**Phosphopantetheinyl transferase (Ppt)-mediated biosynthesis of lysine, but not siderophore or DHN melanin, is required for virulence of *Zymoseptoria tritici* on wheat**

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**Abstract**

*Zymoseptoria tritici* is the causal agent of Septoria tritici blotch (STB) disease of wheat. *Z. tritici* is an apoplastic fungal leaf pathogen which does not penetrate plant cells at any stage of infection and has a long initial period of symptomless colonisation. During this phase, it is unclear to what extent the fungus can access host plant nutrients or communicate with plant cells. Several important primary and secondary metabolite pathways in fungi are regulated by the post-translational activator phosphopantetheinyl transferase (Ppt) which provides an essential co-factor for lysine biosynthesis and the activities of non-ribosomal peptide synthases (NRPS) and polyketide synthases (PKS). To investigate the relative importance of lysine biosynthesis, NRPS-based siderophore production, and PKS-based DHN melanin biosynthesis, we generated deletion mutants of *ZtPpt*. The ∆*ZtPpt* strains were auxotrophic for lysine and iron, non-melanised and non-pathogenic on wheat. Deletion of the three target genes likely affected by ZtPpt loss of function (*Aar*- lysine; *Nrps1*-siderophore and *Pks1*-melanin), highlighted that lysine auxotrophy was the main contributing factor for loss of virulence, with no reduction caused by loss of siderophore production or melanisation. This reveals Ppt, and the lysine biosynthesis pathway, as potential targets for fungicides effective against *Z. tritici*.

Key words: Dothideomycete, Secondary metabolism, *StuA,* transcriptional regulation, phosphopantetheinyl transferase, melanisation, *Mycosphaerella graminicola*, dimorphic fungi, fungicide resistance, host-specific toxin.

**Introduction**

The ascomycete fungus *Zymoseptoria tritici* (Desm.) (Quaedvlieg & Crous)1 causes Septoria tritici blotch (STB) disease of wheat in temperate growing regions worldwide2,3. This disease is widely considered, both in industry and academia, to be the most important wheat disease in Europe3. It is estimated that approximately 70 % of the $1.7 billion a year European fungicide market for wheat is sold for the control of STB4. Extensive use of fungicides has led to numerous cases of resistance in *Z. tritici*, and new chemistries are needed for future disease control 5,6. Knowledge of essential-for-life and important virulence genes in this fungusmay aid the development of these new chemistries7.

 *Z. tritici* is often considered to be a hemibiotrophic fungus, as it undergoes two distinct phases of plant colonisation – an initial symptomless infection lasting at least one week, which is subsequently followed by a necrotrophic phase associated with the formation of leaf lesions. During the symptomless phase, *Z. tritici* hyphae enter leaves through stomata and colonize mesophyll tissue where they grow slowly in the intercellular space, the “apoplast”, without producing haustoria or any other form of plant cell penetrating structures. The onset of leaf necrosis is then rapid and begins with the appearance of chlorotic lesions. These eventually coalesce into necrotic blotches bearing pycnidia, which are asexual fructifications containing rain splash-dispersed pycnidiospores8. This entire infection cycle occurs without fungal penetration of host cells. For this reason, *Z. tritici* is also referred to as an ‘apoplastic’ pathogen, which delineates it from many other plant pathogenic fungi that do physically penetrate plant cells at some point during infection. However, the full repertoire of mechanisms which enable the initial symptomless colonization and the subsequent activation of leaf necrosis by *Z. tritici* have not been fully elucidated.

 Several plant pathogenic fungi produce diffusible secondary metabolites (SMs) to induce host tissue necrosis, which supports colonisation. Many of these SMs are produced by non-ribosomal peptide synthase (NRPS) or polyketide synthase (PKS) enzymes frequently residing in genome clusters with other tailoring genes and transporters. Important examples include the victorin toxin of *Cochliobolus victoriae* and the T-toxin of *Cochliobolus heterostrophus*. Victorin is a non-ribosomal peptide (NRP) that is produced by the NRPS enzyme TOX39. This toxin causes programmed cell death (PCD) in *Arabidopsis thaliana* by interacting with the nucleotide binding site leucine-rich repeat (NB-LRR) receptor protein, LOV110. T-toxin is a polyketide (PK) produced by PKS enzymes encoded at two distinct loci, Tox1A and Tox1B11. It interacts with mitochondria in Texas male sterile cytoplasm corn to cause necrosis12. Both T-toxin and victorin are essential for virulence in their respective pathogens13.

 As bioactive molecules, SMs are often involved in a variety of other processes such as microbial competition, and melanin and siderophores are two key examples of metabolites frequently found to be produced by most fungi 14–16. In the citrus pathogen *Alternaria alternata,* disruption of the NRPS siderophore synthetase-encoding gene *NPS6* leads to hypersensitivity to low iron availability and reactive oxygen species (ROS), which leads to reduced virulence14. This reduction in virulence is likely due to the iron cofactor requirement of enzymes that quench host-derived ROS. In *Z. tritici* several studies have predicted the role of a specific PKS (Pks1) in melanin biosynthesis but this gene has, to date, not been functionally characterised17,18.

 Despite diverse biological roles, all NRPSs and PKSs share a single point of post-translational activation. To function, they must be 4’-phosphopantetheinylated at conserved serine residues within the acyl carrier protein (ACP) domain. The enzyme that carries out this process is 4’-phosphopantetheinyl transferase (Ppt), which was first described for filamentous fungi in *A. nidulans*19,20. In addition to its role in the biosynthesis of SMs and siderophores, Ppt is essential for 4’-phosphopantetheinylation of alpha aminoadipate reductase (Aar), an enzyme crucial for lysine biosynthesis. Hence the *Saccharomyces cerevisiae* Ppt homologue was first described as LYS5, for which cognate mutants were auxotrophic for lysine21.

 Following its discovery in model fungi, *Ppt* homologues were characterised in several plant pathogens with the aim of identifying links to pathogenicity via production of SMs. These include *Colletotrichum graminicola, Magnaporthe oryzae, Fusarium fujikuroi* and *Cochliobolus sativus*22–24. In all these species, *∆Ppt* deletion strains were shown to be auxotrophic for lysine and unable to melanise or produce siderophores. In addition, a Ppt homologue with similar functions has also been characterised in the endophytic fungus *Trichoderma virens*25. The auxotrophy for lysine of *∆Ppt* deletion strains has led to its consideration as a potential antifungal target. For example, in the human pathogens *Candida albicans* and *Aspergillus fumigatus*, high throughput screens have been developed with the aim of identifying inhibitors of this enzyme26,27.

 Though some components of the interaction between *Z. tritici* and *Triticum* *aestivum* have been elucidated28–30, the biological role of SMs in *Z. tritici* and key genes implicated in the regulation of SMs have not been functionally characterised. To gain an insight into the functions of *Z. tritici* SMs and to ascertain the importance of lysine biosynthesis, siderophore biosynthesis and melanisation for the apoplastic lifestyle and virulence, the *Z. tritici* Ppt homologue *ZtPpt* was functionally characterised by gene deletion. In addition, we generated and characterised deletion mutants for key downstream targets of Ppt that are directly responsible for melanisation (*Pks1*), siderophore (*Nrps1*) and lysine (*Aar*) biosynthesis. Further gene deletion strains were generated and characterised for the recently described *Z. tritici* transcriptional regulator *StuA*31, and two additional Polyketide synthases (PKS7 and PKS8) which previous studies had shown to be highly expressed during leaf infection32,33.

**Results**

**Identification of *Zymoseptoria tritici* homologues of Ppt and genes implicated in regulating biosynthesis of lysine, siderophore and DHN-melanin**

A BLASTp analysis was conducted to determine whether *Z. tritici* contains clear homologues of Ppt and associated genes previously characterised in *C. sativus*22. This showed that *Z. tritici* possesses a single Ppt homologue, a single Pks1 homologue, a single Aar homologue and two homologues for NPS6 (Supplementary Table 1). We performed additional Blastp analyses using characterised sequences from the *A. fumigatus* siderophore biosynthesis pathway to more accurately ascribe the best predicted functional homologue of the *Aspergillus fumigatus* proteinNPS6*,* which revealed the protein ZtNrps1 as the best candidate (Supplementary Table 1). The protein encoded by *ZtNrps1* was homologous to both the *sidC* and *sidD* protein families but not to other siderophore biosynthetic proteins that are not generated by NRPSs. ZtNrps1 was in fact most similar to sidD, which is involved in the biosynthesis of an extracellular fusarin C siderophore (Supplementary Table 2).

***Z. tritici* Ppt deletion mutants are lysine auxotrophs, hypersensitive to iron depletion and ROS, and are deficient in melanisation**

*ZtPpt*, *ZtPks1, ZtNrps1* and *ZtAar* gene deletion strains were all generated (Supplementary Figure 1) and firstly assessed for their responses to different stresses *in vitro*. This revealed that Δ*ZtPpt* strainswere hypersensitive to iron depletion, as were the Δ*ZtNrps1* mutants(Figure 1A, 1B). This was evident in the observation that neither of these strains grew in the presence of the iron chelating agent BPS but were rescued with further addition of an exogenous siderophore compound, desferriferrichrome (DFF). Δ*ZtPpt* strainswere also auxotrophic for lysine biosynthesis, as was also observed for the AAR mutant Δ*ZtAar* (Figure 1A and 1C). Finally, Δ*ZtPpt* strains were unable to melanise when exposed to UV light for six days or longer, a phenotype also observed for Δ*ZtPks1* (Figure 1A, 1D). This same phenotype was also recently observed for *ZtStuA* mutants31 (Supplementary Figure 2A and 2B) highlighting that both StuA and Ppt functions are required for melanisation in this fungus. This was further supported by our additional data on ZtStuA, which demonstrated it to be a nuclear localised protein (Supplementary Figure 2C), and to positively influence the expression of several key genes for the DHN- melanin biosynthesis pathway, including the *Pks1* gene itself (Supplementary Figure 2B and Supplementary table 3).

***Zymoseptoria tritici* filamentous growth on nutrient limiting agar requires lysine biosynthesis but not Nrps1-derived siderophores or melanin**

To evaluate whetherΔ*ZtPpt* strains were affected in filamentous growth *in vitro*, mutant strains were spot inoculated onto water agar, a nutrient depleted substrate which stimulates the formation of hyphal filaments from *Z. tritici* spores34–36. This showed that, in the absence of exogenous lysine, Δ*ZtPpt* and Δ*ZtAar* could not extend hyphal filaments (Figure 2 and Supplementary Figure 3). When lysine was added, filamentous growth was rescued in these strains but not to WT levels (Figure 2 and Supplementary Figure 3). Both the Δ*ZtNrps1* and Δ*ZtPks1* strains grew to at least WT levels whether supplemented with lysine or not. One of each of the Δ*ZtNrps1* and Δ*ZtPks1* strains grew significantly faster than the WT without supplements, though this was not consistent across both strains assayed (Figure 2).

**Lysine biosynthesis is a key requirement for *Z. tritici* infection of wheat but the Nrps1-derived siderophore and melanin are not**

We next assayed the virulence of Δ*ZtPpt* and the Δ*ZtPks1,* Δ*ZtNrps1* andΔ*ZtAar* downstream gene deletion mutants, on a susceptible wheat cultivar. This showed that Δ*ZtPpt* was completely avirulent, both with and without lysine supplementation in the starting inoculum, on wheat seedlings (Figure 3A and 3C; Supplementary Fig 4). The same phenotype was also observed for the Δ*ZtAar* strain. In contrast, the Δ*ZtNrps1* strains were not affected in virulence, nor were the Δ*ZtPks1* strains (Figure 3A, 3C). However, Δ*ZtPks1* generated non-melanised pycnidia at the end of the infection period (Figure 3A, 3B). *ΔZtStuA* deletion mutants were recently shown to be attenuated in virulence on wheat31 and our data provided further support for this, which manifested itself in a delayed onset of leaf necrosis and lesions lacking asexual spores (Supplementary Figure 2D).

**Two additional *Zymoseptoria tritici PKS* genes strongly up-regulated during leaf infection are not required for virulence.**

To assess the importance of two previously identified genes, *ZtPks7* and *ZtPks8,* whose gene clusters were transcriptionally up-regulated during infection37, deletion strains were generated and inoculated onto a susceptible wheat cultivar to test for altered virulence. For the single *∆ZtPks7* strain and the three ∆*ZtPks8* strains tested, no effects on virulence were observed (Figure 4). After the full course of infection at 21 days, necrotic symptoms indistinguishable from the WT were observed in all cases (Figure 4A). The number of spores retrieved from infected leaves for each of the mutant strains was also not significantly different to that for the WT strain (Figure 4B-C).

**Discussion**

In this study, we analysed the relative importance of particular primary and secondary metabolites for virulence and other processes in the apoplastic wheat pathogen *Z. tritici*. We functionally characterised several *Z. tritici* genes, some of which were homologues of those known in other fungi to regulate lysine biosynthesis, melanisation, and siderophore biosynthesis,amongst other processes. A summary overview of the deletion strains generated, the downstream processes they regulate, and their contributions (or not) to virulence is presented in Figure 5. The schematic also incorporates data from the recently characterised StuA mutants of *Z. tritici*31 as well as our own novel insights presented here which demonstrated StuA to have a direct impact on the expression of *Pks1* and other melanin pathway genes.

 In agreement with prior studies31 *∆ZtStuA* mutants were defective in melanisation and we have now shown that the *Z. tritici* gene *ZtPks1,* was not expressed in this mutant, neither were other genes predicted to be in the melanin-biosynthesis pathway. *∆ZtPks1* deletion mutants themselves did not exhibit melanisation. These data together provide clear evidence that ZtPks1is involved in melanin biosynthesis and ZtStuA can transcriptionally regulate it. That *ZtPks1* was involved in melanin biosynthesis has been suggested before by Cairns et al. (2017) and Lendenmann et al. (2015)17,18 but here we provide the experimental validation of this prediction. Interestingly, a recent QTL-based analysis of natural variation in the rate of melanisation by different *Z. tritici* strains has also revealed a second transcriptional regulator which positively influences *Pks1* expression, *Zmr1*38. This suggests that either StuA and Zmr1 exert their effects on *Pks1* and melanisation separately, or they may form part of, an as yet undescribed signalling pathway controlling *Pks1* expression. These two possibilities require further testing.

Alike the *∆ZtStuA* and *∆ZtPks1* mutants, *∆ZtPpt* mutants were also unable to melanise suggesting that the protein does indeed function to provide the essential co-factor for PKS and NRPS enzymes, including Pks1. This would indicate that ZtPpt functions similarly to those in other fungi to phosphopantetheinylate PKS and NRPS enzymes, including ZtPks1. ∆Z*tPpt* mutant strains were also hypersensitive to iron depletion and ROS, and auxotrophic for lysine. This further supports the idea that ZtPpt is a functional homologue of the *Ppt* genes characterised in other fungi.

*ZtNrps1* mutant strains were hypersensitive to iron depletion, which would indicate that this gene encodes an enzyme that synthesises an important siderophore in a non-redundant manner under the conditions tested. Growth was restored when extracellular siderophores were added to the medium. In the model fungus *A. fumigatus* two major *NRPS* genes involved in siderophore biosynthesis have been characterised. These are *sidC*, which is required for the biosynthesis of intracellular ferricrocin siderophores, and *sidD*, which is required for biosynthesis of extracellular fusarin C siderophores39. Intracellular siderophores are required for iron storage and transport, whereas extracellular siderophores are required for chelation and uptake of unavailable iron. *ZtNrps1* was more similar to sidD than sidC, supporting the hypothesis that it produces an extracellular siderophore. However, *ZtNrps1* was not necessary for wheat infection by *Z. tritici*. This is in contrast to previous reports on other fungi which have shown homologues of sidD (ZtNrps1) to be important virulence factors40. A notable exception to this was *Ustilago maydis*, where targeted disruption of biosynthesis of all siderophores in this fungus did not reduce virulence41. Since *U. maydis* is a biotrophic fungus and *Z. tritici* exhibits an extensive biotrophic / symptomless phase, it may be that the fungus has sufficient easily accessible iron, which overrides the need for extracellular siderophore production during leaf infection. Alternatively, *Z. tritici* may be able to use stored iron during filamentous growth. This is supported by the ability of *∆ZtNrps1* mutant strains to produce WT levels of hyphal filaments on iron-free water agar.

In contrast to the *∆ZtNrps1 and ∆ZtPks1* strains*,* *∆ZtPpt* and ∆*ZtAar* mutants were unable to grow hyphal filaments *in vitro* or undergo budding growth on minimal medium without lysine supplementation. This indicates that both filamentous and budding growth in *Z. tritici* requires a fully functioning lysine biosynthetic pathway. This is consistent with the lysine auxotrophy observed in Ppt mutant strains in various filamentous fungi studied to date22,23,25. The virulence data on wheat leaves demonstrated that both *∆ZtNrps1 and ∆ZtPks1* exhibited WT virulence, but both *∆ZtPpt* and ∆*ZtAar* were strongly reduced in virulence. Together this suggests that lysine availability (or lack of) forms a major barrier to infection by the latter two mutant strains, most probably as a result of restricting hyphal growth, and it is the loss of *ZtAar* function which is largely responsible for the loss of virulence in *∆ZtPpt.* Hence, both ZtPpt and ZtAar would appear to be good targets for putative fungicides for the future control of STB.

 In a recent study37 it was shown that *Z. tritici* exhibits differential expression of several other large putative SM clusters during infection. Several of these gene clusters contained PKSs highly upregulated coincident with the first appearance of necrotic lesions on wheat leaves. However, functional studies reported here have now shown that two of these PKSs are not essential for infection of the susceptible host cultivar tested, although we cannot rule out possible interactions with other cultivars. Ablation of individual PKS enzymes often has no discernible effect on virulence in plant pathogens. For example, in *Fusarium graminearum*, 15 PKS genes were disrupted in a single study and none affected virulence42. Hence, the functional roles of individual SM clusters in *Z. tritici* remain unknown. Besides their roles in host necrosis as toxins, SMs may be involved in, for example, microbial competition. In *T. reesei*, mutants for a specific PKS were less able to inhibit the growth of competing microbes, including the plant pathogens *Alterneria alternata, Sclerotinia sclerotiorum, Botrytis cinerea*,and *Rhizoctonia solani*43*.* It is possible that *Z. tritici* specifically produces these energetically costly compounds, at a time when necrosis appears, to compete favourably with other microbes that may wish to utilise the dead tissue. However, this, and the roles of several other predicted secondary metabolite clusters in the biology of *Z. trtici,* require further testing.

**Methods**

**Strains, media and growth conditions**

The fully sequenced *Z. tritici* reference strain IPO3236 and its derivative (Δ*ZtKu70*), in which the *Ku70* gene has been disrupted to reduce non-homologous end joining (NHEJ)44, were employed as wild type (WT) and recipient strains for gene deletion and gene disruption. Both strains are highly virulent on the susceptible wheat cultivars Taichung 29 and Riband. The WT and all generated strains were stored at -80 *°*C and re-cultured on potato dextrose agar (PDA) or yeast peptone dextrose (YPD) agar (Sigma-Aldrich Chemie, Steinheim, Germany) at 18 °C. Yeast-like spores were produced on V8 juice medium (Campbell Foods, Puurs, Belgium) or in yeast glucose broth (YGB) medium (yeast extract 10 g/L, glucose 20 g/L) placed in an orbital shaker (Innova 4430; New Brunswick Scientific, Nijmegen, The Netherlands) at 18 °C. To induce mycelial growth, all *Z. tritici* strains were grown under the same conditions but at 24 °C. To assess lysine auxotrophy, 100 mM Lysine was added to the minimal medium Czapek-Dox (CD) agar. To induce filamentous growth *in vitro* strains were grown on 10 % sterile distilled water (SDW) agar.

**Bioinformatic analyses**

Phylogenetic analysis of ZtStuA with homologues from other fungal pathogens was performed using the CLC genomics workbench package (Aarhus, Denmark). All StuA fungal proteins were retrieved from public databases and aligned with a gap opening cost and gap extension penalty of 10 and 1, respectively. The phylogenetic tree was constructed based on the unweighted pair group method with arithmetic average (UPGMA) algorithm, and support for the tree was assessed with 1000 bootstraps.

Additionally, the genome sequence of *Z. tritici* IPO323 (Goodwin et al., 2011) was queried in a BLAST search using the characterised sequences of PPT, NPS6 (a siderophore biosynthesetic enzyme), PKS1 (the fungal melanin biosynthetic enzyme) and AAR (the lysine biosynthesis enzyme) from *C. sativus* (Leng and Zhong, 2012). This was done using the BLASTp program over the NCBI BLAST server (<https://blast.ncbi.nlm.nih.gov/Blast.cgi?PAGE=Proteins>) with default settings. The PKS sequences *ZtPks7* and *ZtPks8* first identified as having a potential role in infection37 were also identified using the *Z. tritici* IPO323 genome sequence.

A further BLASTp analysis was conducted against the *Z. tritici* annotations using the *A. fumigatus* accessions XP\_753088.1, XP\_746662.1, XP\_748660.1 and XP\_748685.1, which correspond to the amino acid predictions for genes *sidC*, *sidD, sidF* and *sigG*, respectively. The former, *sidC* and *sidD,* encode NRPSs that produce intracellular ferricrocin and extracellular fusarin C siderophores, respectively. The latter, *sidF* and *sidG*, encode non-NRPS siderphore synthesis-associated enzymes39.

**Generation of gene replacement, complementation and GFP fusion constructs**

To generate the *ZtStuA* deletion construct, pZtStuAKO, the USER friendly cloning method was used with minor modifications as described previously45,46. Briefly, ZtStuA-F3 and R3, as well as ZtStuA-F4 and R4 primer pairs were used to amplify about 2,000 bp upstream and downstream of ZtStuA using PfuTurbo® Cx Hotstart DNA polymerase (Stratagene, Cedar Creek, TX, US). In parallel, the pRF-HU2 vector possessing the *hygromycin phosphotransferase* (*hph*) gene as a selection marker was digested with two restriction enzymes, PacI and a nicking enzyme Nt.BbvCI, to generate a compatible overhang with the PCR amplicons. Subsequently, the PCR amplicons and the digested vector were mixed and treated with the USER enzyme mix (New England Biolabs, Ipswich, USA) and incubated at 37 °C for 30 min followed by 25 °C for 30 min. The resulting reaction was directly transformed into *Escherichia coli* strain DH5α and subsequently cultured on kanamycin-selective medium.

 To identify bacterial colonies carrying the construct with the insertions in the expected positions, colony PCR was conducted using User-F and User-R primers (located in the middle of the *hph* gene) in combination with ZtStuA-F1 and ZtStuA-R1, respectively.

To produce the *ZtStuA* complementation construct and ZtStuA fused to GFP, the vectors pCZtStuA.com and pCZtStuA.GFP were generated by *in vivo* recombination in the yeast *Saccharomyces cerevisiae* DS94 (MATa, *ura3-52, trp1-1, leu2-3, his3-111,* and *lys2-801*)47 following published procedures48,49. The vector pCZtStuA.com contains the *ZtStuA* ORF under the control of the native promoter (1,025 bp) and terminator sequences (495 bp) for random ectopic integration into the genome of *Z. tritici* by using BASTA as the selection agent, while the vector pCZtStuA.GFP contains ZtStuA.GFP fused to the full-length *ZtStuA* under the control of the constitutive *ZtTub2* promoter and terminator sequences for targeted integration into the *sdi1* locus of *Z. tritici* by using carboxin as the selection agent. A 9,760 bp fragment of pCGEN-YR (digested with *Xba*I and *Zra*I48), 1410 bp fragment covering the BASTA resistance cassette *bar* (amplified with MG-Sep-106 and MG-Sep-107), 3146 bp fragment covering the 1025 bp of *ZtStuA* promoter, 1626 bp pf *ZtStuA* gene and 495 bp of *ZtStuA* terminator (amplified with MG-Sep-113 and MG-Sep-116) were recombined in yeast *S. cerevisiae* to obtain the vector pCZtStuA.com. A 16,597 bp fragment of pCZtGFPSpa2 (digested with *Psh*AI50), 1626 bp full-length *ZtStuA* gene (amplified with SK-Sep-346 and SK-Sep-347) were recombined in yeast *S. cerevisiae* to obtain the vector pCZtGFPStuA.

 To produce the disruption constructs for the Ppt-associated genes *ZtNrps1* and *ZtPks1*, and the PKS genes *ZtPks7* and *ZtPks8*, primers with added 5’ restriction sites and two 5’ adenines were designed for the amplification of the selected flanking sequences using Geneious version 8.1. Flanking sequences were approximately 1 Kb either side of these genes. Flanking regions were amplified in PCR reactions from genomic DNA of isolate IPO323 using RedTaq mastermix according to the manufacturer’s protocol. Thermocycler settings were 95 ̊C 2 min, then 30 cycles of 95 ̊C 30 sec, 60 ̊C 30 sec, 72 ̊C 2 min, and 10 ̊C hold. Flanking sequences were cloned into pCHYG51 either side of *hph* under the *trpC* promoter. For all strains generated in this study, primers and added restriction sites are in Supplementary Table 4.

 The ∆*ZtAar* strain was generated in Ali et al. (2015)52. Briefly, pCambia 0380 was digested with SacII, the 2-micron +ura3 region was amplified by PCR from pYES2 using 2µmURA3 FP and 2µmURA3 RP was used to repair the plasmid by in-yeast recombination. The left and right flanks for *ZtAar* were amplified from genomic DNA by PCR and recombined into BamH1-digested pCambia0380YA along with the hygromycin resistance cassette. Plasmids were purified from yeast using the Zymoprep ITM yeast plasmid extraction kit and rescued into *E. coli* then construction confirmed by restriction digest, PCR and sequencing across recombination sites.

**Fungal transformation**

The gene deletion construct as well as the replacement construct were first cloned into *A. tumefaciens* strain AGL1 via electroporation. *Agrobacterium* mediated transformation was then carried out to delete *ZtStuA* in the Δ*ZtKu70* strain as previously described51. For the complementation assay, the same procedure was applied but the putatively complemented strains were selected on plates with 50 µg mL-1 BASTA. All disrupted PKS and Ppt-associated mutant strains except Δ*ZtAar* were also generated in the Δ*ZtKu70* background44. Δ*ZtAar* was disrupted in a wild type IPO323 background52. For the Ppt and Aar mutants, 100 µM Lysine was added to the selection plates to ensure survival of transformed cells, which were expected to be lysine auxotrophs. Emerging colonies were then subjected to two rounds of selection on hygromycin amended YPD agar plates. Genomic DNA of stable transformants was extracted and used in PCR screens to confirm homologous recombination in the correct positions.

**Epi-fluorescence microscopy**

Fluorescence microscopy was performed as previously described53. In brief, IPO323\_CZtGFPStuA cells were inoculated in YG media and grown at either 18°C with 200 rpm to induce the yeast-like cell growth or 24°C with 100 rpm to induce the hyphal growth for 24 h and placed onto a 2% agar cushion for direct observation using a motorized inverted microscope (IX81; Olympus, Hamburg, Germany), equipped with a PlanApo 100\_/1.45 Oil TIRF objective (Olympus, Hamburg, Germany). Fluorescent tags and dyes were exited using a VS-LMS4 Laser Merge System with solid-state lasers (488 nm/50mW or 75mW and 561 nm/50mW or 75 mW; VisitronSystems, Puchheim, Germany) and single images or z-Stacks, using an objective piezo (Piezosystem Jena GmbH, Jena, Germany), over 6 µm depth with a z resolution of 0.2 µm were captured with150 ms exposure. In addition a DIC image was taken for each cell using a CoolSNAP HQ2 camera (Photometrics/Roper Scientific,Tucson, USA). Overlays of the fluorescent and DIC images were generated using MetaMorph (Molecular Devices, Wokingham, UK). All parts of the system were under the control of the software package MetaMorph (MolecularDevices, Wokingham, UK). The localisation of ZtGFPStuA in the nucleus was confirmed by counterstaining the nucleus with DAPI (Sigma-Aldrich Company Ltd. Dorset, England). To this end, cells of strain IPO323\_CZtGFPStuA were grown for 24 h in YG medium at 18 °C with 200 rpm. DAPI was added the cell culture to a final concentration of 1µg/ml and incubated for 10 minutes at 18 °C with 200 rpm followed by direct observation using a motorized inverted microscope (IX81; Olympus, Hamburg, Germany). The DAPI as well as the ZtGFP were exited using a standard mercury burner and imaged at 500ms exposure time.

**RNA isolation and q-RT-PCR**

*In vitro* and *in planta* expression profiling of *ZtStuA* was performed using quantitative real-time PCR (q-RT-PCR). For *in planta* analyses, the wheat cv. Taichung 29 was inoculated with Δ*ZtKu70* as described previously46 and leaf samples were collected in three biological replicates, flash frozen and ground in liquid nitrogen using a mortar and pestle. Total RNA was extracted either from ground leaves or fungal biomass produced in YGB using the RNeasy plant mini kit (Qiagen, location, USA) and subsequently DNA contamination was removed using the DNAfree kit (Ambion, Cambridgeshire, U.K.). First-strand cDNA was synthesized from approximately two µg of total RNA primed with oligo(dT) using the SuperScript III according to the manufacturer’s instructions. One µl of the resulting cDNA was used in a 25 µl PCR reaction using a QuantiTect SYBR Green PCR Kit and run and analyzed using an ABI 7500 Real-Time PCR System. The relative expression of each gene was initially normalized to the constitutively expressed*Z. tritici β-tubulin* gene54,55 and then calculated based on the comparative C(t) method described previously56. Primers used in the expression profiling analysis are detailed in Supplementary Table 5.

**Analysis of sensitivity to different stressors *in vitro***

To identify *in vitro* defects caused by the disrupted PPT-associated genes, all mutant strains were 10 μl spot-inoculated from 106 spores per ml suspensions in SDW onto YPD and/or CD minimal medium agar plates either with or without supplementation or incubation under UV light. To assess for increased ROS sensitivity, YPD plates were supplemented with 7 mM H2O2; to assess for melanisation deficiency YPD plates were incubated after an initial 6 day period for a further 6 days under UV light at 20 ̊C; to assess for lysine auxotrophy CD plates were supplemented with 100 μM lysine; and to assess for hypersensitivity to iron depletion YPD plates were supplemented with 100 μM bathophenanthroline disulphate (BPS) or 100 μM BPS and 30 μM desferriferrichrome (DFF). BPS is an iron chelating agent that reduces iron availability whereas DFF, a siderophore purified from *U. sphaerogena*, is an iron chelating agent that restores iron availability.

 When placed onto a medium containing zero nutrients such as water agar, *Z. tritici* spores undergo filamentous growth. This contrasts the budding growth observed on nutrient replete media such as YPD agar. To assess defects in filamentous growth, the two WT controls and all targeted disruption strains were spot inoculated onto water agar as described in the previous paragraph. Water agar plates were either amended with 100 µM lysine, 100 µM FeSO4, a combination of both or neither. This was to assess the impact of different PPT-associated phenotypes on filamentous growth. After 20 days, radial growth of spot inoculations was measured vertically and horizontally and the mean value of these measurements was taken.

**Statistical analysis of infection assays and *in vitro* stress response tests for the Ppt-associated and Pks mutant strains**

R version 3.2.3 was used for all statistical tests. For plant infection assays with the Ppt-associated and PKS strains, individual leaves if different seedlings were treated as replicates. Spore count data for leaves inoculated with each independent strain were assessed visually for how well they fit a normal distribution using quantile quantile plots and statistically using the Shapiro test for normality (Supplementary Figure 4). Based on data distributions observed, a log transformation was applied to each sample dataset before further analyses for Ppt associated strains (Supplementary Figure 4). One-way analysis of variance (ANOVA) was carried out on the transformed data, followed by Tukey’s honest significant difference test for pairwise differences. Differences were considered significant at α < 0.05. The analysis was carried out separately for the Δ*ZtPpt* (PPT)*,* Δ*ZtPks1,* Δ*ZtNrps1* andΔ*ZtAar*, and the PKS mutantsΔ*ZtPks7,* Δ*ZtPks8.* For the latter, no data transformations were applied as data mostly fit a normal distribution (Supplementary Figure 5b)*.* Two independent strains (in one case three) were assessed for all but two disruptions, Δ*ZtAar* and Δ*ZtPks7*.The same statistical test was carried out for the radial filamentous growth data from the spot inoculations of water agar. In this instance data were not transformed as they were adequately normally distributed for ANOVA (Supplementary Figure 3).

**Pathogenicity assays**

The susceptible wheat cultivars Taichung 29 and Riband were grown in a greenhouse until the first leaves were fully unfolded or for 17 days, respectively. Inoculum of all strains was produced in YGB (yeast extract 10g/L, Glucose 30g/L) at 18 °C for 7 days in an orbital shaker (Innova 4430; New Brunswick Scientific, Nijmegen, The Netherlands) and yeast-like spores were obtained after centrifugation at 3000 rpm and two washing steps to remove residual medium. Subsequently, the spore concentrations were adjusted to either 107 spores mL-1 for the Δ*ZtStuA* assay or 106 spores mL-1 for the infection assay of all other mutant strains. The resulting suspension was supplemented with 0.15% Tween 20 as a surfactant. Infection assays and counting of yeast-like cells on the infected leaves were conducted as previously described46,57.

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**Competing interests**

The authors have no competing interests to declare.

**Data availability**

All strains generated in this study are available upon request from the corresponding authors.

**Author contributions**

Mark C Derbyshire: design and execution of ZtPpt associated experiments, analysis and writing.

Amir Mirzadi Gohari: design and execution of all ZtStuA associated experiments, analysis and writing.

Rahim Mehrabi: Knockout construct generation, writing.

Sreedhar Kilaru: Fluorescence microscopy analysis, writing.

Gero Steinberg: Fluorescence microscopy analysis, writing.

Solaf Ali: Knockout strain generation.

Andy Bailey: Design of experiments, writing.

Kim Hammond-Kosack: Design of experiments.

Gert HJ Kema: Design of experiments, writing.

Jason J Rudd: Design of experiments, writing.

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**Figure legends**

**Figure 1. Functional characterisation of *∆ZtPpt* and related genes for auxotrophic and stress responses. (A)** From top to bottom, WT (*ΔZtKu70*), *∆ZtPpt-1* and Δ*ZtPpt-2* grown on YPD agar, YPD agar with added BPS to remove available iron, YPD agar supplemented with the BPS and DFF to remove and then restore available iron, YPD agar grown in darkness, YPD agar after a further six days of exposure to UV light, YPD agar with the ROS species hydrogren peroxide added, CD minimal agar without lysine, CD minimal agar with lysine. **(B)** From top to bottom, the two strains Δ*ZtNrps1-1* andΔ*ZtNrps1-2* grown on YPD agar, YPD agar with added BPS to remove available iron and YPD agar with added BPS and DFF to remove then restore available iron. **(C)** From top to bottom, the ∆*ZtAar* strain grown on CD minimal agar with lysine and CD minimal agar without lysine. **(D)** From top to bottom, the two melanin synthetase strains *∆ZtPks1-1* and *∆ZtPks1-2* after six days of growth on YPD in darkness and a further six days of growth under UV light. All YPD plates were grown for six days in darkness apart from those that underwent an additional six days of exposure to UV light.

**Figure 2. *In vitro* growth assays of Ppt-associated deletion strains on water agar with and without supplementation with lysine and iron.** Measurement is radial growth after 20 days. From left to right portions of figure: the two WT strains and all PPT associated strains, ∆*ZtPpt-1,* ∆*ZtPpt-2,* ∆*ZtPks1-1,* ∆*ZtPks1-2,* ∆*ZtNrps1-1,* ∆*ZtNrps1-2* and ∆*ZtAar* on water agar with no supplements; the two PPT mutant strains ∆*ZtPpt-1* and ∆*ZtPpt-2* and the two WTs on water agar with both added available iron (FeSO4) and lysine; The lysine auxotrophic PPT associated strains ∆*ZtPpt-1,* ∆*ZtPpt-2* and ∆*ZtAar* and the two WTs on water agar supplemented with lysine; the siderophore mutant strains ∆*ZtNrps1-1* and ∆*ZtNrps1-2,* the PPT mutant strains ∆*ZtPpt-1* and *,* ∆*ZtPpt-2* and the two WTs on water agar supplemented with FeSO4. Horizontal black bars represent median values, boxes represent second and third quartiles and whiskers represent interquartile range. Asterisks represent significant differences in radial growth relative to the WT that constituted the background strain at α < 0.05.

**Figure 3. *In planta* inoculation assays for Ppt-associated gene deletion strains highlights the key role of lysine biosynthesis for virulence. (A)** Upper panel: All PPT associated disruption strains, ∆*ZtPpt-1,* ∆*ZtPpt-2,* ∆*ZtPks1-1,* ∆*ZtPks1-2,* ∆*ZtNrps1-1,* ∆*ZtNrps1-2* and ∆*ZtAar,* and the two wild type strains IPO323 and ∆Z*tku70* after 22 days post inoculation (dpi) of the susceptible wheat cv, Riband; lower panel: the *Z. tritici* lysine auxotrophic PPT associated strains ∆*ZtPpt-1,* ∆*ZtPpt-2* and ∆*ZtAar,* and the two wildtype strains after 22 (dpi) with lysine added to starting inoculum. **(B)** The WT ∆Z*tku70,* and ∆*ZtPks1-1* and ∆*ZtPks1-2* melanin synthetase mutants viewed under a light microscope after 22 dpi. White arrows mark pycnidia, which in the melanin synthetase mutants are unpigmented. **(C)** Spores collected from infected leaves after 22 dpi. Left portion of plot: all PPT associated strains and the WTs without lysine added to fungal inoculum; right portion of plot: lysine auxotrophic PPT associated strains and the WTs with lysine added to fungal inoculum. Asterisks represent significant differences in spore counts relative to the WT that constituted the background strain (either IPO323 for Δ*ZtAar* or Δ*ZtKu70* for the rest) at α < 0.05. Black horizontal bars represent median values, boxes represent second and third quartiles and whiskers represent interquartile range.

**Figure 4. *In planta* inoculation assays for the two PKS deletion mutant strains ∆*ZtPks7* and ∆*ZtPks8* reveal them to be dispensable for virulence.** **(A)** Symptoms after 22 dpi on the susceptible wheat cv. Riband. Left: The WT (∆Z*tKu70*), the sole ∆*ZtPks7* strain and one of the ∆*ZtPks8* strains, ∆*ZtPks8-1*; right: The WT (∆*Ztku70*) and three ∆*ZtPks8* mutant strains, ∆*ZtPks8-1,* ∆*ZtPks8 -2* and ∆*ZtPks8-3*. **(B)** Spores retrieved from leaves after 22 dpi. Horizontal bars represent median values, boxes represent second and third quartiles and whiskers represent interquartile range. No significant differences were detected.

**Figure 5. Summary of fungal knockout strains generated in this study, downstream processes regulated by them and effects on virulence.**

**Supplementary figure legends**

**Supplementary Figure 1. Generation and validation of** *∆****ZtPpt* and related gene deletion strains. (A)** Genes are represented by red arrows. Above each arrow, the lines represent the two flanking sequences with length in base pairs (bp) given above, the insertion position of the *hph* gene under the *trpC* promoter and positions of primers represented by arrows used to determine successful integration. **(B)** Composite PCR agarose gel showing amplification of regions targeted in mutant strains and WT genomic DNA. A single band of correct size was considered indicative of successful targeted gene disruption. **(C)** Diagram showing the primers used to confirm the Aar2 deletion. PCR amplification hygromycin B resistance gene, Lane 1 Hyperladder 1, lane 2 negative control D.W, lane 3 wild type IPO323. Lane 4 to 12; putative transformants. Lanes 5, 6 and 9 show expected bands of 1.7kb in Δ*Aar* mutants. **(D)** The pair of primers were used to confirm the presence of *Aar*. PCR amplification of *Aar* wild type locus, lane 5, 6 and 9 no band was seen indicating deletion of *Aar* gene.

**Supplementary Figure 2. Generation and functional characterisation of Δ*ZtStuA* gene deletion strains (A)** Validation of *ZtStuA* gene deletion strains**.** Diagram displaying the replacement by the *hygromycin phosphotransferase* (*hph*) resistance cassette through homologous recombination. The Δ*ZtKu70*, complemented strain as well as two independent mutant strains were used for PCR amplification using primers ZtStuA-F1, ZtStuA-R1, ZtStuA-F2 and ZtStuA-R2. Lanes 1 and 2 *Z. tritici* Δ*Ku70* (*Zt* Δ*Ku70*). Lanes 3 and 4 Δ*ZtStuA-C.* Lanes 5 and 6 Δ*ZtStuA-1*. Lanes 7 and 8 Δ*ZtStuA-2*. Lane 1 shows the amplification of *ZtStuA* in *ΔZtKu70* using primers ZtStuA-F1 and ZtStuA-R1 designed to amplify *ZtStuA* ORF whereas no amplicon of *ZtStuA* was observed in lane 2 with primers ZtStuA-F2 and ZtStuA-R2. Lane 3 shows the expected band of 1.6 kb in Δ*ZtStuA-C* amplified by using primers ZtStuA-F1 and ZtStuA-R1 whereas no amplicon of *ZtStuA* was observed in lane 4 with primers ZtStuA-F2 and ZtStuA-R2. Lane 5 indicates no amplicon of *ZtStuA* in Δ*ZtStuA-1* by using primers ZtStuA-F1 and ZtStuA-R1 while the expected band of 1.9 kb in lane 6 by using primers ZtStuA-F2 and ZtStuA-R2 was visualized. Lane 7 indicates no amplicon of *ZtStuA* ORF in Δ*ZtStuA-2* by using primers ZtStuA-F1 and ZtStuA-R1 while the expected band of 1.9 kb in lane 7 by using primers ZtStuA-F2 and ZtStuA-R2 was observed. **(B)** Effects on yeast-like cell production and biosynthesis of melanin. Upper panels- The Δ*ZtKu70* and Δ*ZtStuA-C* strains generated abundant yeast-like cells derived from blastic conidiogenesis in yeast glucose broth medium while Δ*ZtStuA-1* and *2* failed to sporulate and exclusively produced compact hyphal networks(marked with a black arrow)*.*Middle panels- The Δ*ZtKu70* and Δ*ZtStuA-C* strains became melanizedwhereas the strains deleted for *ZtStuA* remained unmelanized. Lower panels- Comparative *in vitro* expression of putative genes involved in the melanization event inΔ*ZtKu70* versus the Δ*ZtStuA-1* strain. Error bars show standard deviation of the mean. **(C)** ZtStuA is localised to the nucleus.Left panels-The subcellular localization of the fluorescent protein ZtStuA::GFP was determined in yeast-like cells. Middle panels- localisation in hyphae; scale bars = 10 μm. Right panels- Yeast-like cells viewed under a fluorescent microscope with 4′,6-diamidino-2-phenylindole (DAPI) staining. Fluorescence co-localizes with the DAPI-stained nucleus; bars = 5 μm. **(D)** The effect of *Zymoseptoria triici StuA* (*ZtStuA*) deletion on disease development in the susceptible wheat cv. Taichung 29. Upper panels- From left to right, first leaves were inoculated with water (as a control) and the Δ*ZtKu70*, Δ*ZtStuA-C,* Δ*ZtStuA-1* and Δ*ZtStuA-2* strains. Final disease levels shown 20 days post inoculation (dpi). Lower panels-Number of spores harvested from replicate leaves for the ΔZ*tKu70*, Δ*ZtStuA-C,* Δ*ZtStuA-1* and Δ*ZtStuA-2* strains. No spores were produced after infection with *ZtStuA* knockout strains. Error bars show standard deviation of the mean.

**Supplementary Figure 3. Quantile plots of data from *in vitro* radial hyphal growth assay.** Data are shown for all *in vitro* conditions tested using water agar for which more than 0 mm of radial growth was observed. Data were not transformed as they did not fail the Shapiro normality test at α = 0.05.

**Supplementary Figure 4. Quantile plots of spore counts from samples of infected leaves.** Data are shown for all strains that generated spores (several did not generate any spores, being completely avirulent, and were therefore omitted from statistical analysis) **(A)** Upper panel: Spore counts for all Ppt-associated strains and the two WTs after 22 dpi without lysine in the fungal inoculum. Grey points indicate a sample that failed the Shapiro normality test at α < 0.05. Black points indicate data that were statistically likely to have come from a normally distributed population α > 0.05. Lower panel: The same plots after data were log transformed. This reduced the number of samples that failed the Shapiro normality test at α < 0.05. Grey points indicate a sample that failed the Shapiro normality test at α < 0.05. Black points indicate data that were statistically likely to have come from a normally distributed population α > 0.05. **(B)** Left: spore counts for the WT strain ∆*ZtKu70* and two of the PKS mutants, ∆*ZtPks7* and ∆*ZtPks11-1* after 22 dpi. Right: spore counts for the WT strain ∆*ZtKu70* and the three ∆*ZtPks11* strains tested. Data were not transformed. Grey points indicate a sample that failed the Shapiro normality test at α < 0.05. Black points indicate data that were statistically likely to have come from a normally distributed population α > 0.05.

**Supplementary table legends**

**Supplementary Table 1. *Zymoseptoria tritici* has homologues of the PPT gene and associated genes characterised in *Cochliobolus sativus*.**

**Supplementary Table 2. Homology of the *Zymoseptoria tritici* siderophore synthetase gene to siderophore synthetases from *Aspergillus fumigatus*.**

**Supplementary Table 3. *Zymoseptoria tritici* hashomologues of *Aspergillus fumigatus* genes encoded the DHN-melanin pathway.**

**Supplementary Table 4. Primers used in knockout construct generation and confirmation of homologous recombination.**

**Supplementary Table 5. Primers used in the expression analysis of melanin biosynthesis genes.**