TF genes were disrupted in Fusarium graminearum using homologous recombination [27]. In that study, the double-joint PCR method [28] was used to generate gene knockout constructs, which were then transformed into fungal protoplasts. In Neurospora crassa, homologous recombination was developed into a high-throughput gene knockout technique [29], in which gene knockout constructs generated by making use of a yeast recombinational cloning method were transformed into conidia by electroporation. Yeast recombinational cloning is a more suitable method than double-joint PCR for high-throughput gene knockout study. However, protoplast transformation is an elaborate, time-consuming and inefficient method, and electroporation of germinating conidia has not been built in the rice blast fungus and many other filamentous fungi. The frequency of homologous recombination after transformation is often quite low in filamentous fungi, typically <10% in *M. oryzae* [30]. In addition, mutant screening is a tedious and inefficient step in highthroughput gene knockout studies. The implementation of KU70/ KU80 (mus-51/mus-52) mutations as gene knockout background strains greatly increases the frequency of homologous recombination in N. crassa, Aspergilli and other filamentous fungi, and reduces the workload in screening [29,31-34]. However, the null mutants obtained from KU70/KU80 deletion background strains need to be complemented with native KU70/KU80 genes before analyzing their mutant phenotypes [29]. These shortcomings limit the application of the above methods in research in fungal functional genomics. Until now, there is still a surge in interest in functional genomics research through the systematic mutagenesis of identified genes sequenced in the genomes of a large number of fungi (http://www.ncbi.nlm.nih.gov/genome/).

To learn the biological functions of TFs at the genome level, we constructed a high-throughput gene knockout method that enables the rapid knockout of large numbers of genes in the rice blast fungus. In this method, gene knockout vectors are built by a yeast recombinational cloning method using a high-throughput way, DNA transformations are performed by Agrobacterium tumefaciens-mediated transformation (ATMT), and the null mutants are identified by the negative (GFP)/positive (a resistant gene) screening system through a novel yeast-Escherichia-Agrobacterium shuttle vector, pKO1B. With this system, we deleted 104 putative fungal-specific Zn_2Cys_6 TF genes in *M. oryzae*. The null mutants were then examined for their phenotypes in development, pathogenicity and responses to stress conditions. In particular, many Zn₂Cys₆ TF genes required for fungal growth, asexual development, conidial germination and appressorium formation, pathogenicity, and response to stress were identified. We further identified the genes regulated by GPF1 and CNF2 via RNAsequencing (RNA-seq) and found that GPF1 and CNF2 have similar mechanisms in the regulation of gene expression related to fungal pathogenicity. Our findings will provide new insights into the transcriptional regulation of fungal development and pathogenicity, and a way to study the biological functions of genes in fungi at the genome level.

Results

Strategy of the high-throughput gene knockout system via pKO1B vector in *M. oryzae*

Three major challenges in high-throughput gene knockout of fungi are 1) how to build gene-deletion cassettes quickly, 2) how to transfer DNA into fungal cells easily, and 3) how to identify null mutants from the large number of transformants efficiently. To solve these problems, we developed a high-throughput gene knockout system using a newly designed yeast-*Escherichia-Agrobacterium* shuttle vector, pKO1B, (Figure 1A). Through this plasmid, three techniques suitable for high-throughput manipulation (yeast recombinational cloning to build gene-deletion cassettes, ATMT to transfer DNA into fungal cells and dual selection system to identify null mutants) were combined into the high-throughput gene knockout system.

pKO1B is a vector built on the framework of the binary vector pCAMBIA1300 (www.cambia.org). It contains the URA3-2micro2_origin sequence from a yeast plasmid pYES2 (Invitrogen, USA) and eGFP gene under the control of a strong promoter of M. oryzae H3 histone gene [35] (Figure 1A). pKO1B has a new characteristic: the ability to be replicated in three organisms, Saccharomyces cerevisiae, Escherichia coli and A. tumefaciens. Through the gapped vector pKO1B, three DNA fragments (5' and 3' flanking fragments of the targeted gene and a resistant gene fragment) could be merged into a gene-deletion cassette in one step by yeast recombinational cloning (Figure 1B). The knockout vectors obtained were transformed into A. tumefaciens without transferring the gene-deletion cassettes from a yeast plasmid to another binary Agrobacterium plasmid. The gene-deletion cassettes in pKO1B were transformed into fungal cells using the ATMT method. When the gene-deletion cassettes were ectopically integrated into fungal genomic DNA, GFP was activated in ectopic insertional transformants and was easily observed under a



Figure 1. Overview of the high-throughput gene knockout system in fungus. (**A**) Features of a new binary yeast-*Escherichia-Agrobacterium* shuttle vector, pKO1B. (**B**) Building of gene-deletion cassettes in pKO1B by yeast recombinational cloning. The 5' and 3' flanking fragments of the targeted genes were separately amplified from genomic DNA with primers 5f/5r and 3f/3r. Primers 5r and 3f have 5' tails homologous to the *SUR* cassette, whereas those for 5f and 3r are homologous to the vector. The two flanks were cotransformed into yeast along with the *SUR* cassette and gapped pKO1B. Homologous recombination created the circular knockout vector, and the final knockout vector was subsequently transformed into A. *tumefaciens*. (**C**) Deletion of the targeted gene. The gene-deletion cassette was transformed into the fungal cells via ATMT. Homologous recombination created three types of transformants: null mutants, ectopic insertion transformants, and null and ectopic insertion mutants. The *GFP* gene was discarded in the null mutants. Primers p1/p2 or p3/p4 were used to identify the unique recombinational DNA fragment, indicating a knockout event. (**D**) The transformants are screened for GFP fluorescence under a microscope. Putative null mutants do have not GFP fluorescence, but ectopic transformants on cetopic transformants are screened by double PCR for the targeted gene using the *β-tubulin* gene as a positive control. The wild-type strain or ectopic transformants produced a characteristic band, indicating the targeted gene, while the null mutants have a 1.2–2.0 kb band on an electrophoretic gel, while the wild-type strain and the ectopic transformants do not. doi:10.1371/journal.ppat.1004432.g001

fluorescence microscope as a negative selective marker, but not in null mutants (Figure 1C and 1D).

To screen null mutants efficiently, the genomic DNAs of the transformants without green fluorescence were isolated using an improved CTAB method in a high-throughput way (shown in Materials and Methods) and then were detected for the targeted

gene and β -tubulin gene by double PCR. If the targeted gene was deleted in a mutant, only one band for β -tubulin appeared as a positive control on an electrophoretic gel; otherwise, there were two bands with one for the targeted gene and the other for β -tubulin in ectopic insertion transformants (Figure 1C and 1E). For those null mutants identified in the above negative screening PCR,



Figure 2. Analysis of Zn₂Cys₆ transcription factor mutant phenotypes in fungal development stages. (**A**) Number of TFs showing multiple mutant phenotypes. (**B**) Number of mutants showing mutant phenotypes in each developmental stage. (**C**) Venn diagram showing the number of mutant phenotypes. The phenotypes included vegetative growth (conidial germination, colony growth, pigmentation and mycelial appearance), conidiation (asexual reproduction), appressorium formation and pathogenicity to rice and barley. doi:10.1371/journal.ppat.1004432.q002

we then continued to search for the unique recombinational DNA fragment that indicated a knockout event and only appeared in the null mutants by PCR. In this PCR, one primer was limited in the genomic DNA outside of the 5' or 3' flanking fragment of the targeted gene, and the other primer was limited in the resistant gene of the gene-deletion cassettes (primers p1+p2 or p3+p4 in Figure 1C). There was one band 1.2-2.0 kb in length on the electrophoretic gel that appeared in null mutants. Otherwise, there was no band appearing in the ectopic transformants and wild-type strain (Figure 1F). Gene knockout and ectopic insertion maybe happened coincidentally (Figure 1C). To identify those mutants that were undetected in previous screening steps, copies of transformed gene deletion cassettes in null mutants were confirmed by qPCR after comparison with the wild-type strain using β -tubulin as a control. The null mutants had one copy of the gene deletion cassette, while the null+ectopic transformants had more than two copies and the wild-type strain none. Finally, the mutants containing a single copy of the gene deletion cassette were considered null mutants.

TFs containing fungal-specific Zn2Cys6 zinc finger and fungal TF MHR Domain are the largest group of TFs in the rice blast fungus. The 163 putative fungal-specific Zn₂Cvs₆ genes, previously annotated at the Fungal Transcription Factor Database (http://ftfd.snu.ac.kr/index.php) [36] or obtained by BLAST searches at NCBI under organism item (rice blast fungus, taxid: 318829) with the Conserved Domain program [37], were selected to generate gene deletion mutants (Table S1 in Text S1). As a result, 104 TF genes were identified to be deleted in M. oryzae (Table S2 and Table S3 in Text S1). The knockout rate of 104 genes was 15.03±12.02% (from 0.68 to 62.07%). The causes for tens of TF genes not deleted in this experiment lie in the following. 1) Knockout vectors of twenty-one genes had not been constructed. 2) For those genes with a knockout rate less than 1%, more transformants need to be screened. 3) Because the SUR gene was selected as a resistance gene, the mutants must be screened on DCM medium. But the mutants of TF genes that are involved with amino acid metabolism cannot grow on DCM medium. 4) Some genes may be essential genes.

Phenotypic analyses of fungal-specific Zn₂Cys₆ transcription factor deletion mutants at developmental stages

We analyzed the phenotypes of the $104 \text{ Zn}_2\text{Cys}_6$ TF null mutants at multiple developmental stages that *M. oryzae* likely encounters during its infection cycle in rice. These phenotypes

analyzed included: developmental characteristics (mycelial growth, colony color and mycelial shape; conidiation; conidial germination; and appressorium formation) and virulence to plants (barley leaf explants, rice leaf explants and rice seedlings). A substantial fraction of the mutants (58.7%, 61/104) clearly displayed visible phenotypes (Figure 2A). The results (Figure 2B) revealed that 27 of the 104 TF genes studied were involved in mycelial growth (Figure S1), 25 TF genes in conidial production, 12 TF genes in conidial germination and 10 TF genes in appressorium formation, and 5 TF genes were involved in pathogenicity in barley (Figure S2), while 7 TF genes were involved in pathogenicity in rice (Figure S3). Table 1 and Table S4 in Text S1 provide a complete list of the phenotypic analyses. Among these mutants displaying clearly visible mutant phenotype changes, 42.6% (26/61) of mutants exhibited multiple mutant phenotypes and 57.4% (35/61) exhibited a single mutant phenotype (Figure 2A). The Zn₂Cys₆ TFs regulating asexual reproduction also often controlled vegetative growth (conidial germination, colony growth, pigmentation and mycelial appearance) of the fungus (Figure 2C). Interestingly, 7 TF genes required for pathogenicity had multiple mutant phenotypes (Figure 2C, Table 1), and this phenomenon implied that the fungal pathogenic process is complex, where it is closely linked with many fungal development stages, and that the pathogenicity-related Zn₂Cys₆ TF genes are also involved in other developmental processes.

We identified the homologs of the 104 *M. oryzae* Zn_2Cys_6 TF genes in *N. crassa* and *F. graminearum* by Blastp (www. broadinstitute.org), and then compared the phenotypes of the knockout mutants with the homologs studied in *N. crassa* [29] or *F. graminearum* [27] (Table S5 in Text S1). Deletion of members of the Zn_2Cys_6 TF family in *F. graminearum* and *N. crassa* resulted in mutant phenotypes at 16% (46/296) and 42% (30/72) of the mutants [27,29]. However, more mutants (58.7%) in *M. oryzae* displayed visible mutant phenotypes. But only two mutants showed defects in growth and asexual and/or sexual development in three species simultaneously (Table S5 in Text S1). This comparison of mutant phenotypes between the three fungi suggested that the large majority of Zn_2Cys_6 TF genes evolved to have a unique, not conserved function in regulating fungal development.

Zn₂Cys₆ transcription factors involved in fungal growth

The null mutants of 27 Zn_2Cys_6 TFs showed significant differences in fungal growth with the wild-type strain (Figure 2B, Table 1). Among these null mutants, three mutants (01C9-1/

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Gene locus (MG8)	Gene	Growth	Conidiation	Conidial germination	Appressorium formation	Virulence	Stress
MGG_04387	FZC1	Normal	Normal	Normal	Normal	Normal	Affected
MGG_01486	FZC2	Normal	Normal	Normal	Reduced	Normal	Affected
MGG_01518	MoNIT4	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_01624	FZC3	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_01734	FZC4	Normal	Normal	Normal	Normal	Normal	Affected
MGG_01833	FZC5	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_02377	FZC6	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_02595	FZC7	Normal	Normal	Normal	Reduced	Normal	Unaffected
MGG_02879	FZC8	Reduced	Normal	Normal	Normal	Normal	Unaffected
MGG_02880	FZC9	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_03463	FZC10	Normal	Normal	Normal	Normal	Normal	Affected
MGG_04108	TAS1	Normal	Normal	Normal	Normal	Normal	Affected
MGG_04843	FZC11	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_05829	FZC12	Normal	Normal	Normal	Normal	Normal	Affected
MGG_05891	FZC13	Reduced	Reduced	Normal	Normal	Normal	Affected
MGG_06355	FZC14	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_07063	ecc1	Reduced	None			Normal	Affected
MGG_07800	FZC15	Normal	Normal	Normal	Normal	Normal	Affected
MGG_08094	FZC16	Normal	Normal	Reduced	Normal	Normal	Affected
MGG_08130	FZC17	Normal	Normal	Normal	Normal	Normal	Affected
MGG_08777	FZC18	Normal	Normal	Normal	Normal	Normal	Affected
MGG_10197	TRA 1	Normal	Normal	Reduced	Normal	Normal	Unaffected
MGG_11116	FZC19	Normal	Normal	Normal	Normal	Normal	Affected
MGG_12037	FZC20	Normal	Normal	Normal	Normal	Normal	Affected
MGG_12424	FZC21	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_16444	FZC22	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_16756	FZC23	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_17060	FZC24	Normal	Normal	Normal	Normal	Normal	Affected
MGG_17669	FZC25	Normal	Reduced	Normal	Normal	Normal	Unaffected
MGG_17821	FZC26	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_17841	GPF1	Reduced	Normal	Reduced	Reduced	None	Affected
MGG_08058	MoACEII	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_00329	FZC27	Normal	Normal	Normal	Reduced	Normal	Affected
MGG_07681	FZC28	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_01779	FZC29	Normal	Normal	Normal	Normal	Normal	Affected

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Table 1. Cont.							
Gene locus (MG8)	Gene	Growth	Conidiation	Conidial germination	Appressorium formation	Virulence	Stress
MGG_01887	FZC30	Normal	Normal	Normal	Normal	Normal	Affected
MGG_02866	FZC31	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_04674	FZC32	Normal	Normal	Normal	Normal	Normal	Affected
MGG_05033	FZC33	Reduced	Normal	Reduced	Normal	Normal	Affected
MGG_05343	MoCOD1	Reduced	None			Reduced	Affected
MGG_06279	FZC34	Normal	Normal	Normal	Normal	Normal	Affected
MGG_06312	FZC35	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_06416	FZC36	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_06626	FZC37	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_06778	FZC38	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_06832	FZC39	Normal	Normal	Normal	Normal	Normal	Affected
MGG_06954	FZC40	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_07149	GTA 1	Reduced	Reduced	Reduced	Normal	Reduced	Affected
MGG_07215	PIG1	Normal	Normal	Normal	Normal	Normal	Affected
MGG_07534	FZC41	Normal	Normal	Normal	Normal	Normal	Affected
MGG_07549	FZC42	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_07777	FZC43	Normal	Reduced	Reduced	Normal	Normal	Affected
MGG_07845	FZC44	Increased	Normal	Normal	Normal	Normal	Unaffected
MGG_08185	FZC45	Increased	Reduced	Normal	Normal	Normal	Affected
MGG_08314	FZC46	Reduced	Normal	Normal	Normal	Normal	Unaffected
MGG_08784	FZC47	Normal	Normal	Normal	Normal	Normal	Affected
MGG_14728	FZC48	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_09027	FZC49	Normal	Normal	Normal	Normal	Normal	Affected
MGG_09273	FZC50	Reduced	Normal	Reduced	Normal	Normal	Affected
MGG_09312	FZC51	Normal	Normal	Normal	Normal	Normal	Affected
MGG_09676	FZC52	Normal	Normal	Normal	Normal	Normal	Affected
MGG_09829	FZC53	Normal	Normal	Normal	Normal	Normal	Affected
MGG_09950	FZC54	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_11764	FZC55	Normal	Normal	Normal	Reduced	Normal	Affected
MGG_12776	FZC56	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_13350	FZC57	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_13360	CNF3	Normal	Increased	Normal	Normal	Normal	Unaffected
MGG_13385	FZC58	Normal	Normal	Normal	Normal	Normal	Affected
MGG_13629	FZC59	Normal	Normal	Normal	Normal	Normal	Affected
MGG_14175	FZC60	Normal	Normal	Normal	Normal	Normal	Affected

Table 1. Cont.							
Gene locus (MG8)	Gene	Growth	Conidiation	Conidial germination	Appressorium formation	Virulence	Stress
MGG_14852	FZC61	Normal	Reduced	Normal	Normal	Normal	Affected
MGG_15023	CNF2	Normal	Increased	Normal	Normal	Reduced	Affected
MGG_17623	PCF1	Normal	Reduced	Reduced	Normal	Reduced	Affected
MGG_00021	FZC62	Increased	Reduced	Normal	Normal	Normal	Affected
MGG_00096	FZC63	Increased	Normal	Normal	Normal	Normal	Affected
MGG_00320	FZC64	Normal	Normal	Normal	Normal	Normal	Affected
MGG_00417	FZC65	Normal	Normal	Normal	Normal	Normal	Affected
MGG_00494	MoPRO1	Normal	Reduced	Normal	Reduced	Normal	Unaffected
MGG_02089	FZC66	Normal	Normal	Normal	Normal	Normal	Affected
MGG_02289	FZC67	Increased	Normal	Normal	Normal	Normal	Affected
MGG_02962	CNF1	Normal	Increased	Reduced	Normal	Reduced	Affected
MGG_03183	CNF4	Normal	Increased	Normal	Normal	Normal	Affected
MGG_03669	FZC68	Normal	Normal	Normal	Normal	Normal	Affected
MGG_04141	FZC69	Reduced	Normal	Normal	Reduced	Normal	Affected
MGG_04326	FZC70	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_04360	FZC71	Normal	Normal	Normal	Normal	Normal	Affected
MGG_04933	FZC72	Normal	Normal	Normal	Normal	Normal	Affected
MGG_05153	FZC73	Reduced	Normal	Normal	Normal	Normal	Affected
MGG_05659	CCA1	Reduced	Reduced	Reduced	Reduced	Reduced	Affected
MGG_05724	FZC74	Normal	Normal	Normal	Normal	Normal	Affected
MGG_05845	FZC75	Normal	Normal	Normal	Reduced	Normal	Affected
MGG_06243	FZC76	Normal	Normal	Normal	Normal	Normal	Affected
MGG_06455	FZC77	Normal	Normal	Normal	Normal	Normal	Affected
MGG_07458	FZC78	Normal	Normal	Normal	Normal	Normal	Unaffected
MGG_07636	FZC79	Increased	Normal	Normal	Normal	Normal	Affected
MGG_08361	FZC80	Normal	Normal	Normal	Normal	Normal	Affected
MGG_08974	FZC81	Normal	Normal	Normal	Normal	Normal	Affected
MGG_12339	FZC82	Normal	Normal	Reduced	Reduced	Normal	Affected
MGG_12349	CONx1	Normal	Reduced			Reduced	Affected
MGG_13927	FZC83	Increased	Normal	Normal	Normal	Normal	Affected
MGG_14816	FZC84	Normal	Normal	Reduced	Normal	Normal	Affected
MGG_15139	FZC85	Normal	Normal	Normal	Normal	Normal	Affected
MGG_17012	FZC86	Normal	Normal	Normal	Normal	Normal	Affected
MGG_03711	FZC87	Reduced	Normal	Normal	Normal	Normal	Affected
Notes: The phenotypes of t <i>FZC1~FZC87</i> (Fungal-specifi doi:10.13271/journal prost 100	he mutants in colony grow c Zn ₂ Cys ₆ transcription fac 4437 ±001	wth, conidiation, conidial ctor 1 to 87) if not desig	l germination, appressorium fr nated in the succeeding text.	ormation, virulence and respor The detailed phenotypic infor	rse to stress were compared with mation of mutants was shown in	the wild-type strain 70- Table S4 in Text S1.	15. The genes are named as

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Figure 3. Colony growth and conidiophores of *M. oryzae* **strains.** (**A**) Colony growth of the wild-type strain 70-15, mutants ($\Delta gcc1$, $\Delta gpf1$ and $\Delta gta1$), and complemented strains (gcc1-c, gta1-c, gta1-c) on CM medium for 8 days. Bar = 1 cm. (**B**) Conidiophores of the wild-type, mutants ($\Delta cnf1$, $\Delta cnf2$, $\Delta pcf1$, $\Delta cca1$, $\Delta conx1$, $\Delta gcc1$ and $\Delta Mocod1$), and complemented strains (cnf1-c, cnf2-c, pcf1-c, cca1-c, conx1-c, gcc1-c and Mocod1-c). Bar = 20 µm.

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 $\Delta MGG 07063$, 01F1-1/ΔMGG 17841 02G4-1/ and $\Delta MGG \ 07149$) showed reduced colony growth on CM medium at 24.7, 26.6 and 43.5%, respectively (Table S4 in Text S1). The TF genes deleted in these mutants were designated GCC1 (growth, conidiation and cell wall regulatory factor 1, MGG_07063), GPF1 (growth and pathogenicity regulatory factor 1, MGG_17841) and GTA1 (growth and tolerance to acidic stress regulatory factor 1, MGG 07149), respectively. To confirm the defects of the mutants in growth were caused by the knockout of the TF genes, we complemented the three mutants $\Delta gcc1$, $\Delta gpf1$ and $\Delta gta1$ with their respective native copies from wild-type strain 70-15 (Figure S4C). The phenotypic analyses showed the GCC1-rescued strain (gcc1-c) and GTA1-rescued strain (gta1-c) all recovered from their defects in growth, and GPF1-rescued strain (gpf1-c) also recovered most defects in growth when compared to the mutants and the wild-type strain (Table 2, Figure 3A). These results implied that Zn₂Cys₆ TFs GCC1, GPF1 and GTA1 were required for the regulation of fungal growth in the rice blast fungus.

Zn₂Cys₆ transcription factors involved in fungal asexual development

Among the 104 Zn_2Cys_6 TFs, the null mutants of 25 genes displayed distinct changes in asexual development (Figure 2B, Table 1). Of these mutants, 21 mutants produced less conidia or even no conidia, but the other 4 mutants produced more conidia (Table 1, Table S4 in Text S1). We selected 2 mutants without conidiation, 4 mutants with less conidiation and 2 mutants with more conidiation to be complemented by their native genes to show the relationship between mutant's phenotype and TF gene deleted (Figure S4C). The genes deleted in these null mutants were CNF1 (conidial production negative regulatory factor 1, MGG_02962 in mutant 06B5-7), CNF2 (conidial production negative regulatory factor 2, MGG_15023 in mutant 03C7-2), PCF1 (pathogenicity and conidiation regulatory factor 1, MGG_17623 in mutant 03C11-4), MoNIT4 (a homolog of the nit-4 gene, which is a pathway-specific regulatory gene that mediates nitrate induction in N. crassa [38], MGG_01518 in mutant 01A7-1), CCA1 (conidiation, conidial germination and appressorium formation required transcription factor 1, MGG_05659 in mutant 06C2-14), CONx1 (conidiation required transcription factor x1, MGG_12349 in mutant 06D5-21), GCC1 (MGG_07063 in mutant 01C9-1) and Mocod1 (homolog of amyR from Aspergillus nidulans [3,39,40], MGG_05343 in mutant 02F2-6). The mutants $\Delta cnf1$ and $\Delta cnf2$ produced more conidia than did the wild-type; mutants $\Delta pcf1$, $\Delta cca1$ and $\Delta conx1$ produced less conidia; and mutants $\Delta gcc1$ and $\Delta Mocod1$ did not produce any conidia. The complementation experiments displayed that the defects of mutants in conidiation were caused by the deletion of the TF genes, since the complemented strains of these mutants all recovered from their abnormal asexual developmental phenomena (Table 2, Figure 3B).

CNF1 is a strong conidial production negative regulatory factor in rice blast fungus, and its deletion may result in greatly increased conidial production (usually 20- to 40-fold of the wild-type strain). Moreover, the mutant $\Delta cnf1$ had significant changes in mycelial appearance, such as deep dark colony color from the gray-white of the wild-type strain, short spore-bearing aerial hyphae form long aerial hyphae of the wild-type strain, and spore-bearing aerial hyphae formed at the rim of mycelial colony earlier than in the wild-type strain (Figure 4A). Although the conidiophore development of $\Delta cnf1$ seemed similar to that of the wild-type strain and CNF1-rescued mutant cnf1-c (Figure 3B, Figure 4C), more aerial hyphae differentiated into conidiaophores in $\Delta cnf1$ (Figure 4B). The increase in conidial production of $\Delta cnf1$ was possibly partly due to the spore-bearing hyphae differentiating earlier and more from the aerial hyphae than in the wild-type strain.

Strain	Growth (mm)	Conidiation (×10 ³ /cm ²)	Conidial germination (%)	Appressorium formation (%)	Strain	Growth (mm)	Conidiation (×10 ³ /cm ²)	Conidial germination (%)	Appressorium formation (%)
Wild-type	48.7±0.6A*	62.5±10.0C	95.2±1.8A	95.9±1.6A					
$\Delta gcc1$	38.7±0.6C	0.0±0.0E			gcc1-c	48.7±0.6A	61.0±8.3C		
∆gpf1	38.0±0.1C		76.4±6.4C	86.5±1.1C	gpf1-c	47.0±1.0B		94.0±1.5A	97.3±0.7A
$\Delta g ta l$	27.3±0.6D		$84.5 \pm 0.5B$	97.8±0.3A	gta1-c	49.7±0.6A		$92.5\pm2.6A$	97.8±1.6A
∆cnf1		1229.7±101.1A	86.7±2.7B	89.9±3.5B	cnf1-c		66.4±8.8C	91.8±2.4A	96.2±0.3A
Δcnf2		188.9±7.6B			cnf2-c		71.0±5.2C		
ΔMonit4		20.2±4.6D			Monit4-c		59.3±15.6C		
$\Delta pcf1$		28.5±6.1D	86.4±2.4B	87.4±2.5B	pcf1-c		58.6±8.0C	94.4±0.9A	97.1±2.1A
Асса 1		7.3±1.7E	3.6±0.6E	10.2±0.8E	cca1-c		65.9±6.4C	94.4±1.7A	96.8±0.7A
Δconx1		0.1±0.0E			conx1-c		65.7±4.4C		
∆Mocod1		0.0±0.0E			Mocod1-c		60.8±10.3C		
Δfzc16			$84.6 \pm 3.0B$	97.1±1.2A	fzc16-c			95.1±1.8A	98.5±0.8A
Notes: The strai suspensions we	ns (5-mm mycelial bloc re incubated on plasti	cks) were grown on CM mei ic slides for 4 hpi (conidial c	dium for 8 days, and the dermination 2	diameter of colonies wied to the diameter of colonies wied to the termination of the diameter	as then measured (gr rmation assav).	owth) or conidia wer	e collected and count	ted (conidiation). 40-µl (1×10 ⁵ conidia/ml) conidia

*Same capital letters incluated on plastic slides for 4 hpi (conidial germination assay) and 24 hpi (appressorium formation assay). *Same capital letters indicate non-significant difference estimated by Duncan's test in each developmental item (P≤0.05). The experimental strains were the wild-type strain, the mutants ($\Delta cca1$, $\Delta cont1$, $\Delta cnf2$, $\Delta fc16$, $\Delta gcc1$, $\Delta gpf1$, $\Delta fa1$, $\Delta Mocd1$, $\Delta Monit4$ and $\Delta pcf1$) and complemented strains (cca1-c, cnt1-c, cnf2-c, fc16-c, gcc1-c, gcc1-c, gcc1-c, gcc1-c, fc10-c, fc16-c, gcc1-c, fc10-c, fc2-c, fc2-c, fc2-c, fc2-c, gcc1-c, gcc1-c, dc1-c, dc2-c, dc1-c, dc1-c, cnt1-c, cnt2-c, fc2-c, gcc1-c, dc1-c, dc2-c, dc1-c, dc1-c, dc1-c, dc2-c, dc1-c, dc1



Figure 4. Mycelial appearance and spore-bearing aerial hyphae of the *M. oryzae* **strains.** (**A**) The mycelial appearance of the wild-type strain, $\Delta cnf1$ and its complemented strain cnf1-c. Stars indicate nonsporulating hyphae. Bar = 1 cm. (**B**) Spore-bearing aerial hyphae of the wild-type strain, $\Delta cnf1$ and its complemented strain cnf1-c on CM medium. Bar = 20 µm. (**C**) Conidial development on conidiophores of the *M. oryzae* strains $\Delta cnf1$ and $\Delta cca1$. The conidiophore pictures of the wild-type, mutants ($\Delta cnf1$ and $\Delta cca1$), and their complemented strains (cnf1-c and cca1-c) were taken on the strains grown on CM medium on microscope slides for 1 day. Bar = 40 µm. doi:10.1371/journal.ppat.1004432.q004

 $\Delta cca1$ produced few and odd, long and vacuolated conidia (Table 2, Figure 5). Single or two tandem spores differentiated from the apex of conidiophores in the mutant $\Delta cca1$, and the CCA1-rescued mutant cca1-c produced normal conidiophores and conidia similar as the wild-type strain (Figure 4C, Figure 5). These results suggested that CCA1 is required for the differentiation of conidiophores and the formation of conidia.

Zn₂Cys₆ transcription factors involved in conidial germination and appressorium formation

When the null mutants of 101 Zn₂Cys₆ TFs were assayed for their conidial germination rate at 4 hpi (hour post inoculation) and their appressorium formation rate at 24 hpi, 12 mutants and 10 mutants displayed significant differences in conidial germination and appressorium formation, respectively (Figure 2B, Table 1). We assayed conidial germination and appressorium formation in the complemented strains of 6 null mutants ($\Delta fzc16$, $\Delta gpf1$, $\Delta gta1$, $\Delta pcf1$, $\Delta cnf1$ and $\Delta cca1$) (Figure S4C) and found that these complemented strains all recovered from their defects in conidial germination or in appressorium formation (Table 2). These experiments implied that Zn₂Cys₆ TFs *GPF1*, *PCF1*, *CNF1*, and *CCA1* are required for conidial germination and appressorium formation and that *FZC16* and *GTA1* are required for conidial germination in the rice blast fungus.

Among these 6 mutants, the mutant $\Delta cca1$ showed marked defects not only in conidial germination and appressorium formation, but also in conidiation (Table 2). The conidial



Figure 5. Conidia, conidial germination and appressorium formation of the *M. oryzae* strain $\Delta cca1$. The conidia (0 h) of the wild-type strain, mutant $\Delta cca1$, and its complemented strain cca1-c were grown in H₂O on plastic cover slides for 4 h (conidial germination) and 24 h (appressorium formation). Bar = 10 µm. doi:10.1371/journal.ppat.1004432.g005

germination rate of the mutant $\Delta cca1$ was 3.6%, and appressorium formation rate was 10.2% (Table 2). The structure and shape of conidium, germinated conidium and appressorium of the mutant $\Delta cca1$ displayed notable differences with the wild-type strain (Figure 5). Therefore, the mutant $\Delta cca1$ produced few multifarious and vacuolated conidia, which germinated and formed not fully melaninized appressoria at a low ratio.

Zn₂Cys₆ transcription factors required for virulence

The pathogenicity of 104 null mutants was tested first on intact barley leaves by placing agar plugs containing mycelia on them. Mutant $\Delta gpf1$ (MGG_17841) was found to lose pathogenicity to barley, and 4 mutants $\Delta cnf1$ (MGG_02962), $\Delta cnf2$ (MGG_15023), $\Delta Mocod1$ (MGG_05343) and $\Delta pcf1$ (MGG_17623) showed reduced pathogenicity (Table S4 in Text S1, Figure S2). We then determined the pathogenicity of 101 mutants on rice seedlings (CO-39) by spraying with conidial suspension $(1 \times 10^5$ spores/ml). The results showed that it was decreased in 3 mutants $\Delta cnf1$ (MGG_02962), $\Delta cnf2$ (MGG_15023) and $\Delta gta1$ (MGG_07149) and that 2 mutants $\Delta g p f 1$ (MGG_17841) and $\Delta cca1$ (MGG_05659) lost their virulence to rice (Table S4 in Text S1, Figure S3A). Impressively, $\Delta cca1$ caused lesions on barley leaves when inoculated with mycelial plugs, but lost virulence on rice seedlings when spayed as a conidial suspension. The other 3 mutants $\Delta conx1$ (MGG_12349), $\Delta gcc1$ (MGG_07063) and $\Delta Mocod1$ (MGG_05343) with little or no conidiation were tested on intact rice leaves with agar plugs containing mycelia. $\Delta conx1$ and $\Delta Mocod1$ were found to have lower virulence (Table S4 in Text S1, Figure S3B).

After complemented with native genes (Figure S4C), we retested the pathogenicity of the 7 mutants with their complemented strains on rice seedlings by spraying with conidial suspensions ($\Delta gpf1$, $\Delta cnf1$, $\Delta cnf2$, $\Delta gta1$, $\Delta cca1$ and gpf1-c, cnf1-c, cnf2-c, gta1-c, and cca1-c) (Figure 6A) or on rice leaf explants by inoculating with mycelial plugs ($\Delta Mocod1$, $\Delta conx1$ and Mocod1-c,



Figure 6. Pathogenicity assay of the *M. oryzae* **strains.** (**A**) Pathogenicity assay of 5 mutants and their complementation strains on rice. These 5 mutants are $\Delta gpf1$, $\Delta gta1$, $\Delta cnf2$, $\Delta cnf1$, $\Delta cca1$ and their complementation strains gpf1-c, gta1-c, cnf2-c, cnf1-c, and cca1-c. The rice seedlings were sprayed with a conidial suspension of *M. oryzae* strains and cultured for 7 days. (**B**) Pathogenicity assay of the mutants on rice leaf explants. The mycelial agar plugs of the mutants $\Delta Mocod1$ and $\Delta conx1$, their complementation strains Mocod1-c and conx1-c, and the wild-type strain 70-15 were placed on intact rice leaves for 4 days. (**C**) Penetration assay. 10-or 20- μ l (5×10^4 conidia/ml) conidial suspensions were inoculated on onion cuticle or barley leaves (intact or abraded) and incubated for 24 or 48 h. The experimental strains were the wild-type strain 70-15, mutants ($\Delta gpf1$, $\Delta gta1$, $\Delta cnf2$, $\Delta cnf1$ and $\Delta cca1$) and complemented strains (gpf1-c, gta1-c, cnf2-c, cnf1-c and cca1-c). Same capital letters in same treatment item indicate non-significant difference estimated by Duncan's test ($P \leq 0.05$). (**D**) Penetration of $\Delta gpf1$. 20 μ l (5×10^4 conidia/ml) conidial suspensions of $\Delta gpf1$, complemented strain gpf1-c or wild-type strain were inoculated on barley leaf explants (intact or abraded) and incubated for 24 or 48 h. Arrows indicate the *in planta* hyphae invaded. Bar = 25 μ m.

conx1-c) (Figure 6B). The facts that the complemented strains recovered from their defects in virulence on rice implied that the loss or weakening of virulence in the mutants was caused by the deletion of the TF genes *GPF1*, *CNF1*, *CNF2*, *GTA1*, *CCA1*, *CONx1* and *MoCOD1*, respectively. To assay the roles of these TF genes in plant penetration, onion cuticle and barley leaves were

inoculated with a conidial suspension of the mutants and their complemented strains. Compared with the wild-type strain, the ability of $\Delta gpf1$, $\Delta gta1$, $\Delta cnf2$, $\Delta cnf1$ and $\Delta cca1$ to penetrate the onion cuticle and barley leaf cuticle was weakened to some extent (Figure 6C). Notably, the mutants $\Delta gpf1$ and $\Delta cca1$ were unable to penetrate barley cuticle, even after barley cuticle was abraded



Figure 7. Growth of the *M. oryzae* **strains under stress conditions.** (A) Growth inhibition rate. 5-mm mycelial blocks of *M. oryzae* strains were inoculated on CM medium and medium with stress conditions for 8 days, and the diameter of colonies was then measured to calculate the growth inhibition rate. The experimental strains were the wild-type strain, mutants ($\Delta Monit4$, $\Delta gpf1$, $\Delta gcc1$, $\Delta tas1$ and $\Delta gta1$), and complemented strains (*Monit4-c, gpf1-c, gcc1-c, tas1-c* and *gta1-c*). Same capital letters in same stress item indicate non-significant difference estimated by Duncan's test ($P \leq 0.05$). (B) Mycelial growth of $\Delta Monit4$ on MM medium. The wild-type strain, $\Delta Monit4$, and complemented strain *Monit4-c* were grown on MM medium at 25°C for 8 days. Bar = 5 mm. doi:10.1371/journal.ppat.1004432.g007



Figure 8. Comparison of differentially expressed genes between $\Delta gpf1$ and $\Delta cnf2$. Green or red arrows are the directions of gene expression regulation in $\Delta gpf1$ or $\Delta cnf2$. doi:10.1371/journal.ppat.1004432.g008

(for $\Delta gpfI$), while the wild-type strain penetrated into the epidermal cells of barley leaves and grew invasively after 24 hpi (Figure 6C, Figure 6D). These results suggested that *GPFI* is required by the rice blast fungus for penetration into the plant cuticle and possibly for invasive growth. Although the conidia of $\Delta cca1$ were unable to penetrate barley cuticle and could not cause lesions in rice, the mycelia of $\Delta cca1$ could cause lesions in barley and rice. The difference in virulence between conidia and mycelia may lie in the conidial defects of $\Delta cca1$ in germination, appressorium formation and host penetration ability.

Phenotypic analyses of Zn₂Cys₆ transcription factor deletion mutants under stress conditions

There are various stress conditions encountered by M. oryzae when it completes its infection cycle in rice. It is possible to predict the biological functions of these deleted Zn₂Cvs₆ TF genes in cell activity regulation by assaying the resistance ability to stress conditions of the mutants. We screened and assaved in a rapid way the mycelial growth of the mutants under 9 abiotic stress conditions (0.5 mM H₂O₂, 0.005% SDS, 0.3 M CaCl₂, 0.5 M NaCl, 1 M sorbitol, olive oil as a single organic carbon source, pH 5.0 and pH 9.0, and minimal medium). Four mycelial inoculation blocks were placed in a 9-cm plate by needle point inoculation and the diameter of mycelial colonies were measured after 6 days culture. The results displayed that many mutants had significant visible phenotypes under those stress conditions when compared with the wild-type strain (Table S4 in Text S1): 14 mutants showed mutant phenotypes in MM medium (nitrogen metabolism), 16 mutants in CM-C medium containing olive oil (carbon metabolism), 22 mutants in CM medium with SDS (cell wall stress), 17 mutants or 19 mutants in CM medium with high concentration of Ca²⁺ or Na⁺ (ionic regulation), 18 mutants in CM medium with 1 M sorbitol (hypertonic pressure), 30 mutants in CM medium with 0.5 mM H₂O₂ (oxidative stress), and 30 mutants or 14 mutants in CM medium at pH 5.0 or pH 9.0 (ambient pH regulation).

After comparison of the relative growth rate of mutants on CM medium, we selected 5 mutants (01A7-1, 01B11-1, 01C9-1, 01F1-1 and 02G4-1) that had significant changes in mycelial growth under stress conditions to be complemented with native genes to see if these phenotypes were caused by the deletion of TF genes

(Figure S4). The TF genes in these 5 mutants were: MoNIT4 in mutant 01A7-1 (MGG_01518), GPF1 in mutant 01F1-1 (MGG_17841), GCC1 in mutant 01C9-1 (MGG_07063), TAS1 (tolerance to acidic stress regulatory factor 1) in mutant 01B11-1 (MGG_04108) and GTA1 in mutant 02G4-1 (MGG_07149). The mycelial growth of $\Delta Monit4$ in minimal medium, $\Delta gpf1$ in CM-C medium containing olive oil, and $\Delta tas1$ at pH 5.0 was reduced (Figure 7A). However, the mycelial growth of $\Delta gcc1$ in CM medium containing SDS and $\Delta gta1$ at pH 5.0 was increased. When rescued with native genes, these mutants all recovered their growth under stress conditions (Figure 7A). Interestingly, the mutant $\Delta Monit4$ grew normally as the wild-type strain on CM medium (Table S4 in Text S1, Figure S1), but grew slowly as a very sparse and thin mycelium layer on MM medium (Figure 7B). As previously reported, MoNIT4 is necessary for the regulation of nitrogen metabolism [38,41]. These data also suggested that TF gene GPF1 is necessary for gene expression regulation relative to fat metabolism, that GCC1 is necessary for gene expression regulation relative to cell wall synthesis, and that TAS1 and GTA1 are necessary for gene expression regulation in response to acidic stress.

Genes regulated by two virulence-required genes GPF1 and CNF2

Eight Zn₂Cys₆ TF genes (CCA1, CNF1, CNF2, CONx1, GPF1, GTA1, MoCOD1 and PCF1) are required for fungal pathogenicity to rice or barley. $\Delta g p f 1$ and $\Delta cn f 2$ are two mutants that have similar phenotypes on virulence with relatively limited differences in other fields, because $\Delta Mocod1$ and $\Delta conx1$ do not produce conidia, $\Delta cca1$ and $\Delta gta1$ have virulence when inoculated on rice leaves with mycelial plugs, $\Delta pcf1$ shows reduced virulence on barley inoculated with mycelial plugs, but normal when sprayed on rice, and $\Delta cnf1$ produces greatly increased conidia. To understand how these TF genes affect fungal pathogenicity, the two genes GPF1 and CNF2, whose mutants lose $(\Delta g p f I)$ or have reduced $(\Delta cn f 2)$ virulence to both barley and rice, were selected to analyze their functions in gene expression regulation. Since $\Delta g p f 1$ could not penetrate rice cuticle, we could not obtain its mRNA during infection. The rice blast fungus experiences temporal nitrogen starvation when it infects the plant, and the gene expression pattern of the mycelia under starvation is similar to that at plant infection [42–45], and the hyphae enduring starvation are similar to those inoculated as mycelial plugs in plant infection. We therefore analyzed the genome-wide gene expression of the mycelia after 4 h starvation of the wild-type strain, $\Delta g p f I$ and $\Delta cnf2$ by RNA-seq. The RNA-seq experiments were performed in biologic triplicate for each strain.

The transcripts of 10,864, 11,048 and 10,681 genes were identified in the starved mycelia of the wild-type strain, $\Delta g p f 1$ and $\Delta cnf2$, respectively. Compared with the wild-type strain, 2641 genes were differentially expressed significantly in $\Delta g p f 1$ (FDR < 0.05), with 1406 genes up-regulated and 1235 down-regulated (Table S6 in Text S1), and 3144 genes in $\Delta cnf2$ (FDR<0.05), with 1668 genes up-regulated and 1476 down-regulated (Table S7 in Text S1). The similarities and differences between genes regulated by GPF1 and CNF2 were analyzed in detail by comparing the DEGs in $\Delta g p f 1$ or $\Delta cn f 2$. The comparison assays showed that the expression of 611 genes or 1114 genes were independently regulated by GPF1 or CNF2, while 2030 genes were regulated by both GPF1 and CNF2 together (Figure 8). Among the 2030 genes commonly regulated by GPF1 and CNF2, 916 genes were downregulated and 1105 genes up-regulated simultaneously in both $\Delta gpf1$ and $\Delta cnf2$, while only 9 genes were regulated in the opposite direction in both mutants (Figure 8, Table S8 in Text S1).

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Table 3. GPF1- or CNF2-dependent genes encoding known virulence factors.

Gene locus (MG8)	Gene	$\Delta gpf1^{a}$	$\Delta cnf2^{a}$	Virulence on plants/Function	Reference
MGG_00365	MAGB	-	-1.58	Reduced virulence/G-protein signaling	[48]
MGG_00466	MoCDC42	-1.52	-2.00	Reduced virulence/CDC42	[74]
MGG_00501	TGD2	-1.67	-2.68	Reduced virulence/Transcription factor	[15,72,83]
MGG_00692	MSTU1	-7.80	-11.85	Reduced virulence/Transcription factor	[8]
MGG_00800	MST7	-2.00	-1.67	No virulence/MAPK pathway	[52]
MGG_00803	MoSNF1	-1.43	-1.79	Reduced virulence/Glucose derepression pathway	[62]
MGG_00883	MoMCK1	-1.39	-1.52	No virulence/MAPK pathway	[53]
MGG_01057	MoLDB1	-1.63	-2.18	Reduced virulence/Transcription factor	[18]
NGG_01204	MoMIG1	-	-1.95	No virulence/Transcription factor	[20]
MGG_01230	MoSSADH	-1.54	-	Reduced virulence/γ-aminobutyric acid degradation	[23]
MGG_01282		1.46	2.26	No virulence/polyubiquitin	[84]
MGG_01558	SET3	-	-1.53	No virulence/Set3 complex	[85]
MGG_01662	MoAAT	-	-1.56	Reduced virulence/4-aminobutyrate aminotransferase	[23]
NGG_01663	HOS2	-1.46	-1.51	No virulence/Set3 complex	[85]
NGG_02531	MoVPR	-3.04	-3.00	Reduced virulence	[23]
NGG_02731	MoRAC1	-	-1.45	Reduced virulence/Small GTPase	[86]
MGG_02848	MC69	-1.57	-	Reduced virulence/Secreted proteins	[87]
MGG_02897	Mossk1	-	-1.81	Reduced virulence/Response regulator	[79]
/IGG_03065	MoVMA11	-1.53	-	No virulence/ATPase	[46]
MGG_03694	MoATG6	-1.39	-	No virulence/Autophagy	[88]
MGG 04050	MoSEC22	-2.81	-3.80	No virulence/R-Snare	[89]
 /IGG 04895	ICL1	-1.72	-1.59	Reduced virulence/Glyoxylate cycle	[63]
 /IGG 04943	MoMPS1	-	-1.75	No virulence/MAPK pathway	[54]
 MGG 05133	MoCRZ1	-	-1.42	Reduced virulence/Ca ²⁺ signaling	[9,90]
MGG 05201	MGB1	-1.67	-1.80	No virulence/G-protein signaling	[51]
 //GG_05287	CON7		-1.93	No virulence/Transcription factor	[12]
/GG_05428	MoVAM7	-	-1.89	No virulence/Snare	[91]
MGG 05664	MoPDEH	-	-1.77	Reduced virulence/cAMP signaling	[92]
MGG 05755	MoMON1	1.38	-	No virulence/Vacuole and vesicle fusion	[93]
MGG 06033	MoMSB2	-	-1.78	Reduced virulence/Signaling	[94]
MGG 06249	MoTCTP	-	1 43	Reduced virulence/Mitochondrial protein	[80]
MGG 06393	MoATG1	-1.36	-1.51	No virulence/Autophagy	[55]
MGG 06439	MoTEA4	-1.88	-2.26	Reduced virulence	[95]
MGG 06564	MoPAC2	-	-1.67	Reduced virulence/Transcription factor	[96]
MGG 06868	II V2	4 74	4.03	Reduced virulence/Amino acid metabolic nathway	[61]
MGG 06971	MoSEL 1	-	-173	Reduced virulence/Transcription factor	[22]
MGG 07460	MoHYR1	1 73	2 3 2	Reduced virulence/Reactive ovvicen species	[59]
MGG 08212	MoATE1	1.75	-	Reduced virulence/Transcription factor	[24]
MGG 08850	MoGTI	-1.58	-2.55	Reduced virulence/Stress	[96]
MGG_08895	GTP1	2.06	1.82	Reduced virulence/Parctive eviden species	[60]
MGG 00125	MosHO1	2.00	- 1 00	Reduced virulence/Signaling	[04]
AGG 00262	Montes	764	- 1.0U	No virulence/Autonbagy	[54]
AGG 00550	MoATCO	. 1 41	0.00	No virulence/Autophagy	[50]
	DMK1	- 1.41	- 1.42	No virulence/Autophagy	[57]
	PIVIK I	- 1.49	- 1.93		[52]
viGG_09912	MOCMK1	-1.75	- 1.46	Reduced virulence/Ca signaling	[97]
NGG_10315	MPG1	-12.49	-15.83	Reduced virulence/Hydrophobin	[75]
/IGG_10323	MoRHO3	-1.46	-1.69	No virulence/Small GTPase	[98]
	MoCDTE1	-	-172	No virulence/Transcription factor	[19]

Table 3. Cont.

Gene locus (MG8)	Gene	∆ <i>gpf1</i> ª	$\Delta cnf2^a$	Virulence on plants/Function	Reference
MGG_12832	SNX41	-1.36	-	No virulence/Autophagy	[58]
MGG_12865	PTH12	2.97	2.09	No virulence/Transcription factor	[16,47]
MGG_12868	ECH1	-	1.47	Reduced virulence/Enoyl-CoA hydratase	[99]
MGG_12958	MST12	-	-1.71	No virulence/Transcription factor	[21]
MGG_13624	MoABC1	-	-1.59	No virulence/ABC transporter	[100]
MGG_14014	MoPLAA	2.12	2.09	Reduced virulence/Lipid metabolism	[64]
MGG_14517	MoFLBA	-	-1.69	Reduced virulence/G-protein signaling	[49,50,101]
MGG_14847	MST11	-	-1.33	No virulence/MAPK pathway	[52]

Note:

^a) The expression of genes increased or decreased in $\Delta qpf1$ and $\Delta cnf2$.

The number means the fold change, and the symbol "-" means no significant change (FDR<0.05) afterwards compared with the wild-type strain.

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It is very surprising that the expression of so many DEGs were regulated in the same direction in $\Delta g p f 1$ and $\Delta cn f 2$ (Figure 8). The correlation coefficient between DEGs of $\Delta gpf1$ and $\Delta cnf2$ was determined by the linear trend model, and the result showed $R^2 = 0.88$ with p < 0.0001. These similar patterns were not mainly caused by the starvation, as we also assayed the mutants of five TF genes containing non-Zn2Cys6 domains by RNA-seq with the same treatment at the same time, but the mutants displayed specific DEG patterns different from $\Delta g p f 1$ and $\Delta c n f 2$. Therefore, these similar patterns in gene expression regulation in two mutants were possibly caused by the fact that TFs Gpf1 and Cnf2 had the same DNA-binding domain (Zn₂Cys₆ domain), and $\Delta gpf1$ or $\Delta cnf2$ had similar phenotypes in pathogenicity. However, GPF1 and CNF2 still showed great differences in gene regulation since 1152 DEGs appeared in $\Delta cnf2$ when compared with $\Delta gpf1$ (Table S9 in Text S1).

To see how GPF1 and CNF2 affect the pathogenicity to plants, we reviewed the functions of the DEGs which were studied in previous reports. Fifty DEGs in $\Delta gpf1$ and 68 DEGs in $\Delta cnf2$ (total of 80 DEGs in both were studied by knockout. Of the 80 genes, the mutants of 57 genes (36 DEGs in $\Delta g p f 1$ and 49 DEGs in $\Delta cnf2$ displayed defects in virulence (Table 3) but the mutants of 23 genes (14 DEGs in $\Delta gpf1$ and 19 DEGs in $\Delta cnf2$) were dispensable for fungal pathogenicity (Table S10 in Text S1). Interestingly, of these reported pathogenicity-required genes, the number of down-regulated genes (45 DEGs) was much higher than that of up-regulated genes (12 DEGs) in the two mutants $\Delta gpf1$ and $\Delta cnf2$ (Table 3). These pathogenicity-required genes are mainly related to transcription (such as CON7, MoLDB1, MoMIG1, MoPAC2, MSTU1, MST12 and PTH12) [8,12,18,20,21,46,47], G-protein signaling (such as MAGB, MoRGS1 and MGB1) [48-51], MAPK pathway (such as MST7, MoMCK1 and MoMPS1) [52-54], autophagy (such as MoATG1, MoATG5, MoATG9 and SNX41) [55-58], regulation of reactive oxygen species (such as MoHYR1 and GTR1) [59,60], and amino acid, lipid and carbohydrate metabolism (such as ILV2, MoSNF1, ICL1 and MoPLAA) [61-64]. We also checked the relationships between Zn2Cys6 TFs and Gpf1 or Cnf2 in transcription regulation. Surprisingly, 60 or 66 Zn₂Cys₆ TF genes were listed as DEGS of $\Delta gpf1$ or $\Delta cnf2$ (total of 79 genes in two mutants) (Table S11 in Text S1). The Zn₂Cys₆ TF genes are also the important targeted genes regulated by two Zn₂Cys₆ TF genes GPF1 and CNF2. These facts implied that the wellbalanced and sophisticated expression regulation of TF genes and pathogenicity-required genes is necessary for the maintenance of pathogenicity in *M. oryzae*.

Discussion

Since the genomes of a large number of fungi have been sequenced and are being sequenced, there is a surge of interest in functional genomics research through the systematic mutagenesis of identified genes. The construction of a genome-wide gene deletion mutant set of fungi based on the homologous recombinational gene knockout procedure is a valuable resource for the analysis of fungal development, pathogenicity, and many aspects of cell biology and biochemistry, such as those done in the budding yeast S. cerevisiae [65], the fission yeast Schizosaccharomyces pombe [66], and the saprobe filamentous fungus N. crassa [29]. However, except for N. crassa, F. graminearum and Aspergilli which are highly efficient in gene deletion experiments, no efficient system has been available to perform high-throughput gene knockout in filamentous fungi. In this study, we present an approach to knockout large numbers of genes that utilizes several methods suitable to high-throughput manipulation. Several timeconsuming steps of the gene knockout procedure could be performed in a high-throughput way. In this procedure, the gene-deletion cassettes were built using a yeast homologous recombination method [29] with a yeast-Escherichia-Agrobacterium shuttle vector pKO1B, which is the first reported artificial plasmid that could be replicated in yeast, E. coli and Agrobacterium cells. The gene-deletion cassettes obtained in a binary vector pKO1B could be directly used to transform fungal cells through the ATMT method [67]. This advantage avoids the tedious and inefficient work to transfer the gene-deletion cassettes from a yeast plasmid to another binary Agrobacterium plasmid. The use of GFP fluorescence as a negative marker to eliminate most ectopic insertion transformants reduced the number of transformants one has to identify by PCR or Southern blot. The employment of the negative (for the targeted gene)/positive (for the unique recombinational DNA) identification PCR and qPCR to identify null mutants instead of Southern blot, which is not high throughput, makes the identification of null mutants a high-throughput procedure. The copies of the gene-deletion cassette in mutant genomic DNA quantified by qPCR has been widely performed in animals and plants [68,69]. The reliability and stability of PCR or qPCR were guaranteed by the extraction of high quality genomic DNA with CTAB [70] performed in a high-throughput way in this

study. The result of the gene knockout events (Figure S4A) were also reconfirmed by Southern blot at the DNA level in 10 mutants (Figure S4B) and at the transcript level in 16 mutants (Figure S4C). As a result, two researchers were able to complete one cycle of the gene knockout experiment for 96 genes in a month, including preparing knockout vectors, performing ATMT transformation, and identifying null mutants, which is a major improvement over the individual gene knockout protocol. More importantly, this procedure can be adapted for knocking out genes in other fungi without any modification or after the substitution of organismspecific promoter to drive the GFP reporter. Besides the GFP gene, a second drug resistance gene (HPH/hygromycin B phosphotransferase gene, NEO/neomycin phosphotransferase II gene or BAR/glufosinate resistance gene, etc.) or a herpes simplex virus thymidine kinase (HSVtk) gene, which converts 5-fluoro-2'deoxyuridine to a toxic compound [71], could be used as an alternative negative selection marker against ectopic transformants. As an alternative vector, pKO1B-HPH (shown in Materials and Methods) is another yeast-Escherichia-Agrobacterium shuttle which could be used in this high-throughput gene knockout system besides pKO1B. To increase knockout efficiency, M. oryzae KU80 null mutants, which are defective for non-homologous end joining (NHEJ) DNA repair [31], could be alternative strains used in targeted gene replacement instead of the wild-type strain.

In this study, we generated 104 fungal-specific Zn₂Cys₆ TF gene-deleted mutants in M. oryzae by a high-throughput gene knockout procedure and analyzed the phenotypes of individual TF mutants. The deletion of Zn_2Cys_6 TF genes resulted in phenotype changes in fungal development and pathogenicity in 58.7% mutants compared to the wild-type strain, while 26 mutants were defective in pleiotropic phenotype. Colony growth and asexual reproduction (conidiation) were the two phenotypic categories most observed in mutants, and most mutants defective in conidiation were often defective in vegetative growth (conidial germination, colony growth, pigmentation and mycelial appearance). Seven Zn_2Cys_6 TF genes functional for pathogenicity in M. oryzae are also especially required for vegetative growth, conidiation or appressorium formation. Comparison of phenotypes between M. oryzae, F. graminearum and N. crassa Zn₂Cys₆ TF gene orthologs seemed to agree with the previously reported viewpoint that Zn₂Cys₆ TFs evolved divergently in how to regulate fungal growth and asexual development, rather than keeping the same function in different fungi [27]. These divergences in functions may be due to the differences in the life cycles between the three fungi. N. crassa is an obligate saprophyte which lives on dead organic material and cannot attack a living host, while M. oryzae and F. graminearum are facultative saprophytic plant pathogens that additionally need specialized structures (such as an appressorium) and functions to infect and obtain nutrients from living plants.

Until now, 6 Zn₂Cys₆ TF genes (*MoCOD1*, *MoCOD2*, *MoNIT4*, *PIG1*, *TRA1* and *XLR1*) have been identified in the rice blast fungus [17,39,41,72,73], and 4 of them were also in our Zn₂Cys₆ TF gene knockout mutant set. The expression of *MoNIT4* and *MoCOD1* were up-regulated during conidiation [41], and their mutants $\Delta Monit4$ and $\Delta Mocod1$ had reduced conidiation [39,41], and $\Delta Mocod1$ also had lower pathogenicity [39]. In our study, the mutants $\Delta Monit4$ and $\Delta Mocod1$ were also defective in conidiation or pathogenicity; furthermore, our mutant $\Delta Mocod1$ did not produce any conidia on CM medium. Similar to a previous report [74], our $\Delta Tra1$ mutant also showed defects in conidial germination. *PIG1* was identified to be involved in melanin biosynthesis, but not confirmed by knockout [73]; however, the $\Delta pig1$ mutant did not show notable phenotypes in fungal development in our study. These small discrepancies in mutant phenotypes with previously reported data may be due to the differences in wild-type strains and experimental conditions.

TFs Mnh6, Moatfl, Mocrzl and Mstul were reported to regulate hyphal growth in the rice blast fungus, and the colony growth of the mutants $\Delta mnh6$, $\Delta Moatfl$, $\Delta Mocrzl$ and $\Delta Mstul$ was about 70–90% of the wild-type strain [7–9,24]. We found 27 Zn₂Cys₆ TF genes involved in fungal growth, and 3 genes GCC1, GPF1 and GTA1 were required for normal colony growth. Interestingly, the colony growth of $\Delta gta1$ was only about 56% of that of wild-type strain, and GTA1 is a gene which affects colony growth the most among known TF genes in the rice blast fungus. GTA1 also functions in conidiation and pathogenicity to plants.

The change in asexual reproduction in a mutant is a phenotype that mostly happens when TF genes or other genes are deleted in the rice blast fungus. The TF gene-deleted mutants ($\Delta com1$, $\Delta con7$, $\Delta cos1$, $\Delta mnh6$, $\Delta Mohox2$) showed reduced conidiation [7,11-13,15,16]. In our study, 25 Zn₂Cys₆ TF genes were identified as being involved in fungal conidiation. Of them, CCA1, GCC1, MoCOD1 and CONx1 are necessary for the differentiation of conidiophores. Unexpectedly, the deletion of four Zn₂Cys₆ TF genes (CNF1, CNF2, CNF3 and CNF4) led to increased conidial production. In particular, Cnf1 is the strongest negative regulatory factor of conidial production identified in the rice blast fungus until now. The deletion of CNF1 led to earlier and more differentiation of spore-bearing hyphae than in the wildtype strain. Impressively, the mutants $\Delta cnf1$ and $\Delta cnf2$ produced more conidia, but their conidia had reduced virulence to barley and rice. Conidia are the main way to spread the rice blast disease. It is clear that a balance exists between the ability to produce conidia and the pathogenicity to plant in the phytopathogenic fungus.

We identified 2641 genes regulated by GPF1 and 3144 genes by CNF2 through RNA-seq. Interestingly, 2021 genes were regulated in the same direction in $\Delta gpf1$ and $\Delta cnf2$. This fact suggested GPF1 and CNF2 have similar mechanisms in the regulation of fungal pathogenicity. Nearly sixty DEGs in $\Delta gpf1$ and $\Delta cnf2$ were confirmed to be required for fungal pathogenicity (Table 3). In particular, the expression of two pathogenicity-required genes, an APSES TF gene MSTU1 [8] and a hydrophobin gene MPG1 [75], were greatly down-regulated in $\Delta g p f 1$ (7.80- and 12.49- fold, respectively) and $\Delta cnf2$ (11.85- and 15.83-fold, respectively). About half of Zn₂Cys₆ TF genes and many other DNA-binding domain TF genes were regulated by GPF1 and CNF2. These data primarily revealed the gene expression network in the regulation of pathogenicity controlled by GPF1 and CNF2. It is necessary to keep the well-balanced expression of TF genes and pathogenicityrequired genes for maintenance of pathogenicity in M. oryzae.

The high-throughput analysis of mutant phenotypes showed that some TF gene mutants shared highly overlapping phenotypes, such as mycelial growth in $\Delta gcc1$, $\Delta gpf1$ and $\Delta gta1$, and pathogenicity in $\Delta gpf1$, $\Delta gta1$, $\Delta cnf1$ and $\Delta cnf2$. On the other hand, several mutant phenotypes were shared by one TF gene mutant. Also, it is fascinating to know how to regulate similar phenotypes by different TF genes. In general, every TF gene regulates the expression of many downstream genes and the expression of every gene is regulated by many TF genes, and therefore each mutant phenotype is controlled by the changes in expression of several genes. When two TF genes show similar mutant phenotype, there are at least 3 types of phenotype regulation mechanisms between them. First, TF gene A regulates another TF gene B, and TF gene B continues to regulate a group of downstream genes. Second, TF genes A and B commonly regulate a group of downstream genes. Third, TF genes A and B

regulate two different groups of downstream genes independently, but they lead to similar phenotypes. However, two or three of these gene regulation types often occur simultaneously. For example, *GPF1* and other TF genes were down-regulated in $\Delta cnf2$, while *CNF2* and other TF genes were also down-regulated in $\Delta gpf1$ (Table S11 in Text S1). A common group of genes (Figure 8) were regulated both by *CNF1* and *GPF1* which led to similar pathogenicity phenotypes in mutants, while two other different groups of genes (Figure 8) were regulated by *CNF1* and *GPF1* independently which led to different phenotypes in conidiation in mutants.

In conclusion, this study represents a major advance in a highthroughput gene knockout system suitable for filamentous fungi, provides the functional characterization of $104 \text{ Zn}_2\text{Cys}_6$ TF genes in the rice blast fungus, and reveals gene expression patterns of two virulence-required TF genes *GPF1* and *CNF2*. These studies will help us build more fungal gene-deletion mutant libraries in an economical way and uncover the transcriptional network in fungi and fungal pathogenic mechanisms.

Materials and Methods

Strains

M. oryzae strain 70-15 and its mutants, *S. cerevisiae* strain FY834, and *E. coli* strain DH5 α and *A. tumefaciens* strain AGL1 were used in this study.

Building of a yeast-*Escherichi-Agrobacterium* shuttle vector pKO1B

A 1184-bp promoter fragment of *M. oryzae H3* histone gene amplified from pKD5 [35] with primers H3sF and H3SR (Table S1 in Text S1) and 720-bp *eGFP* CDS fragment amplified from pEGFP (Clontech, USA) with primers KoGFPF and KoGFPR (Table S1 in Text S1) were inserted into the *XhoI/Eco*R1 sites and the *Eco*R1/*SacI* sites of the binary vector pCAMBIA1300 (Cambia, USA) to produce the vector pKO1. pKO1B was then produced by inserting a 2.9-kb *URA3-*2micro2_origin fragment amplified from pYES2 (Invitrogen, USA) with primers uraf and urar (Table S1 in Text S1) into the *SacII* site of pKO1 by a yeast recombinational cloning method using S. c. EasyComp Transformation Kit (Invitrogen, USA).

Primer design and construction of gene-deletion cassettes

A list of 2-kb regions on both sides of each ORF (12775 ORF totally) was retrieved from the rice blast fungus database MG8 (www.broadinstitute.org) and saved in Excel type files by a program written by Mr. Tan Cheng. 23-nt primers (Table S1 in Text S1) of each gene-specific flank (1000 bp–1500 bp in length and 1200 bp in optimum length) were designed by the BatchPrimer3 program [76]. For each gene, primers were designed and synthesized with the following common 33-nt 5' regions:

5f: GCTGTACAAGTAAGAGCTCGGTACCCGGGGATC... (Homologous to pKO1B)

5r: CCGGGAGATGTGGGGGCACTGTGGCGTTGGCACA... (Homologous to SUR gene)

3f: TTGATTATTGCACGGGAATTGCATGCTCTCACA... (Homologous to *SUR* gene)

3r: TTAAGTTGGGTAACGCCAGGGTTTTCCCAGTCA... (Homologous to pKO1B)

The flank fragments were produced from genomic DNA of *M. oryzae* strain 70-15 using Primer Star or EX Taq (TaKaRa, China) in 96-well PCR plates. The *SUR* cassette fragment was generated by PCR with primers surf and surr (Table S1 in Text S1) from pBS-SUR [35]. All PCR products were verified by agarose gel electrophoresis analysis and then used in subsequent steps without further purification. Among the 163 TF genes, both flanking fragments were successfully amplified for 142 genes. However, one or sometimes both flanking fragments failed to be amplified for 21 genes even after trying several reaction conditions.

The yeast transformation procedure was conducted following a small-scale yeast transformation protocol in the pYES2 user manual (Invitrogen, USA) in 96-well deep well plates as described by Colot *et al* [29]. The cocktail mixture was made by adding 1.8 ml competent yeast cells, 100 μ l linearized pKO1B by *XbaI* and *Hin*dIII (100 ng/ μ l), 210 μ l PCR production of *SUR* cassette, 60 μ l denatured salmon sperm DNA (Sangon, China) and 2.6 ml DMSO to 20.7 ml freshly prepared 1×LiAc/40%PEG-3350/1×TE solution (100 mM lithium acetate, pH 7.5; 50% PEG-3350; 10 mM Tris-HCl, pH 7.5; 1 mM EDTA). Next, 200 μ l of the mixture, followed by 4 μ l of 5' and 3' flank PCR fragments of the targeted gene, were pipetted into each well of a 96-deep-well plate with a multichannel pipette. The plates were sealed and processed further following the user manual.

After 3 days, the yeast cells cultured in SC-Ura liquid medium were collected and disrupted in Fastprep-24 homogenizer (MP, USA) at 4.0 m/s for 2 min. The plasmids were extracted with TIANprep yeast plasmid DNA kit (Tiangen Biotech, China) following the user manual. Next, the yeast plasmids were transformed into the competent cells of *E. coli* strain DH5 α prepared following the Inoue method for "ultra-competent" cells [77]. Four bacterial colonies on each LB plate were placed in the wells of 96-deep-well plates, with each well containing 1 ml of LB liquid medium with 50 µg/ml kanamycin. The plates were shaken at 220 rpm overnight at 37°C.

The correctness of homologous recombinational cloning for gene-deleted cassettes was confirmed by bacterial double PCR. Common primers were designed to amplify the 5' and 3' flanking fragment of each target gene-deleted cassette. Forward primer KO1Bf1 for the 5' flanking fragment and reverse primer KO1Br2 for the 3' flanking fragment were located inside pKO1B, and reverse primer SURr1 for the 5' flanking fragment and forward primer SURf2 for the 3' flanking fragment were located in SURcassette (Table S1 in Text S1). The resulting PCR products in knockout plasmids constructed correctly would be 1.0-1.5 kb for 5' flanking fragment and 1.5–2.0 kb for 3' flanking fragment. The gene deletion cassettes were further confirmed by sequencing with the primers SURf2 and SURr1. PCR and DNA sequencing results showed a success rate of 100% for the recombinant plasmids (Table S2 in Text S1). We found that it was enough to confirm the recombinant plasmids by PCR and not necessary to confirm by sequencing plasmids. The correctly built knockout plasmids were extracted from the bacterial cultures using the AxyPrep-96 plasmid purification kit (Axygen, China) following the protocol in the user manual.

Agrobacterium tumefaciens-mediated transformation with knockout vectors

The knockout plasmids were transformed into the competent cells of *A. tumefaciens* strain AGL1 using the freeze/thaw shock transformation method following the procedure described elsewhere [67]. The plasmids in *Agrobacterium* cell were confirmed by culture PCR with primer set SurP1 and SurP2 (Table S1 in Text S1) for a 368-bp *SUR* gene fragment.

M. oryzae strain 70-15 was grown on CM medium for 12 days at 25° C under constant fluorescent light, and the conidia were harvested and transformed with the knockout plasmids mediated

by A. tumefaciens according to a previously reported procedure [67], whereas performed in groups, usually 24 or 48 genes in a batch each time. The nitrocellulose membrane strips containing the conidia co-cultivated with A. tumefaciens were placed on AIM medium in the dark at 22°C for 2 days and then transferred onto the selection defined complex medium (DCM; 0.17% yeast nitrogen base without amino acids, 0.2% asparagine, 0.1% ammonium nitrate and 1% glucose, pH 6.0 with Na₂HPO₄) plates containing 100 µg/ml sulfonylurea, 50 µg/ml kanamycin, 400 µg/ml cefotaxime and 60 µg/ml streptomycin. Sulfonylurearesistant transformants grown on selection medium were individually transferred onto new selection DCM plates using sterile toothpicks. A total of 8741 primary transformants corresponding to 133 Zn₂Cys₆ TF genes were selected on the basis of sulfonylurea resistance (Table S2 in Text S1). No transformant was obtained for the other nine genes.

Identification of gene-deleted mutants by GFP fluorescence, double PCR and qPCR

After culture for 2 days, a little mycelium of each transformant was picked out and placed onto a glass slide. We usually placed the mycelia of six transformants on a slide in proper order. The green fluorescence emitted by transformants was then observed one by one under a fluorescence microscope. Ectopic transformants emitted green fluorescence and null mutants did not when excited under the fluorescence microscope (Figure 1D). The transformants without green fluorescence were picked out and inoculated on a new selective plate and grown for 3 days. In total, 3191 transformants without green fluorescence were screened from the primary transformants.

Extraction of genomic DNA was performed following the CTAB protocol of Rogers and Bendich [70] with modifications. A small piece (>9 mm²) of mycelium along with some medium in the selective plate was transferred to a 2-ml round-bottom tube, and 400 μ l ddH₂O were added along with 0.1 g porcelain beads. The mycelial cells were disrupted in a Fastprep-24 homogenizer at 4.0 m/s for 2 min; 400 μ l 4×CATB buffer (4% CTAB, 100 mM Trisma base, 20 mM EDTA, 1.4 M NaCl) were then added to each tube and the tubes incubated at 65°C for 30 min. Next, genomic DNA was extracted following the normal CTAB protocol [70]. In total, genomic DNA of the 3191 transformants was extracted with the CTAB method in a high-throughput way.

The gene-deleted mutants from the transformants without green fluorescence were identified by negative screening double PCR. Double PCR was performed using primers CKF and CKR internal to the targeted gene and primers Tbl-gF and Tbl-gR for the β -tubulin gene (Table S1 in Text S1). PCR was performed in 25-µl reaction mixtures in 96-well PCR plates: 0.15 µl primers Tbl-gF and Tbl-gR (20 µM), 0.5 µl primers CKF and CKR (20 µM), 2.5 µl 10×PCR buffer, 0.4 µl dNTP mix (25 µM), 0.3 µl Taq (5 U/ μ l), 19.5 μ l ddH₂O and 1 μ l genomic DNA. The PCR program was: 94°C 3 min followed by 35 cycles of 94°C for 30 s, 57°C for 30 s and 72°C for 30 s, and a final extension at 72°C for 10 min. The PCR products were detected by 1.0% agarose gel electrophoresis. If the targeted gene was deleted in a transformant, there was only one band for β -tubulin with 554 bp in length, appearing as a positive control; otherwise, in ectopic transformants and the wild-type strain there were two bands with one of 300-400 bp (the targeted gene) and another of 554 bp (β -tubulin) (Figure 1E). After checking 2280 genomic DNA samples, we identified 477 transformants with deletions in a total of 104 TF genes (Table S2 in Text S1, Figure S4A). No null mutants were found for the other 29 genes after screening all DNA samples.

For the transformants identified as null mutants in the negative screening double PCR, a second PCR was performed to verify the gene deletion event. One primer p1 or p4 (Table S1 in Text S1) was limited in the genomic DNA outside the 5' or 3' flanking fragment in gene-deletion cassettes, and another primer p2 or p3 (Table S1 in Text S1) was limited in the *SUR* gene in gene-deletion cassettes. If the targeted gene was deleted, there was a band with 1.2-2.0 kb in length appearing in the gel. Otherwise, there was no band for the ectopic transformant and wild-type strain 70-15 (Figure 1F). A total of 477 transformants of 104 genes identified by the preceding negative screening PCR were also identified as null mutants in this positive screening PCR (Table S2 in Text S1, Figure S4A).

The copies of transformed gene-deletion cassettes in null mutants were identified by qPCR. The concentration of DNA samples of null mutants was standardized to 25 ng/µl by DNA fluorometry. The primers for the genomic DNA of SUR gene are qSurF and qSurR (Table S1 in Text S1). The genomic DNA fragment coding for β -tubulin gene (one copy in the genome) was selected as a control. The primers of β -tubulin gene are gtblF and qtblR (Table S1 in Text S1). PCR mixture of 25 µl was prepared: 12.5 µl 2×RT-PCR buffer (SYBR Green, premix Ex Taq, TaKaRa), 0.5 µl forward primer and 0.5 µl reverse primer (20 µM), 2 µl genomic DNA, and 9.5 µl ddH₂O. Real-time PCR was performed in a Mastercycler (Eppendorf, USA) with the following program: 95°C for 2 min, 40 cycles (95°C for 10 s, 60°C for 20 s), and ending with a melting curve step. Each sample was repeated three times. The copies of transformed gene-deletion cassettes in null mutants were calculated by comparing with the data of the β -tubulin gene and wild-type strain. The exogenous gene was inserted into the mutant's genomic DNA in an integral multiple number. If the copy number of the selective marker gene was 1.0 ± 0.2 times the β -tubulin gene in genomic DNA, the mutant was regarded as containing a single insertion of the selective marker gene. As a result, 477 null mutants of 104 genes were identified as single insertion null mutants (Table S2 and Table S3 in Text S1).

The gene deletion events in mutants of ten randomly selected TF genes were reconfirmed by Southern blot (Figure S4B), which was performed according to a previously reported procedure [7].

Complementation of null mutants with native genes

The mutants were complemented with native gene copies of the wild-type strain 70-15. First, the pKO1B-HPH was built by the replacement of Ph3-GFP cassette with a HPH gene from pCB1003 [78] in pKO1B. The copies of the complementation genes were then cloned from the genomic DNA of the wild-type strain with the primers listed in Table S1 in Text S1 and were inserted into the XbaI/HindIII sites of pKO1B-HPH by the yeast recombinational cloning method. The constructed complementation plasmids were transformed into the mutants using the ATMT method, and the transformants were screened on selective medium containing 200 μ g/ml hygromycin B. The gene-rescued transformants were identified by RT-PCR at the mRNA level (Figure S4C) with primers specific for the targeted genes (Table S1 in Text S1).

Phenotypic screening analyses of knockout mutants at development stages

The phenotypic screening analyses were performed by testing 20–30 knockout mutants and one control (the wild-type strain) once in a batch according to previously reported protocols [7,79,80]. Mutant phenotypes were assayed in triplicate with five replicates for each strain. The values of colony growth,

conidiation, conidial germination and appressorium formation of mutants in the different experimental groups were compared after normalization with the wild-type strain 70-15 in the same group.

Colony growth, morphology and conidiation. Mycelial blocks of 5 mm from 9-day-old wild-type strain 70-15 or mutants were inoculated in the center of CM solid medium in 6-cm plates followed by culture at 25°C under constant fluorescent light. For colony growth, the diameter of the mycelial colony was recorded and colony images were captured at 6 days post inoculation (dpi). For conidiation, the whole conidia of a 7-day-old CM-grown culture in 6-cm plates were harvested and counted using a hemacytometer.

Conidialgerminationandappressoriumformation.Conidia were harvested from a 10-day-old colonygrowing on a CM plate by adding 0.5 ml 25 ppm Tween-20solution to the surface of the culture. The suspension was pipettedup and down once more to ensure enough conidia for the assay. A40-µl aliquot of spore suspension was dropped onto a sterilizedplastic coverslip, which was incubated in a moist chamber at 25°Cfor 4 and 24 h under dark conditions. A total of 300–500 conidiawere counted for conidial germination at 4 hpi and for appressorium formation at 24 hpi.

Phenotypic screening analyses of knockout mutants under stress conditions

The phenotypes of the mutants were also screened under different stress conditions (Table S4 in Text S1) with 20–30 knockout mutants once in a batch and the wild-type strain in an experiment. Four pinpoint-like mycelia of 9-day-old strains were inoculated in the solid medium of different stress conditions in a 9cm plate with a space interval between each other and then incubated at 25°C. Colony images were captured and the diameters of the mycelial colonies were recorded at 6 dpi. The growth rates under stress conditions were compared between strains after normalization with the wild-type strain.

The phenotypes of the mutants of 5 TF genes and their complementation strains under stress conditions were reassessed. Mycelial blocks of 5 mm from 9-day-old strains were inoculated in the center of solid medium of different stress conditions in 6-cm plates followed by culture at 25°C under constant fluorescent light, along with the strains grown on CM medium as controls. Colony images were captured and the diameters of the mycelial colonies were recorded at 8 dpi. The experiments were performed in triplicate with five replicates for each strain. Growth inhibition rate of each mutant was calculated as the growth inhibition rate = (colony diameter on CM medium–colony diameter under stress condition)÷ colony diameter on CM medium×100.

Pathogenicity tests on barley and rice

The virulence tests on barley (*Hordeum vulgare*) and rice (*Oryza sativa* cv CO39) were performed following previously reported protocols [7]. For assays with leaf explants of barley or rice, 5-mm mycelium blocks of mutants (along with the wild-type strain 70-15 and mock) were inoculated on the leaf, followed by incubation in a wet box at 25°C for 4 days. For spraying assay on rice seedlings, 4 ml conidial suspension $(1 \times 10^5 \text{ conidia/ml})$ containing 0.2% (w/ v) gelatin were sprayed onto 15–20 rice seedlings between the third and fourth leaf stages using an artist's airbrush. Inoculated plants were placed in a wet box at 25 C for 2 days and then allowed to grow in controlled environment chambers with a photoperiod of 12 h using fluorescent lights for 5 days.

Plant penetration by the wild-type strain or mutant appressoria was assayed on leaf explants of barley or onion cuticles according to previously reported protocols [7]. A droplet of conidia (5×10^4)

conidia/ml) was inoculated on barley leaf cuticles or onion cuticles and incubated at 25°C for 24 or 48 h. The barley leaves were then treated in methanol (overnight, room temperature) to remove chlorophyll, and fixed in alcoholic lactophenol (1 h, 95% alcohol/ lactophenol = 2:1). Appressorium penetration on barley leaf cuticle or onion cuticles was assessed under a microscope.

RNA-sequencing analysis

The wild-type strain, $\Delta gpf1$ and $\Delta cnf2$ were grown in liquid CM medium at 25°C with shaking at 180 rpm for 2 days. The cultures were then collected and incubated in H₂O at 25°C for 4 h. The treatments were repeated in triplicate for each strain. Total RNA was extracted from ground mycelia in liquid nitrogen with the RNeasy Plant Mini Kit (QIAGEN), and mRNA was isolated using AMPure XP beads (Beckman). RNA-seq libraries were constructed using NEBNext RNA sample preparation kit (NEB) in accordance with the standard low-throughput protocol. Samples were sequenced in a 1×100 nt way on an Illumina Hiseq2500 instrument using the TruSeq PE Cluster Kit v3 - cBot -HS (Illumina) and TruSeq SBS Kit v3-HS (Illumina). The clean reads were generated by removing adaptor sequences, tags with> 10% "N", and low quality tags by FastQC (http://www. bioinformatics.babraham.ac.uk), and were mapped to the M. oryzae genome database (MG8) (www.broadinstitute.org) using Tophat software [81]. The data from triple biological replicates were then analyzed using Cufflinks software and resulted in quantified genome-wide transcript levels of genes (expressed in fragments per kilobase of exon model per million mapped fragments - FPKM) [82]. The significant differences in FPKM between different samples were assayed by the Cuffdiff component of the Cufflinks package [82].

List of genes deleted in this study

The functions of 104 Zn_2Cys_6 TF genes were studied by knockout and phenotypic analysis in this study (Table 1).

Accession numbers

RNA-sequencing data were deposited in NCBI's Gene Expression Omnibus (GEO accession number GSE57146).

Supporting Information

Figure S1 The colonies of *M. oryzae* strains on CM medium. The mutants of $104 \text{ Zn}_2\text{Cys}_6$ transcription factor genes and the wild-type strain 70-15 were cultured at 25°C for 6 days. (PDF)

Figure S2 Pathogenicity screening assay of the mutants on barley leaf explants. The mycelial agar plugs of the mutants of $104 \text{ Zn}_2\text{Cys}_6$ transcription factor genes and the wildtype strain 70-15 were placed on intact barley leaves for 4 days. (PDF)

Figure S3 Pathogenicity screening assay of the mutants of 104 Zn_2Cys_6 transcription factor genes on rice. (A) The rice seedlings were sprayed with conidial suspension $(1 \times 10^5 \text{ spores/ml})$ of 101 *M. oryzae* mutants and cultured for 7 days. (B) The mycelial agar plugs of the mutants of 3 TF genes and the wild-type strain were placed on intact rice leaves for 4 days. (PDF)

Figure S4 Knockout and complementation of Zn_2Cys_6 transcription factor genes in *M. oryzae*. (A) Knockout event of null mutants confirmed by PCR. One mutant of each TF gene (which was assayed in mutant phenotype) was selected as a representative to show the PCR identification results. The size of DNA standards are indicated on the right of lanes (M). WT, M. oryzae strain 70-15; **a**, bands for β -tubulin; **b**, bands for the targeted genes; c, bands for unique recombinational DNA fragments relative to the target gene-deletion event. (B) Null mutants of ten randomly selected TF genes confirmed by Southern blot. Genomic DNAs were digested with restriction enzymes shown in Figure S4B and separated on 0.7% agarose gels. The DNAs were individually hybridized with the probes (indicated in Figure S4B). Only one band was detected in mutants and its size was different from that in the wild-type strain, indicating that homologous recombination occurred at a single site. (\mathbf{C}) Complementation of 16 Zn₂Cys₆ transcription factor gene-deleted mutants. The mutants were rescued with their native copy of gene in M. oryzae strain 70-15. RT-PCR results were shown after amplification with 35 cycles. RNA was isolated from the mycelia of the wild-type strain, mutants and complemented strains grown on CM medium. a, the targeted genes; b, β -tubulin gene; 1, mutants; 2, complemented strains; and 3, wild-type strain 70-15. (PDF)

Text S1 Supplemental tables. Table S1. Primers used in this study. **Table S2**. Summary of the gene knockout of 163 Zn_2Cys_6 transcription factor genes in the rice blast fungus. **Table S3**. List of fungal-specific Zn_2Cys_6 zinc finger genes being knocked out and identification of copies of *SUR* gene in the mutants in this study. **Table S4**. Phenotypic analyses of the null mutants of 104

References

- Dean R, Van Kan JA, Pretorius ZA, Hammond-Kosack KE, Di Pietro A, et al. (2012) The Top 10 fungal pathogens in molecular plant pathology. Mol Plant Pathol 13: 414–430.
- Ou SH (1980) Pathogen variability and host resistance in rice blast disease. Annu Rev Phytopathol 18: 167–187.
- Dean RA, Talbot NJ, Ebbole DJ, Farman ML, Mitchell TK, et al. (2005) The genome sequence of the rice blast fungus *Magnaporthe grisea*. Nature 434: 980–986.
- Valent B, Chumley FG (1991) Molecular genetic analysis of the rice blast fungus, Magnaporthe grisea. Annu Rev Phytopathol 29: 443–467.
- Talbot NJ (2003) On the trail of a cereal killer: Exploring the biology of Magnaporthe grisea. Annu Rev Microbiol 57: 177–202.
- Jeon J, Park SY, Chi MH, Choi J, Park J, et al. (2007) Genome-wide functional analysis of pathogenicity genes in the rice blast fungus. Nat Genet 39: 561–565.
- Lu JP, Feng XX, Liu XH, Lu Q, Wang HK, et al. (2007) Mnh6, a nonhistone protein, is required for fungal development and pathogenicity of *Magnaporthe* grisea. Fungal Genet Biol 44: 819–829.
- Nishimura M, Fukada J, Moriwaki A, Fujikawa T, Ohashi M, et al. (2009) Mstu1, an APSES transcription factor, is required for appressorium-mediated infection in *Magnaporthe grisea*. Biosci Biotechnol Biochem 73: 1779–1786.
- Choi J, Kim Y, Kim S, Park J, Lee YH (2009) MoCRZ1, a gene encoding a calcineurin-responsive transcription factor, regulates fungal growth and pathogenicity of *Magnaporthe oryzae*. Fungal Genet Biol 46: 243–254.
- Qi Z, Wang Q, Dou X, Wang W, Zhao Q, et al. (2012) MoSwi6, an APSES family transcription factor, interacts with MoMps1 and is required for hyphal and conidial morphogenesis, appressorial function and pathogenicity of *Magnaporthe oryzae*. Mol Plant Pathol 13: 677–689.
- Yang J, Zhao X, Sun J, Kang Z, Ding S, et al. (2010) A novel protein Com1 is required for normal conidium morphology and full virulence in *Magnaporthe* oryzae. Mol Plant Microbe Interact 23: 112–123.
- Odenbach D, Breth B, Thines E, Weber RW, Anke H, et al. (2007) The transcription factor Con7p is a central regulator of infection-related morphogenesis in the rice blast fungus *Magnaporthe grisea*. Mol Microbiol 64: 293–307.
- Zhou Z, Li G, Lin C, He C (2009) Conidiophore stalk-less1 encodes a putative zinc-finger protein involved in the early stage of conidiation and mycelial infection in *Magnaporthe oryzae*. Mol Plant Microbe Interact 22: 402–410.
- Kramer B, Thines E, Foster AJ (2009) MAP kinase signalling pathway components and targets conserved between the distantly related plant pathogenic fungi *Mycosphaerella graminicola* and *Magnaporthe grisea*. Fungal Genet Biol 46: 667–681.
- Liu W, Xie S, Zhao X, Chen X, Zheng W, et al. (2010) A homeobox gene is essential for conidiogenesis of the rice blast fungus *Magnaporthe oryzae*. Mol Plant Microbe Interact 23: 366–375.
- Kim S, Park SY, Kim KS, Rho HS, Chi MH, et al. (2009) Homeobox transcription factors are required for conidiation and appressorium development in the rice blast fungus *Magnaporthe oryzae*. PLoS Genet 5: e1000757.

Zn₂Cys₆ transcription factor genes. **Table S5**. Comparison of phenotypes of previously characterized transcription factors between three fungal mutants (*M. oryzae*, *F. graminearum* and *N. crassa*). **Table S6**. Differentially expressed genes in $\Delta gpf1$ when compared with the wild-type strain (FDR<0.05). **Table S7**. Differentially expressed genes in $\Delta cnf2$ when compared with the wild-type strain (FDR<0.05). **Table S8**. Differentially expressed genes shared by both $\Delta gpf1$ and $\Delta cnf2$. **Table S9**. Differentially expressed genes in $\Delta cnf2$ when compared with $\Delta gpf1$ (FDR<0.05). **Table S10**. *GPF1*- or *CNF2*-dependent genes that are dispensable for pathogenicity reported in previous studies. **Table S11**. Zn2Cys6 transcription factor genes that were listed as DEGs of $\Delta gpf1$ or $\Delta cnf2$.

(XLSX)

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Author Contributions

Conceived and designed the experiments: JL. Performed the experiments: HC JL LZ PH. Analyzed the data: JL HC LZ. Contributed reagents/ materials/analysis tools: FL. Wrote the paper: JL.

- Battaglia E, Klaubauf S, Vallet J, Ribot C, Lebrun MH, et al. (2013) Xlr1 is involved in the transcriptional control of the pentose catabolic pathway, but not hemi-cellulolytic enzymes in *Magnaporthe oryzae*. Fungal Genet Biol 57: 76– 84.
- Li Y, Liang S, Yan X, Wang H, Li D, et al. (2010) Characterization of MoLDB1 required for vegetative growth, infection-related morphogenesis, and pathogenicity in the rice blast fungus *Magnaporthe oryzae*. Mol Plant Microbe Interact 23: 1260–1274.
- Yan X, Li Y, Yue X, Wang C, Que Y, et al. (2011) Two novel transcriptional regulators are essential for infection-related morphogenesis and pathogenicity of the rice blast fungus *Magnaporthe oryzae*. PLoS Pathog 7: e1002385.
- Mehrabi R, Ding S, Xu JR (2008) MADS-box transcription factor migl is required for infectious growth in Magnaporthe grisea. Eukaryot Cell 7: 791–799.
- Park G, Xue C, Zheng L, Lam S, Xu JR (2002) MST12 regulates infectious growth but not appressorium formation in the rice blast fungus *Magnaporthe* grisea. Mol Plant Microbe Interact 15: 183–192.
- Li G, Zhou X, Kong L, Wang Y, Zhang H, et al. (2011) MoSfil is important for virulence and heat tolerance in *Magnaporthe oryzae*. PLoS One 6: e19951.
- Guo M, Chen Y, Du Y, Dong Y, Guo W, et al. (2011) The bZIP transcription factor MoAP1 mediates the oxidative stress response and is critical for pathogenicity of the rice blast fungus *Magnaporthe oryzae*. PLoS pathogens 7: e1001302.
- 24. Guo M, Guo W, Chen Y, Dong S, Zhang X, et al. (2010) The basic leucine zipper transcription factor Moatf1 mediates oxidative stress responses and is necessary for full virulence of the rice blast fungus *Magnaporthe oryzae*. Molecular plant-microbe interactions 23: 1053–1068.
- Lee K, Singh P, Chung WC, Ash J, Kim TS, et al. (2006) Light regulation of asexual development in the rice blast fungus, *Magnaporthe oryzae*. Fungal Genet Biol 43: 694–706.
- Nguyen QB, Kadotani N, Kasahara S, Tosa Y, Mayama S, et al. (2008) Systematic functional analysis of calcium-signalling proteins in the genome of the rice-blast fungus, *Magnaporthe oryzae*, using a high-throughput RNAsilencing system. Mol Microbiol 68: 1348–1365.
- Son H, Seo YS, Min K, Park AR, Lee J, et al. (2011) A phenome-based functional analysis of transcription factors in the cereal head blight fungus, *Fusarium graminearum*. PLoS Pathog 7: e1002310.
- Yu JH, Hamari Z, Han KH, Seo JA, Reyes-Dominguez Y, et al. (2004) Double-joint PCR: a PCR-based molecular tool for gene manipulations in filamentous fungi. Fungal Genet Biol 41: 973–981.
- Colot HV, Park G, Turner GE, Ringelberg C, Crew CM, et al. (2006) A highthroughput gene knockout procedure for Neurospora reveals functions for multiple transcription factors. Proc Natl Acad Sci U S A 103: 10352–10357.
- Talbot NJ, Foster AJ (2001) Genetics and genomics of the rice blast fungus Magnaporthe grisea: developing an experimental model for understanding fungal diseases of cereals. Advances in Botanical Research 34: 263–287.
- Villalba F, Collemare J, Landraud P, Lambou K, Brozek V, et al. (2008) Improved gene targeting in *Magnaporthe grisea* by inactivation of MgKU80

required for non-homologous end joining. Fungal Genetics and Biology 45: 68-75.

- Takahashi T, Masuda T, Koyama Y (2006) Enhanced gene targeting frequency in ku70 and ku80 disruption mutants of Aspergillus sojae and Aspergillus oryzae. Molecular Genetics and Genomics 275: 460–470.
- Nayak T, Szewczyk E, Oakley CE, Osmani A, et al. (2006) A Versatile and Efficient Gene-Targeting System for Aspergillus nidulans. Genetics 172: 1557– 1566.
- 34. da Silva Ferreira ME, Kress MR, Savoldi M, Goldman MH, Hartl A, et al. (2006) The akuB(KU80) mutant deficient for nonhomologous end joining is a powerful tool for analyzing pathogenicity in *Aspergillus fumigatus*. Eukaryot Cell 5: 207–211.
- Li HJ, Lu JP, Liu XH, Zhang LL, Lin FC (2012) Vectors building and usage for gene knockout, protein expression and fluorescent fusion protein in the rice blast fungus. Journal of Agricultural Biotechnology 20: 94–104.
- Park J, Jang S, Kim S, Kong S, Choi J, et al. (2008) FTFD: an informatics pipeline supporting phylogenomic analysis of fungal transcription factors. Bioinformatics 24: 1024–1025.
- Marchler-Bauer A, Lu S, Anderson JB, Chitsaz F, Derbyshire MK, et al. (2010) CDD: a Conserved Domain Database for the functional annotation of proteins. Nucleic Acids Res 39: D225–229.
- Yuan GF, Fu YH, Marzluf GA (1991) nit-4, a pathway-specific regulatory gene of *Neurospora crassa*, encodes a protein with a putative binuclear zinc DNAbinding domain. Mol Cell Biol 11: 5735–5745.
- Chung H, Choi J, Park S-Y, Jeon J, Lee YH (2013) Two conidiation-related Zn(II)₂Cys₆ transcription factor genes in the rice blast fungus. Fungal Genetics and Biology 61: 133–141.
- Tani S, Katsuyama Y, Hayashi T, Suzuki H, Kato M, et al. (2001) Characterization of the amyR gene encoding a transcriptional activator for the amylase genes in *Aspergillus nidulans*. Curr Genet 39: 10–15.
- Park SY, Choi J, Lim SE, Lee GW, Park J, et al. (2013) Global expression profiling of transcription factor genes provides new insights into pathogenicity and stress responses in the rice blast fungus. PLoS Pathog 9: e1003350.
- Donofrio NM, Oh Y, Lundy R, Pan H, Brown DE, et al. (2006) Global gene expression during nitrogen starvation in the rice blast fungus, *Magnaporthe* grisea. Fungal Genet Biol 43: 605–617.
- 43. Talbot NJ, McCafferty HRK, Ma M, Moore K, Hamer JE (1997) Nitrogen starvation of the rice blast fungus *Magnaporthe grisea* may act as an environmental cue for disease symptom expression. Physiological and Molecular Plant Pathology 50: 179–195.
- Ebbole DJ, Jin Y, Thon M, Pan H, Bhattarai E, et al. (2004) Gene discovery and gene expression in the rice blast fungus, *Magnaporthe grisea*: analysis of expressed sequence tags. Mol Plant Microbe Interact 17: 1337–1347.
- Wang Y, Wu J, Park ZY, Kim SG, Rakwal R, et al. (2011) Comparative secretome investigation of *Magnaporthe oryzae* proteins responsive to nitrogen starvation. J Proteome Res 10: 3136–3148.
- Chen GQ, Liu XH, Zhang LL, Cao HJ, Lu JP, et al. (2013) Involvement of MoVMA11, a putative vacuolar atpase c' subunit, in vacuolar acidification and infection-related morphogenesis of *Magnaporthe oryzae*. PLoS ONE 8: e67804.
- Sweigard JA, Carroll AM, Farrall L, Chumley FG, Valent B (1998) Magnaporthe grisea pathogenicity genes obtained through insertional mutagenesis. Mol Plant Microbe Interact 11: 404–412.
- Liu S, Dean RA (1997) G protein alpha subunit genes control growth, development, and pathogenicity of *Magnaporthe grisea*. Mol Plant Microbe Interact 10: 1075–1086.
- Zhang H, Tang W, Liu K, Huang Q, Zhang X, et al. (2011) Eight RGS and RGS-like proteins orchestrate growth, differentiation, and pathogenicity of *Magnaporthe oryzae*. PLoS Pathog 7: e1002450.
- Ramanujam R, Yishi X, Liu H, Naqvi NI (2012) Structure-function analysis of Rgs1 in *Magnaporthe oryzae*: role of DEP domains in subcellular targeting. PLoS One 7: e41084.
- Nishimura M, Park G, Xu JR (2003) The G-beta subunit MGB1 is involved in regulating multiple steps of infection-related morphogenesis in *Magnaporthe* grisea. Mol Microbiol 50: 231–243.
- Zhao X, Kim Y, Park G, Xu JR (2005) A mitogen-activated protein kinase cascade regulating infection-related morphogenesis in *Magnaporthe grisea*. Plant Cell 17: 1317–1329.
- Jeon J, Goh J, Yoo S, Chi MH, Choi J, et al. (2008) A putative MAP kinase kinase, MCK1, is required for cell wall integrity and pathogenicity of the rice blast fungus, *Magnaporthe oryzae*. Mol Plant Microbe Interact 21: 525– 534.
- Xu JR, Staiger CJ, Hamer JE (1998) Inactivation of the mitogen-activated protein kinase Mps1 from the rice blast fungus prevents penetration of host cells but allows activation of plant defense responses. Proc Natl Acad Sci U S A 95: 12713–12718.
- Liu XH, Lu JP, Zhang L, Dong B, Min H, et al. (2007) Involvement of a Magnaporthe grisea serine/threonine kinase gene, MgATG1, in appressorium turgor and pathogenesis. Eukaryotic Cell 6: 997–1005.
- Lu JP, Liu XH, Feng XX, Min H, Lin FC (2009) An autophagy gene, MgATG5, is required for cell differentiation and pathogenesis in *Magnaporthe* oryzae. Curr Genet 55: 461–473.
- Dong B, Liu XH, Lu JP, Zhang FS, Gao HM, et al. (2009) MgAtg9 trafficking in *Magnaporthe oryzae*. Autophagy 5: 946–953.

- Deng YZ, Qu Z, He Y, Naqvi NI (2012) Sorting nexin Snx41 is essential for conidiation and mediates glutathione-based antioxidant defense during invasive growth in *Magnaporthe oryzae*. Autophagy 8: 1058.
- Huang K, Czymmek KJ, Caplan JL, Sweigard JA, Donofrio NM (2011) HYR1-mediated detoxification of reactive oxygen species is required for full virulence in the rice blast fungus. PLoS Pathog 7: e1001335.
- Fernandez J, Wilson RA (2014) Characterizing roles for the glutathione reductase, thioredoxin reductase and thioredoxin peroxidase-encoding genes of *Magnaporthe oryzae* during rice blast disease. PLoS One 9: e87300.
- Du Y, Zhang H, Hong L, Wang J, Zheng X, et al. (2013) Acetolactate synthases MoIIv2 and MoIIv6 are required for infection-related morphogenesis in *Magnaporthe oryzae*. Mol Plant Pathol 14: 870–884.
- Yi M, Park JH, Ahn JH, Lee YH (2008) MoSNF1 regulates sporulation and pathogenicity in the rice blast fungus *Magnaporthe oryzae*. Fungal Genet Biol 45: 1172–1181.
- Wang ZY, Thornton CR, Kershaw MJ, Debao L, Talbot NJ (2003) The glyoxylate cycle is required for temporal regulation of virulence by the plant pathogenic fungus *Magnaporthe grisea*. Mol Microbiol 47: 1601–1612.
- Liu XH, Zhuang FL, Lu JP, Lin FC (2011) Identification and molecular cloning Moplaa gene, a homologue of Homo sapiens PLAA, in *Magnaporthe* oryzae. Microbiological research 167: 8–13.
- Tong AH, Evangelista M, Parsons AB, Xu H, Bader GD, et al. (2001) Systematic genetic analysis with ordered arrays of yeast deletion mutants. Science 294: 2364–2368.
- Kim DU, Hayles J, Kim D, Wood V, Park HO, et al. (2010) Analysis of a genome-wide set of gene deletions in the fission yeast *Schizosaccharomyces* pombe. Nat Biotechnol 28: 617–623.
- Rho HS, Kang S, Lee YH (2001) Agrobacterium tumefaciens-mediated transformation of the plant pathogenic fungus, Magnaporthe grisea. Mol Cells 12: 407–411.
- Ginzinger DG (2002) Gene quantification using real-time quantitative PCR: an emerging technology hits the mainstream. Exp Hematol 30: 503–512.
- Ingham DJ, Beer S, Money S, Hansen G (2001) Quantitative real-time PCR assay for determining transgene copy number in transformed plants. Biotechniques 31: 132–134, 136–140.
- Rogers SO, Bendich AJ (1985) Extraction of DNA from milligram amounts of fresh, herbarium and mummified plant tissues. Plant Molecular Biology 5: 69– 76.
- Khang CH, Park SY, Lee YH, Kang S (2005) A dual selection based, targeted gene replacement tool for *Magnaporthe grisea* and *Fusarium oxysporum*. Fungal Genet Biol 42: 483–492.
- Breth B, Odenbach D, Yemelin A, Schlinck N, Schroder M, et al. (2013) The role of the Tra1p transcription factor of *Magnaporthe oryzae* in spore adhesion and pathogenic development. Fungal Genet Biol 57: 11–22.
- 73. Tsuji G, Kenmochi Y, Takano Y, Sweigard J, Farrall L, et al. (2000) Novel fungal transcriptional activators, Cmr1p of *Colletotrichum lagenarium* and pig1p of *Magnaporthe grisea*, contain Cys2His2 zinc finger and Zn(II)₂Cys₆ binuclear cluster DNA-binding motifs and regulate transcription of melanin biosynthesis genes in a developmentally specific manner. Mol Microbiol 38: 940–954.
- Zheng W, Zhao Z, Chen J, Liu W, Ke H, et al. (2009) A Cdc42 ortholog is required for penetration and virulence of *Magnaporthe grisea*. Fungal Genet Biol 46: 450–460.
- Talbot NJ, Ebbole DJ, Hamer JE (1993) Identification and characterization of MPG1, a gene involved in pathogenicity from the rice blast fungus Magnaporthe grisea. Plant Cell 5: 1575–1590.
- You FM, Huo N, Gu YQ, Luo MC, Ma Y, et al. (2008) BatchPrimer3: a high throughput web application for PCR and sequencing primer design. BMC Bioinformatics 9: 253.
- Sambrook J, Russell DW (2001) Molecular cloning, a labortary manual (3nd ed). Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York, USA.
- Carroll AM, Sweigard JA, Valent B (1994) Improved vectors for selecting resistance to hygromycin. Fungal Genet Newsl 41.
- Motoyama T, Ochiai N, Morita M, Iida Y, Usami R, et al. (2008) Involvement of putative response regulator genes of the rice blast fungus *Magnaporthe oryzae* in osmotic stress response, fungicide action, and pathogenicity. Curr Genet 54: 185–195.
- Zhang LL, Cao HJ, Li XD, Lin FC, Feng XX, et al. (2013) MoTCTP, a homolog of translationally controlled tumor protein, is required for fungal growth and conidiation in *Magnaporthe oryzae*. Chinese J Cell Biol 35: 1141– 1154.
- Trapnell, Pachter L, Salzberg SL (2009) TopHat: discovering splice junctions with RNA-Seq. Bioinformatics 25: 1105–1111.
- Trapnell C, Williams BA, Pertea G, Mortazavi A, Kwan G, et al. (2010) Transcript assembly and quantification by RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. Nat Biotechnol 28: 511–515.
- Zhang H, Zhao Q, Guo X, Guo M, Qi Z, et al. (2014) Pleiotropic Function of the Putative Zinc-Finger Protein MoMsn2 in *Magnaporthe oryzae*. Mol Plant Microbe Interact 27: 446–460.
- Oh Y, Franck WL, Han S-O, Shows A, Gokce E, et al. (2012) Polyubiquitin Is Required for Growth, Development and Pathogenicity in the Rice Blast Fungus Magnaporthe oryzae. PloS one 7: e42868.

- Chen J, Zheng W, Zheng S, Zhang D, Sang W, et al. (2008) Rac1 is required for pathogenicity and Chm1-dependent conidiogenesis in rice fungal pathogen *Magnaporthe grisea*. PLoS Pathog 4: e1000202.
- Saitoh H, Fujisawa S, Mitsuoka C, Ito A, Hirabuchi A, et al. (2012) Large-scale gene disruption in *Magnaporthe oryzae* identifies MC69, a secreted protein required for infection by monocot and dicot fungal pathogens. PLoS Pathog 8: e1002711.
- Kershaw MJ, Talbot NJ (2009) Genome-wide functional analysis reveals that infection-associated fungal autophagy is necessary for rice blast disease. Proceedings of the National Academy of Sciences 106: 15967–15972.
- Song W, Dou X, Qi Z, Wang Q, Zhang X, et al. (2010) R-SNARE homolog MoSec22 is required for conidiogenesis, cell wall integrity, and pathogenesis of Magnaporthe oryzae. PLoS One 5: e13193.
- Zhang H, Zhao Q, Liu K, Zhang Z, Wang Y, et al. (2009) MgCRZ1, a transcription factor of *Magnaporthe grisea*, controls growth, development and is involved in full virulence. FEMS Microbiol Lett 293: 160–169.
- Dou X, Wang Q, Qi Z, Song W, Wang W, et al. (2011) MoVam7, a conserved SNARE involved in vacuole assembly, is required for growth, endocytosis, ROS accumulation, and pathogenesis of *Magnaporthe oryzae*. PloS one 6: e16439.
- Ramanujam R, Naqvi NI (2010) PdeH, a high-affinity cAMP phosphodiesterase, is a key regulator of asexual and pathogenic differentiation in *Magnaporthe* oryzae. PLoS pathogens 6: e1000897.

- Gao HM, Liu XG, Shi HB, Lu JP, Yang J, et al. (2013) MoMon1 is required for vacuolar assembly, conidiogenesis and pathogenicity in the rice blast fungus *Magnaporthe oryzae*. Res Microbiol 164: 300–309.
- Liu W, Zhou X, Li G, Li L, Kong L, et al. (2011) Multiple plant surface signals are sensed by different mechanisms in the rice blast fungus for appressorium formation. PLoS Pathog 7: e1001261.
- Patkar RN, Suresh A, Naqvi NI (2010) MoTea4-mediated polarized growth is essential for proper asexual development and pathogenesis in *Magnaporthe* oryzae. Eukaryotic cell 9: 1029–1038.
- Chen Y, Zhai S, Zhang H, Zuo R, Wang J, et al. (2013) Shared and distinct functions of two Gtil/Pac2 family proteins in growth, morphogenesis and pathogenicity of *Magnaporthe oryzae*. Environ Microbiol 16: 788–801.
- Liu XH, Lu JP, Dong B, Gu Y, Lin FC (2010) Disruption of MoCMK1, encoding a putative calcium/calmodulin-dependent kinase, in Magnaporthe oryzae. Microbiological research 165: 402–410.
- Zheng W, Chen J, Liu W, Zheng S, Zhou J, et al. (2007) A Rho3 homolog is essential for appressorium development and pathogenicity of *Magnaporthe* grisea. Eukaryotic cell 6: 2240–2250.
- Patkar RN, Ramos-Pamplona M, Gupta AP, Fan Y, Naqvi NI (2012) Mitochondrial beta-oxidation regulates organellar integrity and is necessary for conidial germination and invasive growth in *Magnaporthe oryzae*. Mol Microbiol 86: 1345–1363.
- Urban M, Bhargava T, Hamer JE (1999) An ATP-driven efflux pump is a novel pathogenicity factor in rice blast disease. EMBO J 18: 512–521.
- 101. Liu H, Suresh A, Willard FS, Siderovski DP, Lu S, et al. (2007) Rgs1 regulates multiple Gα subunits in Magnaporthe pathogenesis, asexual growth and thigmotropism. The EMBO journal 26: 690–700.