

# An NADPH-dependent genetic switch regulates plant infection by the rice blast fungus

Richard A. Wilson<sup>a,b</sup>, Robert P. Gibson<sup>a</sup>, Cristian F. Quispe<sup>b</sup>, Jennifer A. Littlechild<sup>a</sup>, and Nicholas J. Talbot<sup>a,1</sup>

<sup>a</sup>School of Biosciences, University of Exeter, Exeter EX4 4QD, United Kingdom; and <sup>b</sup>Department of Plant Pathology, University of Nebraska, Lincoln, NE 68583-0722

Edited by Jeffery L. Dangl, University of North Carolina, Chapel Hill, NC, and approved November 2, 2010 (received for review May 16, 2010)

To cause rice blast disease, the fungus *Magnaporthe oryzae* breaches the tough outer cuticle of the rice leaf by using specialized infection structures called appressoria. These cells allow the fungus to invade the host plant and proliferate rapidly within leaf tissue. Here, we show that a unique NADPH-dependent genetic switch regulates plant infection in response to the changing nutritional and redox conditions encountered by the pathogen. The biosynthetic enzyme trehalose-6-phosphate synthase (Tps1) integrates control of glucose-6-phosphate metabolism and nitrogen source utilization by regulating the oxidative pentose phosphate pathway, the generation of NADPH, and the activity of nitrate reductase. We report that Tps1 directly binds to NADPH and, thereby, regulates a set of related transcriptional corepressors, comprising three proteins, Nmr1, Nmr2, and Nmr3, which can each bind NADP. Targeted deletion of any of the Nmr-encoding genes partially suppresses the nonpathogenic phenotype of a  $\Delta$ tps1 mutant. Tps1-dependent Nmr corepressors control the expression of a set of virulence-associated genes that are derepressed during appressorium-mediated plant infection. When considered together, these results suggest that initiation of rice blast disease by *M. oryzae* requires a regulatory mechanism involving an NADPH sensor protein, Tps1, a set of NADP-dependent transcriptional corepressors, and the nonconsuming interconversion of NADPH and NADP acting as signal transducer.

fungal pathogenicity | ascomycete | cofactor

Rice blast disease represents a significant constraint on worldwide rice production, resulting in severe epidemics and overall global yield losses of 10–30% each year (1). Rice constitutes 23% of the calories consumed annually by the global human population, so understanding and controlling rice blast disease could play an important role in ensuring global food security in the future (2). To infect rice plants, the blast fungus *Magnaporthe oryzae* produces specialized infection cells called appressoria, which rupture the leaf cuticle and allow fungal hyphae to invade and colonize the host. The fungus is able to proliferate rapidly within rice cells, deriving nutrition from living tissue while evading or suppressing plant defenses (1). Understanding the regulatory mechanisms that allow the fungus to undergo these developmental transitions and to grow so effectively within its host may provide new means to control rice blast disease.

In this study, we set out to investigate the gene regulatory mechanisms that condition the ability of *M. oryzae* to respond to the nutrient status of its environment during plant infection—moving from the nutrient-free conditions of the rice leaf surface to the relatively nutrient-rich interior of the leaf. Previously, we observed a pivotal role for the biosynthetic enzyme trehalose-6-phosphate synthase (Tps1) in the regulation of carbon and nitrogen metabolism in *M. oryzae* (3, 4). Tps1 is required for production of the nonreducing disaccharide trehalose from glucose-6-phosphate (G6P) and uridine diphosphate (UDP)-glucose (3). Mutants lacking Tps1 are able to produce appressoria, but these cells do not function correctly and the fungus cannot colonize rice tissue (3). A  $\Delta$ tps1 gene deletion mutant does not produce trehalose, or the intermediate trehalose-6-phosphate (T6P), but by introducing mutations to the G6P binding pocket of Tps1, we showed that impairment of pathogenicity in *M. oryzae* by  $\Delta$ tps1 strains is not due simply to loss of trehalose biosynthesis. Introducing amino acid

substitutions into Tps1, which prevent G6P binding, rendered the fungus nonpathogenic, whereas mutations that solely affected the catalytic activity of the enzyme did not affect its role in rice blast disease, suggesting that the inability of  $\Delta$ tps1 mutants to sense intracellular G6P may be associated with their loss of virulence (4).

Tps1 appears to integrate control of both carbon and nitrogen metabolism in *M. oryzae* because without the ability to sense G6P,  $\Delta$ tps1 mutants are unable to grow on nitrate as a sole source of nitrogen. Filamentous fungi carry out nitrate reduction to ammonium via the sequential reactions of nitrate reductase (NR) and nitrite reductase. NADPH provides the reducing power for the first step of this pathway and is a cofactor for NR. In *M. oryzae*, growth on nitrate normally results in increased hexokinase (Hxk1) activity and a rise in G6P production compared with growth on ammonium as nitrogen source (4). Because both G6P dehydrogenase (G6PDH) activity and NADPH levels are significantly reduced in  $\Delta$ tps1 mutants, we reasoned that sensing of G6P by Tps1 might lead to increased NADPH production in the oxidative pentose phosphate pathway (PPP) via activation of G6P dehydrogenase (G6PDH), thus ensuring that sufficient reducing power (in the form of NADPH) is available to NR to allow growth of the fungus on nitrate. Consistent with this idea,  $\Delta$ tps1 mutants are depleted for NADPH and lack the reducing power to metabolize nitrate as a nitrogen source (4). We could not, however, determine the precise mechanism that allowed Tps1 to exert such a profound effect on the establishment of rice blast disease based solely on its role as a metabolic regulator.

In this report, we show that the virulence-associated function of Tps1 is mediated by a family of NADP-dependent transcriptional corepressors, linking its metabolic function to a wider role in the control of gene expression. This transcriptional regulatory mechanism involves the interaction of up to three GATA-factor transcriptional activators with cognate NADP-binding corepressors and is necessary for regulation of a set of genes that are expressed during appressorium-mediated plant infection in *M. oryzae*. Furthermore, we demonstrate that Tps1 directly binds NADPH, consistent with the operation of a unique NADPH-dependent genetic switch that allows the rice blast fungus to regulate gene expression in response to rapidly fluctuating changes in nutrient and redox status during plant infection.

## Results

**Tps1 Regulates Nitrogen Source Utilization Through the Oxidative Pentose Phosphate Pathway.** We initially set out to investigate the mechanism by which *M. oryzae* Tps1 regulates nitrogen source utilization and to determine whether this was important for its role in rice blast disease. Tps1 is essential for *M. oryzae* to grow on nitrate as a sole nitrogen source and we therefore needed to establish whether nitrate metabolism was necessary for fungal

Author contributions: N.J.T. designed research; R.A.W. and C.F.Q. performed research; R.P.G., C.F.Q., and J.A.L. contributed new reagents/analytic tools; R.A.W., R.P.G., J.A.L., and N.J.T. analyzed data; and R.A.W. and N.J.T. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

<sup>1</sup>To whom correspondence should be addressed. E-mail: N.J.Talbot@exeter.ac.uk.

This article contains supporting information online at [www.pnas.org/lookup/suppl/doi:10.1073/pnas.1006839107/-DCSupplemental](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1006839107/-DCSupplemental).

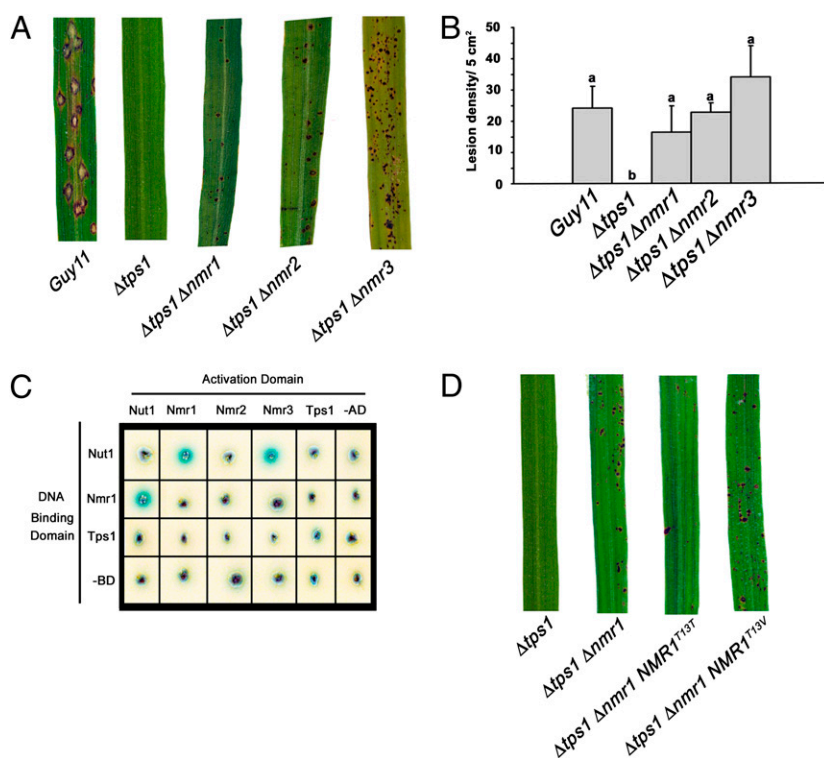


Interestingly, overexpression of *G6PDH* in a  $\Delta tps1$  mutant background partially restored the ability to cause rice blast disease in a susceptible cultivar, CO-39, as shown in Fig. 1A. We conclude that elevation of G6PDH activity, which consumes G6P and generates NADPH, can at least partially compensate for the loss of trehalose-6-phosphate synthase in  $\Delta tps1$  mutant of *M. oryzae*.

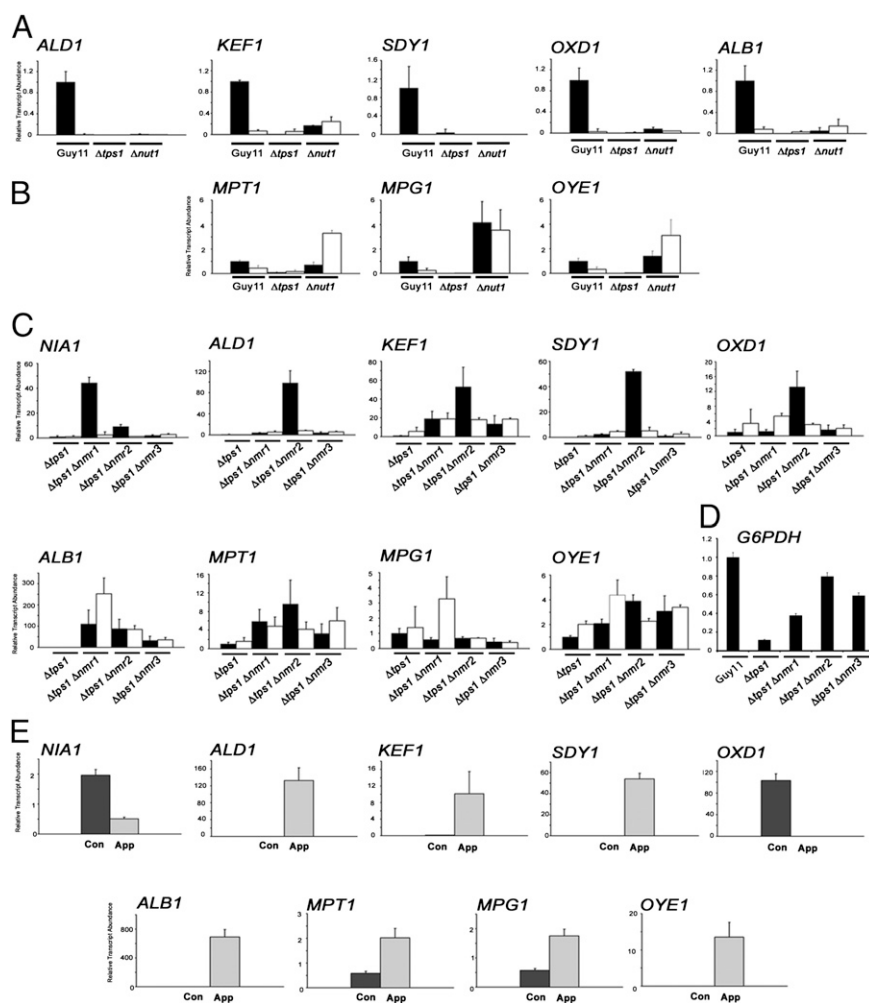
**Tps1 Can Directly Bind NADPH.** As a result of the importance of G6PDH activity in the generation of NADPH, we wondered whether Tps1 might be directly regulated by the ratio of NADP/NADPH. We therefore purified recombinant Tps1 protein and titrated it against NADPH to establish whether Tps1 was able to bind the cofactor. A specific interaction between Tps1 and NADPH was detected, as monitored by a quenching of NADPH fluorescence at 465 nm in the presence of increasing concentrations of Tps1 (Fig. 1B). Furthermore, we found evidence that NADPH could be displaced from Tps1 by the presence of G6P because the Tps1-mediated reduction in NADPH fluorescence could be reversed by the addition of excess G6P, as shown in Fig. 1C. We investigated the specificity of NADPH binding in more detail by measuring the intrinsic tryptophan fluorescence of Tps1 in the presence or absence of NADPH. Fig. S3.4 demonstrates the tryptophan quenching occurs in the active site of Tps1 in the presence of NADPH. From these data, the calculated Hill coefficient, used to determine the dissociation constant for a substrate bound to a macromolecule, gave a binding constant of  $3.96 \pm 0.06 \mu\text{M}$  and a Hill slope of  $5.18 \pm 0.35$ . This value is consistent with cooperative binding and demonstrates that binding of NADPH by Tps1 is a specific interaction. The high Hill coefficient for Tps1 and NADPH also implies that although only one molecule is present in the active site of each protein, NADPH binding might promote higher-order protein conformations, such as Tps1 tetramers. Using our modeled structure of Tps1, based on the solution structure of the bacterial Tps1 homolog *otsA* (4), we sought a structural explanation of the interaction of NADPH with Tps1. The native substrates of Tps1, UDP-glucose, and G6P, when modeled into the Tps1 active site, have calculated energies of binding of  $-19$  and  $-14$  kcal/mol, respectively, as shown in Fig. 1D. However, NADPH can also be modeled into the active site

with a calculated energy of binding of  $-27$  kcal/mol (Fig. 1D). The structural similarity between NADPH and the native Tps1 substrates, coupled with the similarity in predicted energies of binding, is consistent with a mechanism whereby NADPH can reversibly displace G6P and UDP-glucose from the Tps1 active site to inhibit Tps1 activity during conditions of elevated NADPH levels. Conversely, when NADPH is consumed by NADPH-requiring enzymes (such as NR), G6P and UDP-glucose can displace NADPH at the active site and, through G6P sensing, activate NADPH production by G6PDH. This process is consistent with a feedback loop model controlling Tps1-dependent G6PDH activity in response to available G6P and NADPH.

**Identification and Functional Characterization of NADP-Dependent Nmr Corepressors.** To investigate how a Tps1/NADPH-dependent regulatory process might regulate fungal virulence, we next investigated how Tps1 regulates gene expression.  $\Delta tps1$  mutants do not express the NR-encoding gene *NIA1* when exposed to nitrate and instead adopt an ammonium-responsive status (including the expression of genes encoding ammonium transporters), irrespective of the nitrogen source to which the fungus is exposed (Fig. S4.4). In filamentous fungi, such as *Aspergillus nidulans* and *Neurospora crassa*, the presence of ammonium represses the expression of genes encoding enzymes to metabolize alternative nitrogen sources (7). This is due in part to a repressor protein NmrA/Nmr1, which prevents the activity of a GATA-factor transcription factor, AreA/Nit2, the central regulator of gene expression when ammonium is depleted (4). We therefore reasoned that in *M. oryzae*, Tps1-dependent transcriptional regulation of *NIA1* is likely to be mediated via control of the equivalent areA/Nit2-like transcription factor (TF), Nut1, by the repressor Nmr1 (ref. 4 and Fig. S4.4). In contrast to previously characterized free-living fungal species, however, the genome of *M. oryzae* has two additional homologs of *NMR1* (*NMR2* and *NMR3*) (8), as shown in Fig. S4B. Low stringency yeast two-hybrid analysis, using  $\beta$ -galactosidase activity as the detection assay, provided evidence of a physical interaction between Nmr1 and Nmr3 with Nut1, suggesting that two of the *M. oryzae* Nmr proteins are functional



**Fig. 2.** Fungal virulence is controlled by Tps1 via the Nmr corepressor proteins. (A) Partial restoration of virulence to  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr2$ , and  $\Delta tps1 \Delta nmr3$  double mutants of *M. oryzae*. Rice seedlings were inoculated with  $1 \times 10^4$  spores  $\text{mL}^{-1}$ . (B) Bar charts of lesion densities of Guy11,  $\Delta tps1$ ,  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr2$ , and  $\Delta tps1 \Delta nmr3$  mutants. Results are the average of three independent replicates, and error bars are the SD. Bars with the same letter are not significantly different (Student's *t* test,  $P \leq 0.05$ ). (C) Low-stringency yeast two-hybrid analysis of physical interactions among Nut1, Nmr1, Nmr2, Nmr3, and Tps1 compared with the empty vector controls. -BD are strains carrying an empty *GAL4* DNA binding domain vector. -AD are strains carrying an empty *GAL4* activation domain vector. X-Gal is supplemented in the media, and low stringency protein-protein interactions result in  $\beta$ -galactosidase gene expression and formation of blue colonies. (D) Rice infections were carried out with  $\Delta tps1$ ,  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr1:NMR1^{T13T}$ ,  $\Delta tps1 \Delta nmr1:NMR1^{T13V}$ , and  $\Delta tps1 \Delta nmr1:NMR1^{T13V}$ . Expression of the *NMR1*<sup>T13V</sup> allele does not complement the  $\Delta tps1 \Delta nmr1$  mutant phenotype. Seedlings were inoculated with  $1 \times 10^4$  spores  $\text{mL}^{-1}$ .



**Fig. 3.** qPCR analysis of *TPS1*-dependent gene expression. Genes encoding putative NAD(P)-dependent enzymes (*NIA1*, *ALD1*, *KEF1*, *SDY1*, *OXD1*, and *OYE1*), known virulence-associated genes (*MPG1* and *ALB1*), a putative metalloprotease (*MPT1*), and glucose-6-phosphate dehydrogenase (*G6PDH*) (for details, see Table S1) were analyzed for expression. Strains were grown in CM for 48 h followed by 16 h of growth in MM containing either 10 mM nitrate (filled bars) or 10 mM ammonium (open bars) as the sole nitrogen source. Gene expression results were normalized against expression of the  $\beta$ -tubulin gene (*TUB2*). Results are the average of at least three independent replicates, and error bars are the SD. (A) Bar charts showing gene expression in Guy11,  $\Delta tps1$ , and  $\Delta nut1$  mutant. *ALD1*, *KEF1*, *SDY1*, *OXD1*, and *ALB1* are nitrate-inducible and dependent on the presence of *TPS1* and *NUT1*. Expression is relative to gene expression in the wild-type grown on nitrate. (B) Bar charts showing gene expression in Guy11,  $\Delta tps1$ , and  $\Delta nut1$  mutant. *MPT1*, *MPG1*, and *OYE1* appear nitrate-inducible and *TPS1*-dependent, but independent of *NUT1*. Expression is relative to gene expression in the wild type grown on nitrate. (C) Bar charts showing that *TPS1*-dependent gene expression is partially restored in a  $\Delta tps1$  mutant by introduction of  $\Delta nmr1$ ,  $\Delta nmr2$ , or  $\Delta nmr3$  gene deletions. Expression is relative to gene expression in  $\Delta tps1$  strains grown on nitrate. (D) Bar charts showing that *G6PDH* gene expression is partially restored in a  $\Delta tps1$  mutant by introduction of  $\Delta nmr1$ ,  $\Delta nmr2$ , or  $\Delta nmr3$  gene deletions. (E) Bar charts showing gene expression of all nine genes in conidia (black bars) and appressoria (gray bars) of Guy11. Expression is relative to *TUB2* gene expression.

homologs of NMRA (Fig. 2). To test the role of these genes in Tps1 regulation, we deleted *NMR1*, *NMR2*, and *NMR3* in a  $\Delta tps1$  mutant background. Strikingly, targeted deletion of *NMR1*, *NMR2*, or *NMR3* all partially restored the ability of the  $\Delta tps1$  mutant to cause rice blast disease (Fig. 2). No significant differences in lesion number were observed between leaves inoculated with the wild-type and  $\Delta tps1$   $\Delta nmr$  suppressor strains (Fig. 2B), but relative lesion sizes were reduced in  $\Delta tps1$   $\Delta nmr1$  (mean relative lesion size =  $49 \pm 22$  pixel<sup>2</sup>,  $n = 12$  leaves),  $\Delta tps1$   $\Delta nmr2$  ( $30 \pm 19$  pixel<sup>2</sup>,  $n = 13$ ), and  $\Delta tps1$   $\Delta nmr3$  ( $19 \pm 7$  pixel<sup>2</sup>,  $n = 25$ ) compared with Guy11 ( $109 \pm 33$  pixel<sup>2</sup>,  $n = 12$ ). By contrast, deletion of the *NMR* genes singly, or together, in a wild-type strain did not significantly affect fungal pathogenicity (Fig. S4C). Deletion of the *Nmr* genes also resulted in partial derepression of nitrate reductase activity in wild-type and  $\Delta tps1$  mutant (see Fig. S4D and Fig. S4E) and partial remediation of sporulation of  $\Delta tps1$  strains on nitrate media (Fig. S4F). However, although partially remediated for virulence,  $\Delta tps1$   $\Delta nmr3$  suppressor strains were

not restored for sporulation on nitrate media (Fig. S4F). This observation indicates that the regulation of virulence-associated gene expression by Nmr proteins is both complex and independent of nitrate utilization.

We can therefore conclude that the three Nmr proteins act as corepressors to control virulence-associated gene expression downstream of Tps1. Low-stringency yeast two-hybrid studies did not detect a physical interaction between the Nmr proteins and Tps1 (Fig. 2C), suggesting that direct modulation of Nmr activity by Tps1 is unlikely. However, each Nmr protein possesses a putative NADP-binding Rossmann fold, which has been shown in *A. nidulans* NMRA to bind NADP (9) (Fig. S4B). We therefore investigated the role of NADP binding in Nmr function and Tps1 regulation in *M. oryzae*. Focusing on Nmr1, we found that introducing a native copy of *NMR1* into a  $\Delta tps1$   $\Delta nmr1$  double mutant complemented the  $\Delta nmr1$  gene deletion, as expected, and resulted in strains that, like  $\Delta tps1$  single mutants, were non-pathogenic (Fig. 2). However, we also constructed a mutant allele

of *NMR1* encoding a T13V substitution. This change has been shown in *A. nidulans* NmrA and *Dictyostelium discoideum* PadA to impair NADP binding (10, 11). Expression of the *NMR1*<sup>T13V</sup> allele in a  $\Delta tps1 \Delta nmr1$  double mutant did not restore the  $\Delta tps1$  non-pathogenic phenotype (Fig. 2D). We conclude that the non-pathogenic mutant phenotype of  $\Delta tps1$  mutants is associated with the role of Nmr1, Nmr2, and Nmr3 in repressing virulence-associated gene expression. We can also infer that the function of Nmr1 to carry out this role depends on its ability to bind NADP.

#### Tps1-Dependent Gene Expression Is Modulated by the Presence of Nmr Corepressors.

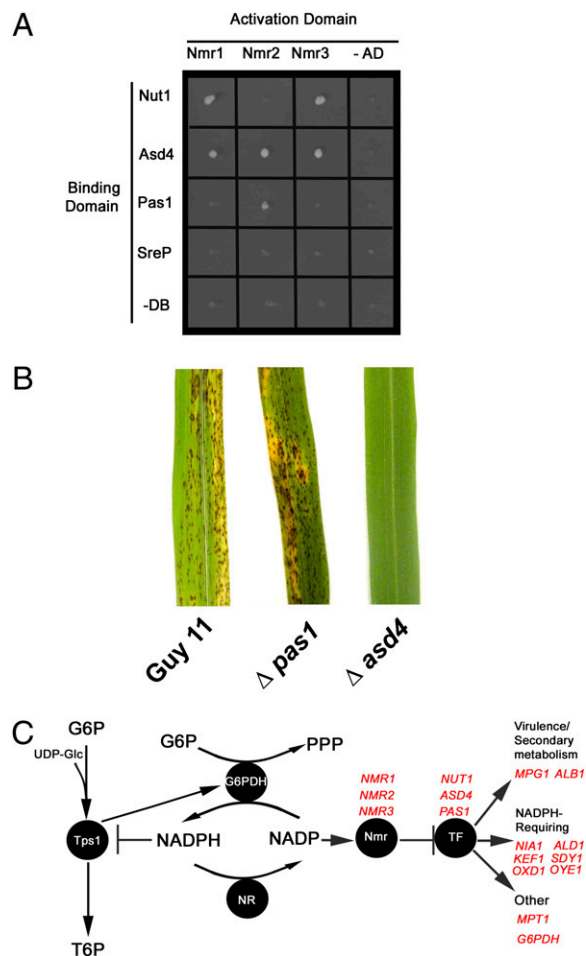
To understand how Tps1 might control virulence-associated gene expression during plant infection, we carried out comparative gene expression analysis of the wild-type,  $\Delta tps1$  mutant and the  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr2$ , and  $\Delta tps1 \Delta nmr3$  suppressor strains. During vegetative growth on nitrate compared with ammonium, we found that two known virulence genes *MPG1* and *ALB1* (12, 13), six genes encoding putative NADPH-dependent enzymes (*NIA1*, *ALD1*, *KEF1*, *SDY1*, *OXD1*, and *OYE1*), and a putative metalloprotease (*MPT1*) were expressed in a Tps1-dependent manner in response to nitrate (Fig. 3A and B, Table S1, and Fig. S3A). Expression of these genes was at least partially restored in the  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr2$ , or  $\Delta tps1 \Delta nmr3$  suppressor strains compared with the  $\Delta tps1$  mutant, suggesting they are controlled via NADP-dependent Nmr inhibition of transcription (Fig. 3C). In addition, we found that *G6PDH* gene expression was also elevated in the  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr2$ , and  $\Delta tps1 \Delta nmr3$  suppressor strains, compared with the  $\Delta tps1$  mutant, during growth on nitrate (Fig. 3D). *G6PDH* and a subset of the genes analyzed in Fig. 3C were elevated in expression during appressorium development, consistent with the operation of this regulatory mechanism during plant infection (Fig. S2B and Fig. 3E). Moreover, some genes, like *ALB1* and *G6PDH*, were derepressed in the  $\Delta tps1 \Delta nmr1$ ,  $\Delta tps1 \Delta nmr2$ , and  $\Delta tps1 \Delta nmr3$  suppressor strains compared with  $\Delta tps1$ , whereas other genes are derepressed in some suppressor strains but not others. The expression of *ALD1*, for instance, was only affected in the  $\Delta tps1 \Delta nmr2$  mutant. This result suggests that Nmr corepressors may work either individually or cooperatively to regulate gene expression, which may result from their regulation of additional GATA factors. Consistent with this idea, a subset of the genes was expressed in a *NUT1*-dependent manner (Fig. 3A), whereas others were expressed independently of *NUT1* (Fig. 3B), indicating that the Nmr proteins may regulate other GATA family transcription factors. To test this idea, we carried out high stringency yeast two-hybrid experiments to determine whether Nmr proteins can interact with other GATA family transcription factors including *Asd4*, a putative homolog of a *Neurospora crassa* morphogenetic regulator (14); *Pas1*, a GATA factor containing a pas-fold domain involved in redox (15) and/or light sensing, which is a putative homolog of white collar-2 from *N. crassa* (8), and *Sre1*, a GATA factor required for siderophore metabolism (16). We found evidence that the Nmr1, Nmr2, and Nmr3 can physically interact with *Asd4*, whereas Nmr2 interacts with *Pas1*, as shown in Fig. 4A. No interaction was observed with *Sre1*. Based on their potential interaction with Nmr corepressors, we decided to investigate the function of *ASD4* and *PAS1*. We carried out targeted gene replacements and found that  $\Delta asd4$  mutants displayed reduced hyphal growth (Fig. S5A) and sporulation. By contrast,  $\Delta pas1$  mutants showed enhanced conidiation (Fig. S5B). Significantly,  $\Delta asd4$  mutants were unable to cause rice blast disease (Fig. 4B), which was associated with their inability to form appressoria (Fig. S5C).

We conclude that Tps1 affects virulence-associated gene expression via its modulation of a set of NADP-dependent transcriptional repressors, which target GATA factors implicated in both fungal development and pathogenicity.

#### Discussion

In this study we set out to understand the role of trehalose-6-phosphate synthase as a key regulator in the establishment of rice blast disease (4). When considered together, our results are

consistent with a role for Tps1 as part of an NADPH-dependent genetic switch in *M. oryzae*, which is essential for virulence and integrates the intracellular monitoring of nutritional and redox status with control of fungal gene expression. At the heart of this regulatory mechanism is the nonconsuming interconversion of NADPH and NADP, which bind to Tps1 and the Nmr transcriptional corepressor proteins, respectively, as shown in the



**Fig. 4.** A unique NADP(H)-dependent genetic switch is essential for fungal virulence in the rice blast fungus. (A) The Nmr repressors interact with three GATA transcription factors in *M. oryzae*. High stringency yeast two-hybrid analysis of physical interactions between Nut1, Asd4, Pas1, and Sre1 with Nmr1, Nmr2, and Nmr3. -BD are strains carrying an empty *GAL4* DNA-binding domain vector. -AD are strains carrying an empty *GAL4* activation domain vector. High stringency selection was on media lacking adenine, histidine, tryptophan, and leucine. (B) Role of the GATA-factors Pas1 and Asd4 in pathogenicity.  $\Delta pas1$  strains are fully pathogenic, whereas  $\Delta asd4$  strains are unable to infect host tissue. Rice seedlings were inoculated with  $1 \times 10^5$  spores  $\text{mL}^{-1}$ . (C) A model for the action of Tps1 in virulence of *M. oryzae*. In response to G6P sensing, Tps1 activates G6PDH to produce NADPH from G6P and NADP in the oxidative pentose phosphate pathway. NADPH can bind to Tps1 to maintain a balance between G6P consumption and NADPH production. The conversion of NADP to NADPH inactivates the Nmr inhibitor proteins and results in the derepression of genes encoding at least two known virulence factors, at least six NADPH-dependent proteins, and G6PDH. NADPH is subsequently consumed by NADPH-dependent proteins, such as NR. Under starvation conditions, such as found during appressorium formation on the surface of the leaf, the homeostatic balance between G6P consumption and NADPH production is not maintained, NADP levels are elevated, and the Nmr inhibitor proteins become activated leading to repression of the genes required for virulence. Proteins are indicated as black circles. TF, transcription factor(s). The corresponding genes are shown in red. UDP-glc, UDP-glucose.

model presented in Fig. 4C. Nmr proteins interact with at least three GATA family transcription factors to regulate expression of genes necessary for virulence, in addition to a number of genes encoding putative NADPH-dependent enzymes. The switch has two distinct modes, an “on” and a default “off” status, depending on the nutrient condition of the fungal cell. When G6P is available and the NADPH/NADP ratio is high, the switch is on and dynamically links G6P availability to gene expression with (i) NADPH depletion by enzymes such as NR being balanced by Tps1-dependent NADPH production in response to G6P sensing, (ii) NADPH production and G6P consumption maintained in equilibrium by competition for Tps1-binding, and (iii) the expression of virulence-associated genes and genes encoding NADPH-requiring enzymes, such as NR, induced only under NADPH replete conditions due to inactivation of the Nmr corepressors resulting from low levels of NADP. Trehalose would also be produced as a storage compound under these conditions. However, when G6P levels and the NADPH/NADP ratio are low, the system cannot maintain dynamic equilibrium. Tps1 becomes inactive because it cannot bind either NADPH or its native substrates. G6PDH activity and gene expression is reduced, and the default off position of the pathway is therefore NADP activation of the Nmr corepressor proteins. Further feedback to the system is provided by the likely control of *G6PDH* gene expression exerted by the GATA factor/Nmr transcriptional regulators. The default off position is also seen in  $\Delta$ *tps1* mutants, because the Nmr proteins are constitutively active, repressing virulence-associated gene expression.

The involvement of dinucleotide cofactors in transcriptional regulation has not been widely reported, but it was recently predicted from a study in *Saccharomyces cerevisiae* where NAD(P) was shown to have the potential to act as a cofactor in the interaction between Gal80p and Gal4p during induction of galactose metabolism (17). In mammals, meanwhile, a family of histone deacetylases, the sirtuins, are involved in regulation of diverse cellular processes and appear to depend on NAD for their activity (reviewed in ref. 18), ensuring a tight link between sirtuin activity and the underlying metabolic status of the cell. However, the results presented here suggest that *M. oryzae* Tps1 acts as part of an NADP(H) homeostatic model, where the expression of genes encoding NADP(H)-requiring enzymes directly influences NADPH production and corresponding gene expression. We speculate that this adaptability to fluctuating nutritional environments may be key to the success of plant pathogens such as *M. oryzae*, which must quickly adapt from nutrient-free leaf surfaces to intracellular propagation within living plant tissue. Once in the plant, the fungus must, for instance, undergo rapid genetic reprogramming to elaborate specialized invasive hyphae from the

thin penetration hyphae and evade or suppress the host defense response to establish disease (19). The Tps1 genetic switch described in this study may be responsible for postpenetration genetic reprogramming because  $\Delta$ *tps1* strains form appressoria, but these cells are unable to develop into invasive hyphae to establish disease (3). Significantly, T6P synthases have been shown to have important roles in developmental biology in an increasing range of organisms, being necessary for plant embryonic development, flowering, and other morphogenetic processes (20). The genetic mechanism defined in this study may therefore be a more widespread mechanism for integrating metabolic and genetic regulation during eukaryotic development.

## Materials and Methods

**Fungal Strains, Growth Conditions, and DNA Analysis.** *M. oryzae* strains used in this study are derived from Guy11 (1, 3, 4). Standard procedures of *M. oryzae* growth, maintenance, appressorium formation, and transformation were performed, as described (12). Rice plant infections were performed as described (12). DNA and RNA extractions were as described in ref. 4. Gel electrophoresis, restriction enzyme digestions, DNA ligations, and PCR were performed by using standard procedures (21).

**Transcript and Protein Analysis.** cDNA synthesis and qRT-PCR was performed as described in *SI Materials and Methods*. Yeast two-hybrid analysis was performed by using the Matchmaker GAL4 Two-Hybrid System 3 kit from Clontech. Purified Tps1 recombinant protein was obtained by using the *Arcticexpress* (Stratagene) gene expression system. Whole-cell protein extraction and nitrate reductase assays were performed as described (4). Activities were determined on a Pharmacia Biotech spectrophotometer in triplicate. All assay components were purchased from Sigma, except NADPH (Calbiochem). NADPH fluorescence was recorded on a 200 series microplate reader (Tecan Group). Protein modeling of *M. oryzae* Tps1 was performed as described in *SI Materials and Methods*. Statistics were performed by using Student's *t* test. Protein sequence alignments were performed by using ClustalW and illustrated using BoxShade.

**Generation of *M. oryzae* Gene Deletion Mutants.** Targeted gene replacement of *M. oryzae* genes were performed based on the split marker strategy (22), as described in *SI Materials and Methods* and Fig. S4. *M. oryzae* gene sequence information was acquired from [www.broad.mit.edu/annotation/genome/magnaporthe\\_grisea/MultiHome.html](http://www.broad.mit.edu/annotation/genome/magnaporthe_grisea/MultiHome.html). For full experimental protocols, see *SI Materials and Methods*.

**ACKNOWLEDGMENTS.** This work was supported by a grant to N.J.T. from the Biotechnology and Biological Sciences Research Council. C.F.Q. was supported by a graduate student assistantship from the Department of Plant Pathology, University of Nebraska, Lincoln, NE.

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