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The chitin synthase FgChs2 and other FgChss co-regulate vegetative development and virulence in *F. graminearum*

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Fusarium graminearum contains eight chitin synthase (*Chs*) genes belonging to seven classes. Previous studies have found that deletion of *FgChs3b* is lethal to *F. graminearum*, and deletion of *FgChs1*, *FgChs2*, *FgChs7* and *FgChs5* caused diverse defects in chitin content, mycelial growth, conidiation, virulence or stress responses. However, little is known about the functional relationships among these *FgChss*. In this study, *FgChs2* deletion mutant Δ *FgChs2* exhibited reduced mycelial growth and virulence as reported previously. In addition, we found that the mutant produced thickened and “wavy” septa. Quantitative real-time PCR (qRT-PCR) assays showed that the expression levels of *FgChs1*, *FgChs3a*, *FgChs4*, *FgChs7*, *FgChs5* and *FgChs6* in Δ *FgChs2* were significantly higher than those in the wild type. Therefore, we generated six double deletion mutants of *FgChs2* and each of the above six *FgChss*, and found that *FgChs2* shares a function with *FgChs1* in regulating mycelial growth, and co-regulates conidiation with *FgChs1*, *FgChs4*, *FgChs7* and *FgChs5*. Furthermore, *FgChs2* and other six *FgChss* have overlapped functions in virulence, DON production and septum formation. Taken together, these results indicate that although each chitin synthase of *F. graminearum* plays certain roles, *FgChss* may co-regulate various cellular processes in *F. graminearum*.

Chitin, a β (1, 4)-linked homopolymer of *N*-acetylglucosamine (GlcNAc), is an essential component of cell walls and septa of all fungi studied to date^{1,2}. The synthesis of chitin is mediated by membrane-bound chitin synthases (*Chss*), which were divided into seven classes³. There are three *Chs* genes in budding yeast *Saccharomyces cerevisiae*, while filamentous fungi generally contain seven or eight *Chs* genes. Chitin synthases belonging to classes III, V, VI, and VII are only identified in filamentous fungi and some dimorphic yeasts³, which may result in higher chitin content and greater complexity of growth and development of these fungi than the budding yeast. In filamentous fungi, chitin accounts for 10–20% of dry weight content of cell wall in vegetative cells, which is much higher than 1–2% in *S. cerevisiae*^{4,5}.

In *S. cerevisiae*, three *Chss* have been extensively studied and their functions have been well understood. The Class I *Chs* (*ScChs1*) repairs the weakened cell wall of daughter cells after separation⁶. *ScChs2* (II) is essential for both septum formation and cell division⁷. *ScChs3* (IV) synthesizes 90% of chitin in the cell walls and is required for chitin ring formation at the base of emerging buds and chitin synthesis in the lateral cell⁸. However, the functions of individual *Chss* and their specific involvements and interactions remain poorly understood in filamentous fungi. One of main reasons might be functional overlap among multiple *Chss* in filamentous fungi.

Fusarium graminearum (teleomorph *Gibberella zeae*) causes Fusarium head blight (FHB), which is a devastating disease of cereal crops worldwide. Infection of cereal crops with *F. graminearum* may not only lead to huge yield losses in severe epidemic years, but also pose a serious threat to human and animal health owing to deoxynivalenol (DON) and other mycotoxins in infested grains^{9,10}. Despite the serious damage caused by FHB, efficient strategies for the management of FHB are not available to date¹¹. Previous studies on *Magnaporthe oryzae* and *Botrytis cinerea* have showed that chitin synthases play important roles in fungal growth and pathogenicity^{12–16}.

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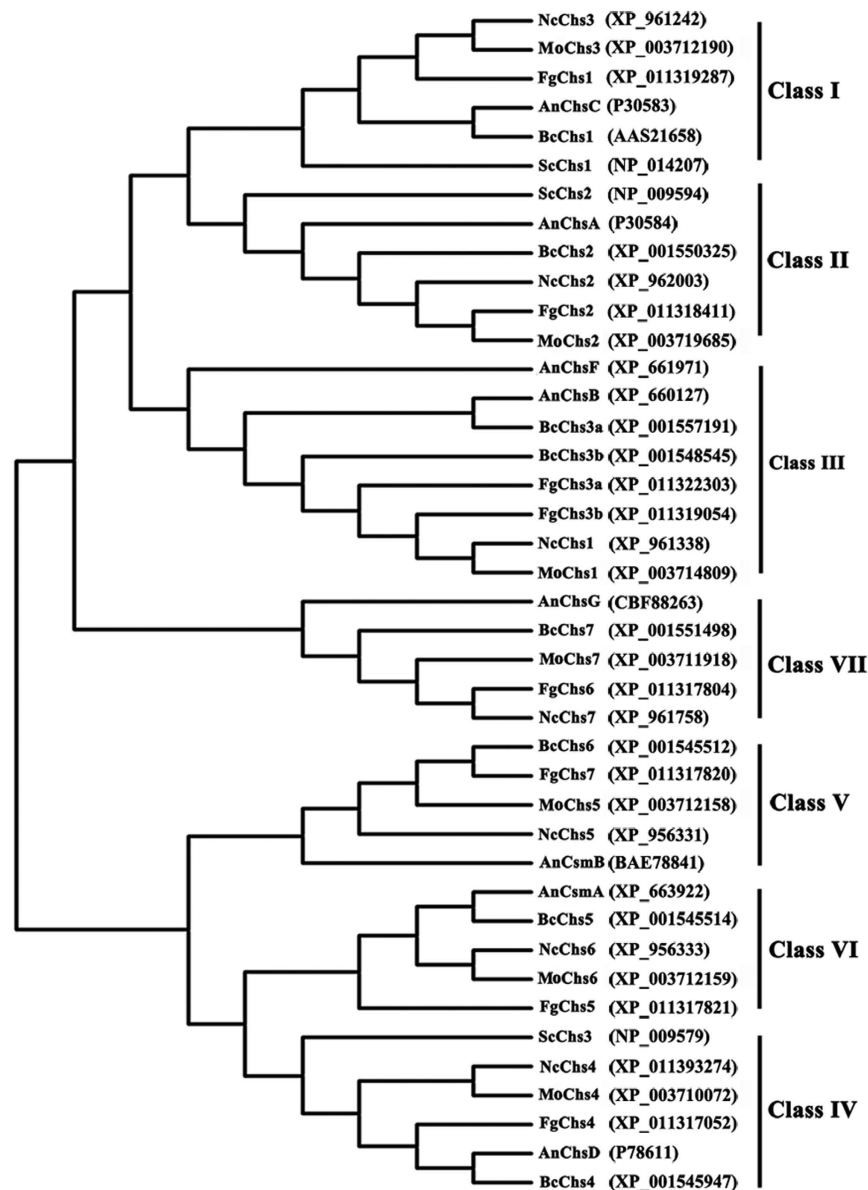


Figure 1. Phylogenetic tree of fungal chitin synthases. Phylogenetic tree generated using the neighbor-joining method with Mega 5.0 software on the basis of deduced amino acid sequences of chitin synthases from different fungi. FgChs1, FgChs2, FgChs3a, FgChs3b, FgChs4, FgChs5, FgChs6 and FgChs7 from *Fusarium graminearum*; AnChsA, AnChsB, AnChsC, AnChsD, AnChsF, AnChsG, AnCsmA and AnCsmB from *Aspergillus nidulans*; BcChs1, BcChs2, BcChs3A, BcChs3B, BcChs4, BcChs5, BcChs6 and BcChs7 from *Botrytis cinerea*; NcChs1, NcChs2, NcChs3, NcChs4, NcChs5, NcChs6 and NcChs7 from *Neurospora crassa*; MoChs1, MoChs2, MoChs3, MoChs4, MoChs5, MoChs6 and MoChs7 from *Magnaporthe oryzae*; ScChs1, ScChs2 and ScChs3 from *Saccharomyces cerevisiae*. GenBank accession no. of each protein was presented in brackets.

Thus, deep understanding the biological functions of Chs in plant pathogenic fungi can provide the basis for the development of chitin synthase-targeted antifungal agents. What's more, such antifungal agents might be safe to high eukaryotes since chitin and chitin synthases are not present in animals and plants^{1,3}.

In silico analyses showed that the *F. graminearum* genome contains eight *FgChs* genes (Fig. 1). Following the classification proposed by Chigira *et al.* and Choquer *et al.*^{17,18}, these genes are referred as *FgChs1* (I), *FgChs2* (II), *FgChs3a* (III), *FgChs3b* (III), *FgChs4* (IV), *FgChs7* (V), *FgChs5* (VI), and *FgChs6* (VII) respectively in this study (Fig. 1). Previous reports have indicated that *FgChs3b* is essential, and the deletion of *FgChs1*, *FgChs2*, *FgChs7* and *FgChs5* led to reduced mycelial growth, virulence or increased sensitivity to various stresses^{19–21}. The deletion of *FgChs3a*, *FgChs4* and *FgChs6* genes did not cause significant differences from the wild type²¹. Previous studies were conducted with the different genetic background strains, we therefore constructed various *Chs* deletion strains in a single progenitor in current study in order to characterize *FgChs*s systemically. Results of this study indicated that *FgChs2* and other *FgChs* genes co-regulate various cellular processes in *F. graminearum*.

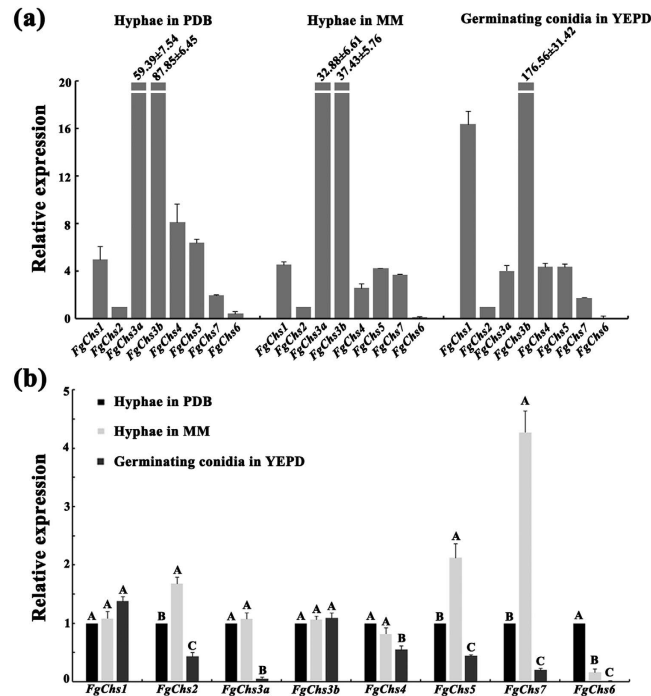


Figure 2. Expression profiles of eight *FgChs* genes assayed by qRT-PCR. RNA samples of the wild-type PH-1 were isolated from vegetative hyphae grown in PDB or MM, and from germinating conidia cultured in YEPD. The relative expression level of individual *FgChs* gene was analyzed with the $2^{-\Delta\Delta C_t}$ method with the *ACTIN* gene as the internal control for normalization. (a) Comparison of the transcript abundance of eight *FgChs* genes in hyphae grown in PDB and MM, and germinating conidia cultured in YEPD. The expression level of *FgChs2* was referred to 1. (b) Comparison of the transcript abundance of individual *FgChs* genes in hyphae and germinating conidia. The expression level of each *FgChs* gene in vegetative hyphae grown in PDA was referred to 1. Mean and standard errors were determined with data from three independent replicates. Values on the bars followed by the same letter are not significantly different according to a least significant difference (LSD) test at $P=0.05$.

Results

Eight chitin synthase genes of *F. graminearum* are differentially expressed in both mycelia and conidia. The expression levels of *FgChs* genes in mycelia cultured in PDB and MM, and in germinating conidia grown in YEPD were determined by quantitative real-time PCR (qRT-PCR) assays. Among the eight genes, the abundance of *FgChs6* transcripts was the lowest in both vegetative hyphae and germinating conidia (Fig. 2a). In contrast, the *FgChs3b* had the highest expression levels (Fig. 2a). The *FgChs2*, *FgChs7* and *FgChs5* genes had similar expression profiles with higher expression levels in mycelia grown in MM than in PDB (Fig. 2b). With the exception of *FgChs1* and *FgChs3b*, other *FgChs* genes exhibited higher expression levels in hyphae than in germinating conidia (Fig. 2b).

The expression levels of other seven *FgChs* genes in $\Delta FgChs2$. To explore the relationships of *FgChs2* and other *FgChs*, we determined the expression levels of other seven *FgChs* genes in the *FgChs2* deletion mutant $\Delta FgChs2$ by qRT-PCR assays. As shown in Fig. 3, the expression levels of *FgChs1*, *FgChs3a*, *FgChs4*, *FgChs7*, *FgChs5* and *FgChs6* in $\Delta FgChs2$ were significantly higher than those in the wild-type PH-1. Based on the results of qRT-PCR assays, the double mutants of $\Delta FgChs2/1$, $\Delta FgChs2/3a$, $\Delta FgChs2/4$, $\Delta FgChs2/7$, $\Delta FgChs2/5$ and $\Delta FgChs2/6$ were constructed by deletion of *FgChs1*, *FgChs3a*, *FgChs4*, *FgChs7*, *FgChs5* and *FgChs6* in $\Delta FgChs2$, respectively (Fig. S1).

***FgChs2* shares a function with *FgChs1* in regulating mycelial growth.** Previous studies have found that deletion of *FgChs2*, *FgChs7* or *FgChs5* caused the reduction of mycelial growth in *F. graminearum*, and the deletion mutants $\Delta FgChs3a$, $\Delta FgChs1$, $\Delta FgChs4$ and $\Delta FgChs6$ had an undistinguishable growth rate compared with that of the wild type^{19–21}. In this study, we found that the double mutant $\Delta FgChs2/1$ grew much slower than the single deletion mutants $\Delta FgChs1$ and $\Delta FgChs2$ on both PDA and MM media (Fig. 4a,b). Moreover, aerial hyphae of double mutant $\Delta FgChs2/1$ were developed poorly (Fig. 4a). To a lesser extent, the double mutants $\Delta FgChs2/7$ and $\Delta FgChs2/5$ showed reduced mycelial growth in comparison with the single mutants $\Delta FgChs2$, $\Delta FgChs7$ and $\Delta FgChs5$ (Fig. 4a,b). In contrast, the double mutants $\Delta FgChs2/3a$, $\Delta FgChs2/4$ and $\Delta FgChs2/6$ exhibited similar growth rate with the single mutant $\Delta FgChs2$. These results indicate that *FgChs2* has an overlapping function with *FgChs1* in regulating mycelial growth in *F. graminearum*.

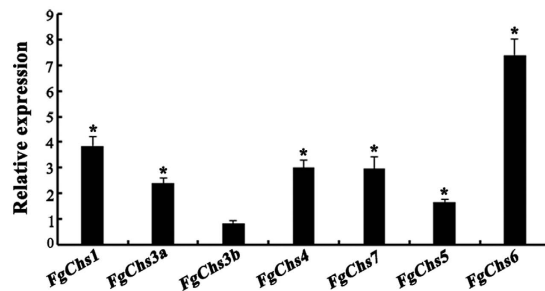


Figure 3. Effect of *FgChs2* deletion on the transcription of other *FgChs* genes assayed by qRT-PCR. The relative expression level of each *FgChs* gene in the *FgChs2* deletion mutant $\Delta FgChs2$ is the relative amount of mRNA in the wild type. Line bars in each column denote standard errors of three repeated experiments. A *t* test was performed to determine significant differences, *significant difference for each gene at a 95% coincidence interval.

Overlapping function in conidiation between *FgChs2* and *FgChs1*, *FgChs4*, *FgChs7* or *FgChs5*.

According to the previous studies^{19,20}, conidial production of the mutants $\Delta FgChs1$, $\Delta FgChs7$ and $\Delta FgChs5$ was dramatically reduced. In this study, we determined the phenotypes of asexual development for the single mutants $\Delta FgChs2$, $\Delta FgChs3a$, $\Delta FgChs4$ and $\Delta FgChs6$ and the double mutants $\Delta FgChs2/1$, $\Delta FgChs2/3a$, $\Delta FgChs2/4$, $\Delta FgChs2/7$, $\Delta FgChs2/5$ and $\Delta FgChs2/6$. The mutant $\Delta FgChs2$ showed a decrease of 45.2% in conidiation and produced smaller conidial spores with less septation (Fig. 5a–d). Deletion of *FgChs3a*, *FgChs4* and *FgChs6* genes did not cause significant difference in asexual development in comparison with the wild type (Fig. 5a–d). The mutants $\Delta FgChs2/3a$ and $\Delta FgChs2/6$ produced similar amount of conidia as $\Delta FgChs2$ (Fig. 5a,b). Whereas, the double mutant $\Delta FgChs2/1$ was unable to produce conidium in CMC even 15 days after inoculation. The mutants $\Delta FgChs2/4$, $\Delta FgChs2/7$ and $\Delta FgChs2/5$ exhibited a significantly reduced conidiation compared with the corresponding single mutants. Moreover, microscopic examination showed that the mutants $\Delta FgChs2/4$, $\Delta FgChs2/7$ and $\Delta FgChs2/5$ had more severe defects in conidium length and septation in comparison with the corresponding single mutants (Fig. 5a,c,d). These results indicated that the functions of *FgChs2* in regulating conidiation are partially exchangeable with those of *FgChs1*, *FgChs4*, *FgChs7*, and *FgChs5*.

Although the mutants $\Delta FgChs1$, $\Delta FgChs2$, $\Delta FgChs7$, $\Delta FgChs5$, $\Delta FgChs2/4$, $\Delta FgChs2/7$ and $\Delta FgChs2/5$ produced more shortened conidia with less septa, more than 90% conidia of each mutant as well as the wild type, could germinate after 4 h of incubation in 2% (w/v) sucrose solution (Fig. S2).

FgChs2 co-regulates virulence and DON biosynthesis with *FgChs3a*, *FgChs1*, *FgChs4*, and *FgChs6*.

Among the eight *FgChs*s in *F. graminearum*, *FgChs1*, *FgChs2*, *FgChs7* and *FgChs5* have been found to play important roles in virulence^{19–21}. The mutants $\Delta FgChs7$ and $\Delta FgChs5$ almost lost virulence on flowering wheat head¹⁹, and the mutants $\Delta FgChs2$ and $\Delta FgChs1$ showed significantly decreased virulence^{20,21}. Here, we determined the virulence of the double mutants on wheat head, and found that the double mutants $\Delta FgChs2/3a$, $\Delta FgChs2/4$ and $\Delta FgChs2/6$ caused the scab symptoms only in the inoculated spikelets and $\Delta FgChs2/1$, $\Delta FgChs2/7$, and $\Delta FgChs2/5$ could not cause any scab symptom in the inoculated spikelets 15 days after inoculation (Fig. 6a). After 25 days of inoculation, the scab symptoms caused by the single mutants $\Delta FgChs7$ and $\Delta FgChs5$ could be observed only in the inoculated spikelets, but no visible scab symptoms were found for the double mutants $\Delta FgChs2/1$, $\Delta FgChs2/7$ and $\Delta FgChs2/5$ (data not shown). These results indicate that there are overlapping functions between *FgChs2* and *FgChs1*, *FgChs3a*, *FgChs4*, *FgChs7*, *FgChs5* or *FgChs6* in regulating virulence in *F. graminearum*. Furthermore, cellophane penetration assays were performed for the mutants $\Delta FgChs7$, $\Delta FgChs5$, $\Delta FgChs2/1$, $\Delta FgChs2/7$ and $\Delta FgChs2/5$. As shown in Fig. S3, these mutants could penetrate cellophane sheets, indicating that the dramatically reduced virulence of these mutants might mainly be owing to other factors than host penetration.

DON is an important virulence factor of *F. graminearum*^{22–24}. Therefore, we were interested in analyzing DON biosynthesis in each mutant since studies on functions of *FgChs*s in DON production have not been conducted previously. As shown in Fig. 6b, after culture on sterilized wheat kernels for 30 days, single mutants $\Delta FgChs1$, $\Delta FgChs2$, $\Delta FgChs4$, $\Delta FgChs7$ and $\Delta FgChs5$ showed reduced DON production by 13.7 to 84.5% (Fig. 6b). Double deletion of *FgChs2* and any other *FgChs* gene intensified the decrease of DON production (Fig. 6b).

The trichothecene (*TRI*) genes are responsible for DON biosynthesis^{25,26} and the expression of *TRI4*, *TRI5* and *TRI6* have a positive correlation with the DON production in *F. graminearum*²⁷. To confirm the decreased DON in *FgChs* mutants, we further assayed the expressions of *TRI4*, *TRI5* and *TRI6* genes by qRT-PCR assays. The expression levels of three *TRI* genes in six double mutants were dramatically lower than those in all single mutants (Fig. 6c).

FgChs1, *FgChs3a*, *FgChs4*, *FgChs7*, *FgChs5* and *FgChs6* have additive effects in septum formation.

In *F. graminearum*, only one previous study clearly reported that the septal pores in the mutant $\Delta FgChs7$ ($\Delta FgChs7$) was observed to be plugged by a woronin body-like structure through transmission electron microscopy examination¹⁹. Here, we found that deletion of *FgChs2* caused thickened (Fig. 7, left panel of $\Delta FgChs2$) and “wavy” septa (Fig. 7, middle panel of $\Delta FgChs2$) occasionally with a larger central pore (Fig. 7, right panel of $\Delta FgChs2$), although deletion of other single *FgChs* could not result in recognizable changes in

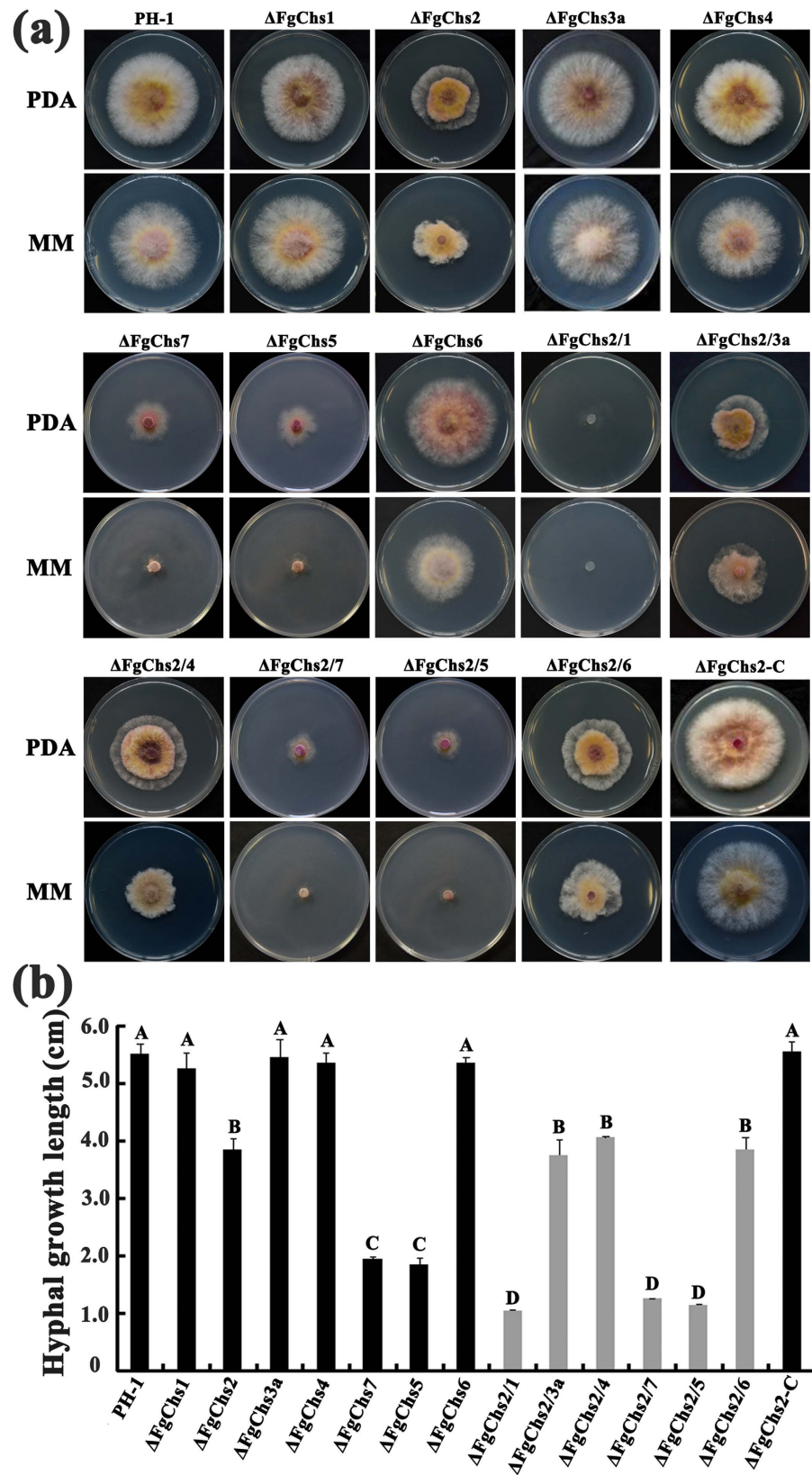


Figure 4. Impacts of *FgChs* single and double deletion on *F. graminearum* hyphal growth . (a) Colony morphology of *FgChs* single and double deletion mutants. The wild-type PH-1, *FgChs* deletion mutants (Δ FgChs1-7), double deletion mutants of *FgChs2* and other *FgChs*s (Δ FgChs2/1-7), and the complemented strain Δ FgChs2-C were grown on PDA or MM at 25 °C for 3 days. **(b)** Colony diameter of each strain cultured on PDA at 25 °C for 3 days. Line bars in each column denote standard errors of three repeated experiments. Values on the bars followed by the same letter are not significantly different according to a least significant difference (LSD) test at $P=0.05$.

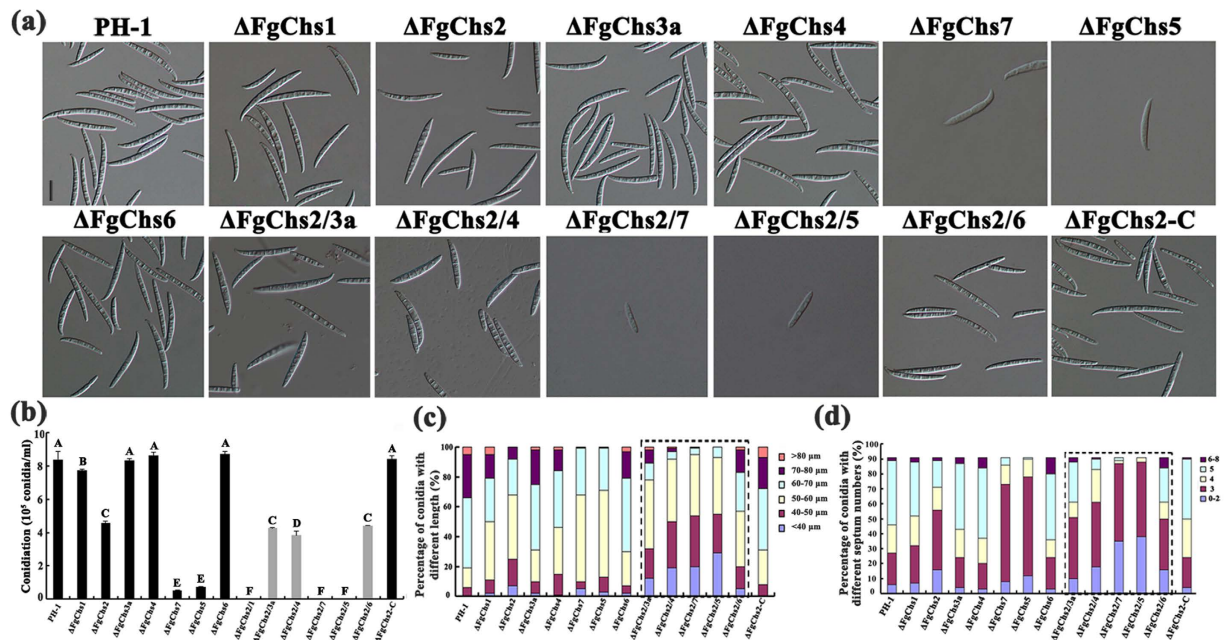


Figure 5. Involvement of *FgChs* in regulating conidiation in *F. graminearum*. (a) Conidial morphology of the wild type, *FgChs* deletion mutants ($\Delta FgChs1-7$), double deletion mutants of *FgChs2* and other *FgChs* ($\Delta FgChs2/1-7$), and the complemented strain $\Delta FgChs2-C$. The differential interference contrast (DIC) images of conidia were captured with an electronic microscope. Bar = 20 μm . (b) The quantity of conidia produced by each strain in carboxymethyl cellulose liquid medium (CMC) for 4.5 days in a shaker. Values on the bars followed by the same letter are not significantly different according to a least significant difference (LSD) test at $P = 0.05$. (c) Comparisons of conidial length among the above strains. A total of 200 conidia were examined for each strain. (d) Comparisons of the percentage of conidia with different septum numbers among the above strains. A total of 200 conidia were examined for each strain.

septum morphology (data not shown). Interestingly, the double mutants $\Delta FgChs2/1$, $\Delta FgChs2/3a$, $\Delta FgChs2/4$, $\Delta FgChs2/7$, $\Delta FgChs2/5$ and $\Delta FgChs2/6$ produced more thickened septa than $\Delta FgChs2$ (Fig. 7). To a great extent, the double mutant $\Delta Chs2/1$ could not form complete septum structure with aberrant thickness and abnormally large pores. These results indicate that although *FgChs2* plays an important role in septation, *FgChs1*, *FgChs3a*, *FgChs4*, *FgChs7*, *FgChs5* or *FgChs6* have additive effects in septum formation in *F. graminearum*.

***FgChs2*, *FgChs5* and *FgChs7* play an important role in the response to cell wall stress.** Among the eight *FgChs*s in *F. graminearum*, *FgChs1*, *FgChs2*, *FgChs7* and *FgChs5* have been found to be involved in the response to various stresses^{19–21}. To explore the function of *FgChs*s in cell wall stress response, serial dilutions of conidial suspension of each strain were spotted on PDA amended with 0.2 g/l of cell wall-damaging agent congo red (CR). After incubation at 25 °C for 3 days, the single gene mutants $\Delta FgChs2$, $\Delta FgChs5$, $\Delta FgChs7$, but not $\Delta FgChs1$, $\Delta FgChs3a$, $\Delta FgChs4$ and $\Delta FgChs6$, showed increased sensitivity to CR dramatically (Fig. 8a). In addition, we determined the sensitivity of $\Delta FgChs2/1$ to CR using mycelial plugs since this mutant was unable to produce conidia. As shown in Fig. 8b, $\Delta FgChs2/1$ and $\Delta FgChs2$, presented similar sensitivity to CR. To further verify this finding, we tested the sensitivity of each strain to cell wall-degrading enzymes. As shown in Fig. 8c, hyphae of the single gene mutants $\Delta FgChs2$, $\Delta FgChs7$, $\Delta FgChs5$, and the double mutants $\Delta FgChs2/1$, $\Delta FgChs2/3a$, $\Delta FgChs2/4$, $\Delta FgChs2/7$, $\Delta FgChs2/5$ and $\Delta FgChs2/6$ all were well digested and released abundant protoplasts after treatment with cellulase, lysozyme and snailase at 30 °C for 30 min. However, the single gene mutants $\Delta FgChs1$, $\Delta FgChs3a$, $\Delta FgChs4$ and $\Delta FgChs6$ could not be digested adequately. These results indicated that $\Delta FgChs2$, $\Delta FgChs5$, $\Delta FgChs7$, but not $\Delta FgChs1$, $\Delta FgChs3a$, $\Delta FgChs4$ and $\Delta FgChs6$, play an important role in response to cell wall stress.

Discussion

Chitin synthases from various fungi have been grouped into seven classes³. *Neurospora crassa* and *M. oryzae* contain seven chitin synthase genes. However, *F. graminearum* contains eight predicted *FgChs* genes (Fig. 1). In this study, we found that eight *FgChs*s exhibited different expression patterns in hypha and conidia. In comparison with other *FgChs*s, *FgChs3b* exhibited the highest expression levels (Fig. 2a). Consistent with a previous report²¹, we were unable to knockout *FgChs3b*, indicating that it might be essential in *F. graminearum*. Additionally, *FgChs2*, *FgChs7* and *FgChs5* exhibited higher expression in hyphae grown in MM than in PDA. Mycelial growth assays found that the growth rates of $\Delta FgChs2$, $\Delta FgChs7$ and $\Delta FgChs5$ on MM plates was much slower than those on PDA plates, in contrast, the growth rate of the wild type was similar on both plates (Fig. 4a). These

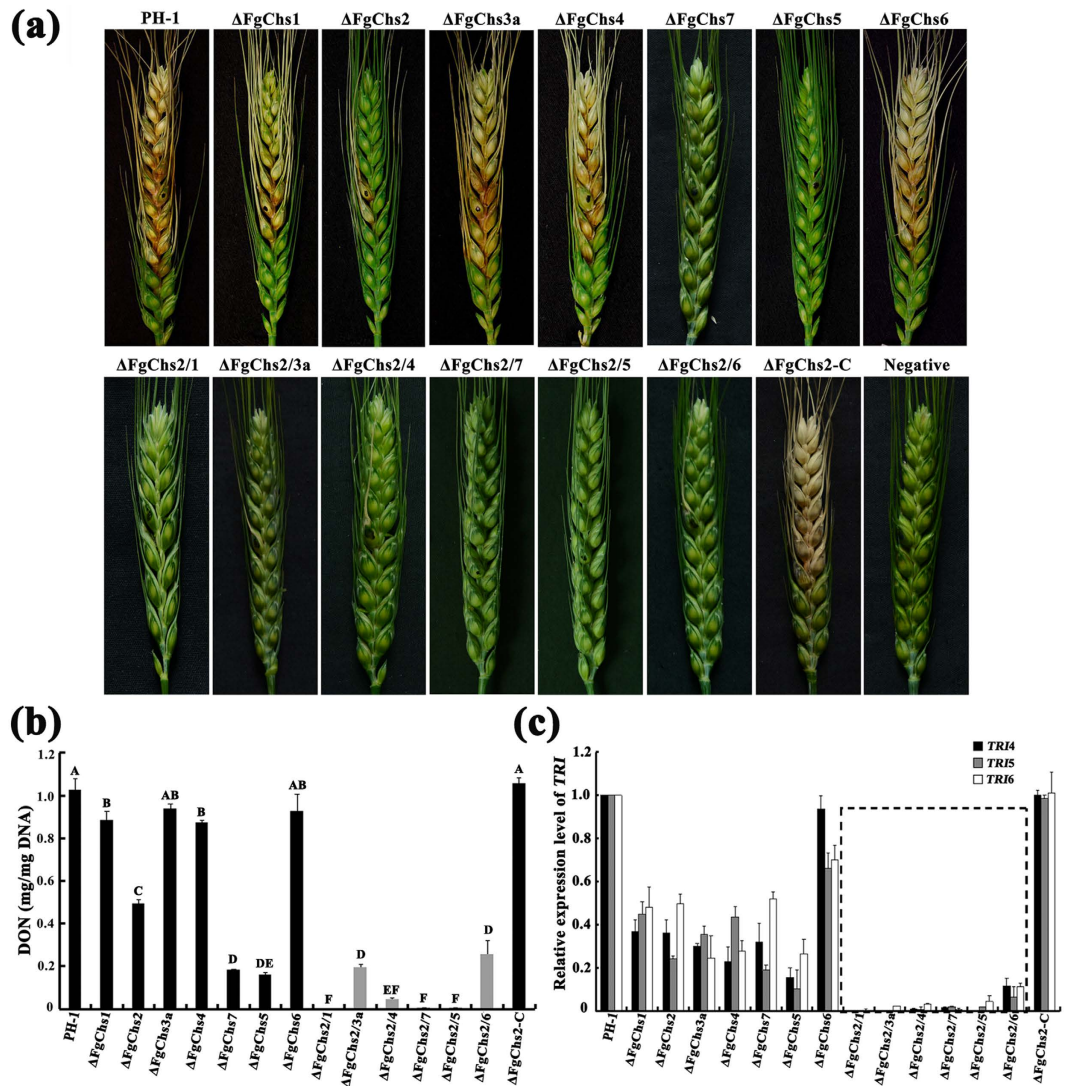


Figure 6. Impacts of *FgChs* deletion on virulence and DON biosynthesis. (a) Flowering wheat heads were point inoculated with a conidial suspension at 10^5 conidia/ml of the wild-type PH-1, *FgChs* deletion mutants ($\Delta FgChs1-7$), double deletion mutants of *FgChs2* and other *FgChss* ($\Delta FgChs2/1-7$), and the complemented strain $\Delta FgChs2-C$. The infected wheat heads were photographed 15 days after inoculation. (b) The amount of DON (per mg fungal DNA) produced by each strain in infected wheat kernels was determined after 30 days of inoculation. Line bars in each column denote standard errors of three replicated experiments. Values on the bars followed by the same letter are not significantly different according to a least significant difference (LSD) test at $P=0.05$. (c) Relative expression of DON biosynthetic *TRI* genes in each strain and bars denote standard errors from three repeated experiments.

results indicate that the three *FgChss* (*FgChs2*, *FgChs7* and *FgChs5*) might play more important roles in nutrient deficiency conditions.

All *Chss* responsible for the polymerization of GlcNAc contain conserved chitin synthase and transmembrane domains. Additionally, classes V and VI *Chss* have the myosin motor domain (MMD) at their N-terminal end²⁷. Several studies focusing on the functions of *Chss* in fungi have found that *Chss* belonging to different classes jointly play roles in hyphal growth, asexual and sexual development, pathogenicity and response to stresses^{12,27-31}, which may indicate functional redundancy in various facets among *Chss*. In *F. graminearum*, the individual *FgChss* have been characterized, the interactions of *FgChs2* with other *Chss* were therefore the main point explored in this study. We found *FgChs2* shares functions in mycelial growth with *FgChs1*, and to a lesser extent, with *FgChs7* and *FgChs5* in *F. graminearum* (Fig. 4a,b). The double mutant $\Delta FgChs2/1$ grew dramatically slow and produced fewer aerial hyphae. Although the previous study found that deletion of *FgChs1* does not affect biomass and the hyphal growth rate²⁰, our study showed that *FgChs1* still plays an important role in hyphal growth. A similar finding has been reported in *Aspergillus nidulans* and *M. oryzae*. Fujiwara *et al.* (2000) reported that the *AnchsC* (I) *AnchsA* (II) double mutant of *A. nidulans* showed fewer aerial hyphae and lower hyphal density²⁹, which is similar to the phenotypes of $\Delta FgChs2/1$ in this study. The hyphae of the *AnchsB* (III) *AnchsD* (IV)

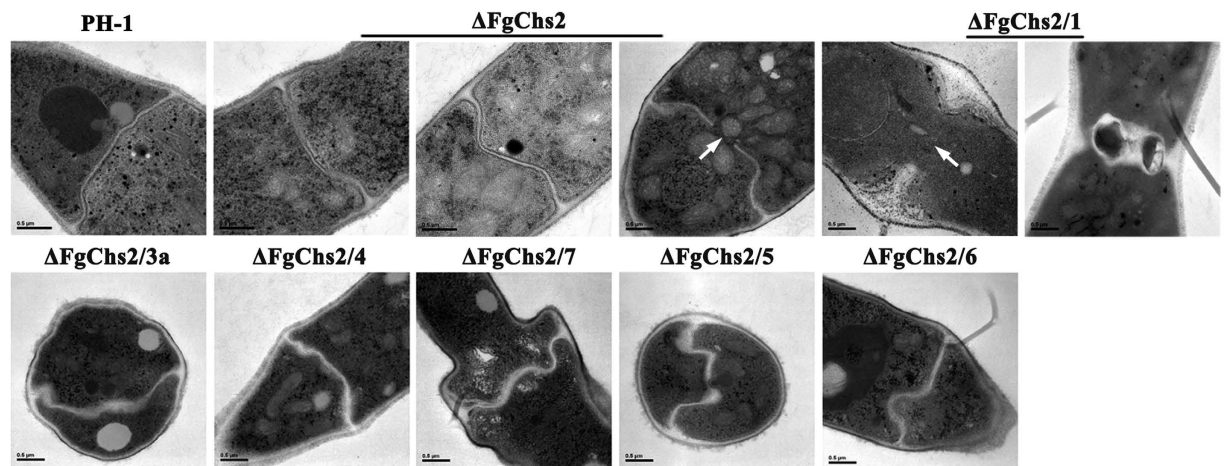


Figure 7. Involvement of FgChs in septum formation in *F. graminearum*. Transmission electron microscopic examination of hyphae from the wild-type PH-1, *FgChs2* deletion mutant $\Delta FgChs2$, double deletion mutants of *FgChs2* and other *FgChs*, $\Delta FgChs2/1$, $\Delta FgChs2/3a$, $\Delta FgChs2/4$, $\Delta FgChs2/7$, $\Delta FgChs2/5$ and $\Delta FgChs2/6$. Septal pores are indicated by white arrows.

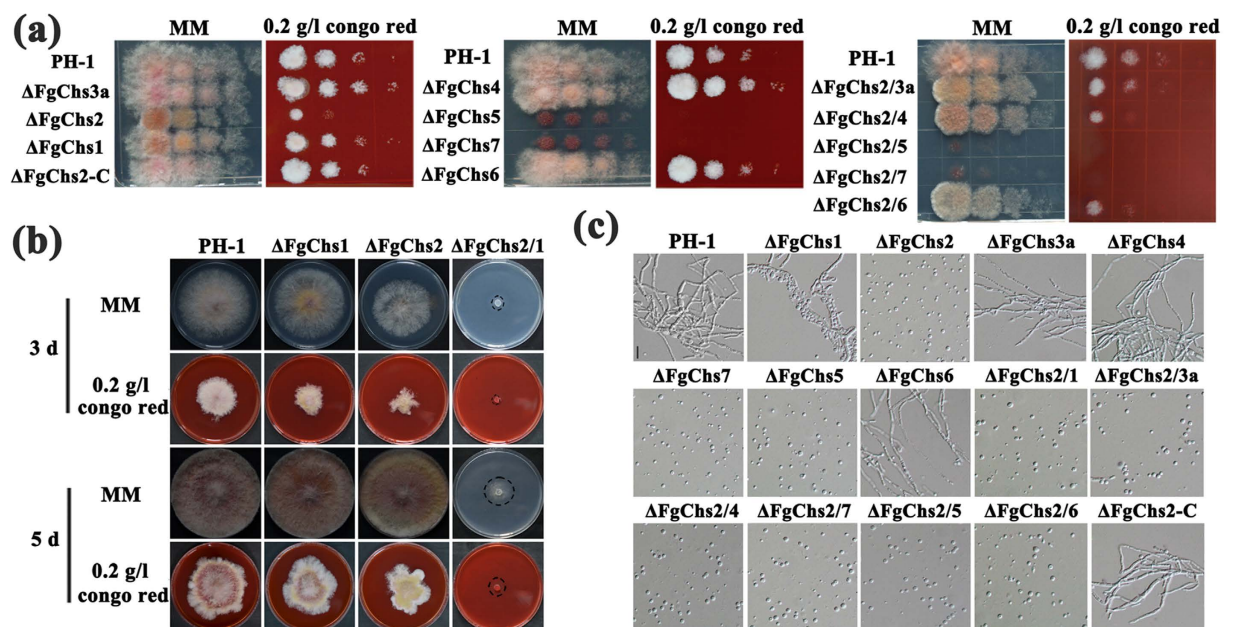


Figure 8. Sensitivity of *FgChs* single and double deletion mutants to cell wall stress agents. (a) Serial dilutions of conidial suspension of each strain were spotted on MM (CK) and MM supplemented with 0.2 g/l congo red (CR). All the plates were incubated at 25 °C for 3 days. (b) Mycelial plugs of PH-1, $\Delta FgChs1$, $\Delta FgChs2$ and $\Delta FgChs2/1$ were inoculated on MM (CK) and MM supplemented with 0.2 g/l CR. All the plates were incubated at 25 °C for 3 and 5 days. (c) After treatment with cellulase, lysozyme and snailase at 30 °C for 30 min, mycelia of the mutants $\Delta FgChs1$, $\Delta FgChs2$, $\Delta FgChs7$, $\Delta FgChs5$ and all the double deletion mutants were well digested and released abundant protoplasts. Bar = 10 μ m.

double mutant showed more disorganized than those of the *AnchsB* single mutant³². Double deletion of *AnchsmA* (VI) and *AnchsmB* (V) impeded the elongation of germ tubes or hyphae, whereas single deletion of any one rarely caused such defects²⁷. In *M. oryzae*, *MoChs5* (V) and *MoChs6* (VI) were also reported to have overlapping functions in maintaining polarized growth in vegetative tissue¹². Results of these studies indicated that Chss of different classes may co-regulate vegetative growth in filamentous fungi.

In *A. nidulans*, a previous study showed that the double disruption of *AnchsA* and *AnchsD* caused a severe defect in conidial formation although each single disruptant showed no obvious decrease in conidiation²⁸. The *AnchsC* and *AnchsA* double null mutant showed drastically reduced conidiophore population and occasionally produced secondary conidiophores²⁹. These studies indicated that class II Chs shares functions with class I and

IV Chss in regulating conidiation in *A. nidulans*. Similarly, in this study, the double mutant $\Delta FgChs2/1$ was unable to produce conidia and the mutant $\Delta FgChs2/4$ showed significantly less conidiation than those of each single gene deletion mutants $\Delta FgChs2$ and $\Delta FgChs4$ (Fig. 5a,b). Importantly, we found that *FgChs2* also has overlapping functions in conidiation with *FgChs7* and *FgChs5*, indicating that the class II *FgChs* can co-regulate conidiation with multiple classes of Chss in *F. graminearum*.

A previous study on *Wangiella dermatitidis* found that the double disruption of *WdCHS2* (I) and *WdCHS3* (III) caused marked virulence defects although the single gene mutants showed no loss of virulence³³. Zheng *et al.* (2006) reported that disruption mutant of both *WdCHS2* and *WdCHS1* (II) grew abnormally and almost lost virulence at 25 °C, while single gene disruption strains remained virulence as the wild type, indicating overlapping functions in virulence between these *Chs* genes³⁴. Functional overlap in virulence of Chss has also been found in our study. *FgChs2* shares functions in pathogenicity not only with *FgChs1* and *FgChs3a*, but also with *FgChs4*, *FgChs7*, *FgChs5* and *FgChs6*. In addition, our study found that all chitin synthases except *FgChs3a* and *FgChs6* are involved in regulating the production of DON. The reduced DON production in the *FgChs* deletion mutants further verify the involvement of *FgChss* in virulence since DON plays an important role in the extension of *F. graminearum* in plant tissue²³. In *M. oryzae*, *Mochs6* mutant was non-pathogenic, and *Mochs1* (III) and *Mochs7* (VII) single gene mutants were reported to cause only rare lesions on rice seedlings¹². In *B. cinerea*, *BcChs1* (I), *BcChs3a* (III), *BcChs6* (V) or *BcChs7* (VII) deletion mutant exhibited reduced virulence^{13–16}. Muszkieta *et al.* (2014) found that *AfcsmA* (VI) and *AfcsmB* (V) mutants of *Aspergillus fumigatus* were responsible for the virulence to *Galleria mellonella* and mouse³⁵. These studies indicate that Chss play an important role in virulence in pathogenic fungi.

In this study, we found that the mutant $\Delta FgChs2$ showed thickened and “wavy” septa occasionally with a larger central pore (Fig. 7). In *S. cerevisiae*, *C. albicans* and *W. dermatitidis*, class II Chss are also found to be responsible for septum formation^{7,34,36}, which is consistent with our finding. Unexpectedly, all the double mutants of *FgChs2* and other *FgChss*, especially $\Delta FgChs2/1$, showed more serious defects on septal morphology than the single mutants, indicating that all classes of *FgChss* are involved in septation. The *FgChs1 FgChs2* double mutant produced aberrantly thick septa with an abnormally large pore, which is very similar to those of the double mutant *AnchsC* and *AnchsA* of *A. nidulans*³⁰. However, $\Delta AnchsA$ and $\Delta AnchsC$ single mutants did not show different appearances in comparison with the wild type³⁰. In $\Delta AncsmB$ and $\Delta AncsmA$, the generation of intrahyphal hyphae was associated with the closing of septal pores, indicating that class V and VI chss in *A. nidulans* might be involved in the formation of septal pores³⁷. But in our study, the single deletion mutants $\Delta FgChs7$ and $\Delta FgChs5$ did not show obvious defects on septal morphology, moreover, the double mutants $\Delta FgChs2/7$ and $\Delta FgChs2/5$ did not exhibit additional defects on septal pores in comparison with $\Delta FgChs2$, which might indicate the functions in septum formation of classes V and VI Chss in *F. graminearum* may differ from those in *A. nidulans*.

Similar to our case, localization analysis of seven chitin synthases in *N. crassa* showed that all of them localize at septa indicating all Chss might be involved in septum formation^{38–40}. However, only the class VI heterokaryotic KA6 strain produced aberrant and particularly “wavy” septa in *B. cinerea*¹⁶. Characterization of all Chss of *M. oryzae* showed that only class III *Chs* deletion mutant displayed more than 90% of abnormal conidia without any septum¹². In *Fusarium oxysporum*, the *FochsVb* (V) single and *FochsV* (VI) *FochsVb* double mutants exhibited morphological abnormalities in septum formation and distribution³¹, whereas other *FoChs1* (I), *FoChs2* (II) and *FoChs7* (IV) single mutants showed similar septation with the wild type⁴¹. These studies strongly indicated that different classes of Chss may be responsible for septum formation in different filamentous fungi.

In *M. oryzae*, the *Mochs1 Mochs3* (I) double mutant exhibited increased susceptibility to high osmotic and oxidative stresses¹². In *A. nidulans*, the double disruptant of *AnchsC* and *AnchsA* showed high sensitivity to SDS, chitin-binding dyes and chitin synthase inhibitors, although the single *AnchsC* and *AnchsA* mutants did not show defects in responses to these stresses²⁹, indicating that *AnchsC* and *AnchsA* may have compensatory functions in responses to stresses. In contrast to what is seen in *A. nidulans* and *M. oryzae*, our study found that *FgChss* do not co-regulate the response to cell wall-damaging stress in *F. graminearum* (Fig. 8). These results indicate functions of Chss in stress responses are species-specific.

Experimental Procedures

Strains and culture conditions. *F. graminearum* strain PH-1 was used as the wild-type progenitor for the construction of *FgChs* deletion mutants. The wild type, resultant mutants and complemented strains generated in this study were grown on potato dextrose agar (PDA) (200 g potato, 20 g glucose, 20 g agar, and 1 l water) or minimal medium (MM) (10 mM K₂HPO₄, 10 mM KH₂PO₄, 4 mM (NH₄)₂SO₄, 2.5 mM NaCl, 2 mM MgSO₄, 0.45 mM CaCl₂, 9 mM FeSO₄, 10 mM glucose, and 1 l water, pH 6.9) for mycelial growth tests, carboxymethyl cellulose liquid medium (CMC; 15 g carboxymethyl cellulose, 1 g yeast extract, 0.5 g MgSO₄, 1 g NH₄NO₃, 1 g KH₂PO₄ and 1 l water) for conidiation tests, and 2% sugar water and yeast extract peptone dextrose liquid medium (YEPD; 1% yeast extract, 2% peptone, 2% dextrose, and 1 l water, pH 6.7) for conidial germination tests.

Generation of gene deletion and complementation mutants. The double-joint PCR approach⁴² was used to generate the gene replacement construct for each target gene (Fig. S1a,d). In briefly, the 5' and 3' flanking regions of each gene were amplified with the primer pairs listed in Table S1, and the amplified sequences were then fused with the appropriate resistance gene cassette. The resulting PCR products for each gene were transformed into protoplasts of the wild-type progenitor PH-1 respectively, as described previously^{23,43}. Hygromycin B (Calbiochem, La Jolla, CA) was added to a final concentration of 100 mg/l for transformant selection. When other *FgChs* gene deletion mutants were constructed in the *FgChs2* deletion background, the geneticin resistance gene cassette (*NEO*) was used as a second marker. In order to complement the *FgChs2* deletion mutant with the entire wild-type *FgChs2*, the entire *FgChs2* was inserted into pYF11 vector which contained *NEO* by the yeast homologous recombination approach⁴⁴.

Putative gene deletion mutants were identified by PCR assays with the relevant primers (Table S1), and were further analyzed by the Southern blotting assays (Fig. S1). DNA of each strain was extracted and then digested by the appropriate restriction endonucleases, as indicated in Fig. S1a,d. The probes used for each strain (Fig. S1a,d) were labeled with digoxigenin (DIG) using a High Prime DNA Labeling and Detection Starter kit II according to the manufacturer's instructions (Roche Diagnostics; Mannheim, Germany).

In this study, we totally obtained seven single mutants $\Delta FgChs2$, $\Delta FgChs1$, $\Delta FgChs3a$, $\Delta FgChs4$, $\Delta FgChs7$, $\Delta FgChs5$ and $\Delta FgChs6$, six double mutants of *FgChs2* and each of other *FgChss*, $\Delta FgChs2/1$, $\Delta FgChs2/3a$, $\Delta FgChs2/4$, $\Delta FgChs2/7$, $\Delta FgChs2/5$ and $\Delta FgChs2/6$, and one complemented strain $\Delta FgChs2-C$ (Fig. S1b,c,e,f). We failed to obtain *FgChs3b* mutant although we had obtained more than 100 ectopic transformants from 4 transformation experiments independently, which indicates that the deletion of this gene may be lethal. All of the mutants generated in this study were preserved in 15% glycerol at -80°C .

RNA extraction and quantitative real-time PCR (qRT-PCR). Total RNA of the wild type was extracted from mycelia grown in potato dextrose broth (PDB) and MM at 25°C for 2 days in the dark, and from germinating conidia in YEPD at 25°C for 6 hours, by using the TaKaRa RNAiso Reagent (TaKaRa Biotechnology Co., Dalian, China). Ten mg of each RNA sample was used for reverse transcription with a RevertAid H Minus First Strand cDNA Synthesis Kit employing the oligo(dT)₁₈ primer (Fermentas Life Sciences, Burlington, ON, Canada). The expression levels of *FgChs* genes were determined by qRT-PCR with the primers listed in Table S1. For each sample, PCR amplification with the primer pair Actin-F + Actin-R (Table S1) for quantification of transcription of *ACTIN* gene was performed as a reference. The expression level of each gene in each strain was calculated using the $2^{-\Delta\Delta C_t}$ method⁴⁵. The experiment was repeated three times.

To assay the expression levels of in *FgChs* genes in the *FgChs2* deletion mutant $\Delta FgChs2$, the wild type and $\Delta FgChs2$ were grown in PDB at 25°C for 2 days in the dark. To determine the expression levels of *TRI* genes, the wild type and deletion mutants were inoculated into mycotoxin synthetic (MS) medium⁴⁶ (0.5 g KH_2PO_4 , 0.6 g K_2HPO_4 , 0.017 g MgSO_4 , 1 g $(\text{NH}_4)_2\text{SO}_4$, 20 g glucose, 0.1 ml Vogel's trace elements stock solution and 1 l water) and cultured for 4 days at 25°C in the dark. RNA extraction and qRT-PCR were performed as described above. The experiment was repeated three times.

Growth and conidiation tests. To measure hyphal growth of each strain, mycelial plugs were taken from the edge of 3-day-old colony and placed on the center of PDA and MM plates at 25°C in the dark. After incubation for 3 days, colony diameter in each plate was measured in two perpendicular directions with the original mycelial plug diameter (5 mm) subtracted from each measurement. The experiment was repeated three times independently.

For conidiation assays, fresh mycelia (50 mg) of each strain were inoculated in a 50-ml flask containing 20 ml CMC liquid media. The flasks were incubated at 25°C for 4.5 and 15 days in a shaker (180 rpm). Then the conidial number in each flask was determined using a hemacytometer. Conidial morphology was observed with a Nikon ECLIPSE E100 microscope (Nikon Co., Tokyo, Japan). Furthermore, conidia (approximately 10 conidia/ μl) of each strain were incubated in 2% sugar water at 25°C for 4 hours, and conidium germination was examined under a Nikon ECLIPSE E100 microscope (Nikon Co., Tokyo, Japan). The experiment was repeated three times independently.

Pathogenicity assays. Pathogenicity of each strain on flowering wheat heads was performed as described previously⁴⁷. Briefly, a 10- μl aliquot of fresh conidial suspension was injected into a floret in the central section spikelet of single flowering wheat heads of susceptible cultivar Jimai 22 and the control heads were inoculated with 10- μl of sterilized water. Fifteen replicates were experimented for each strain. After inoculation, the plants were kept at $22 \pm 2^{\circ}\text{C}$ under 95–100% humidity with 12 h of daylight. After inoculating for 15 and 25 days, the infected spikelets of each inoculated wheat head were recorded. The experiment was repeated four times.

To further analyze the virulence defects of the mutants in details, penetration behavior of each strain was examined on cellophane membranes as described previously⁴⁸. Briefly, each strain was grown on minimal medium covered with a cellophane membrane. After 2 days of incubation, the cellophane membrane with the colony was removed from each plate. After the plates were incubated for two additional days, mycelial growth on each plate was examined. The presence of mycelial growth on the plate indicates penetration of the cellophane membrane. The experiment was repeated three times.

Determination of DON production. To determine DON biosynthesis, a 50-g aliquot of healthy wheat kernels was sterilized and inoculated with five mycelial plugs of each strain. After incubation at 25°C for 30 days, DON was extracted using a previously described protocol⁴⁹. The DON extracts were purified with PuriToxSR DON column TC-T200 (Trilogy analytical laboratory), and the amount of DON (per mg fungal DNA) in each sample was determined by using a HPLC system Waters 1525 (Waters Co., America)⁵⁰. Additionally, the amount of *F. graminearum* DNA of each sample was determined using qRT-PCR assays⁴⁷. The experiment was repeated three times, and data were analyzed using analysis of variance (SAS version 8.0; SAS Institute, Cary, NC).

Determination of sensitivity to cell wall stress agents. Serial dilutions of conidial suspension of each strain were spotted on MM amended with the cell wall-damaging agent CR (0.2 g/l). After the plates were incubated at 25°C for 3 days, the growth of each strain in each plate was examined. Give that the mutant $\Delta FgChs2/1$ could not produce conidium, we determined the sensitivity to CR using mycelial plugs. Mycelial plugs (5-mm in diameter) of each strain taken from the periphery of a 3- or 5-day-old colony were incubated on MM amended 0.2 g/l CR. After the plates were incubated at 25°C for 3 and 5 days, the growth of each strain in each plate was examined. Each experiment was repeated three times independently.

For each strain, fresh hyphae of each strain were harvested, and treated with cellulase, lysozyme and snailase (2% w/v, each) (Kaiyang Co., Shanghai, China) for 30 min in 0.7 M NaCl at 30 °C. The resulting protoplast of each strain was examined under a Nikon ECLIPSE E100 microscope (Nikon Co., Tokyo, Japan). Each experiment was repeated three times independently.

Transmission electron microscopy (TEM) assays. For the transmission electron microscopy assay, the 1.5-day-old mycelia cultured in PDB were fixed with 2.5% (v/v) glutaraldehyde. The specimens were dehydrated in a graded series of ethanol and embedded in Epon812. Ultrathin sections were cut with an ultramicrotome (LKB-V, Sweden), stained with uranyl acetate and lead citrate, and observed with an H-7650 transmission electron microscope (Hitachi, Japan).

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Author Contributions

Z.L., X.Z., X.L., C.F. and X.H. carried out the experiments. Y.Y. and Z.M. proposed the hypothesis, figured out strategy, designed experiments, supervised the project, and wrote the manuscript. All authors contributed to the data collection and analysis and the manuscript preparation.

Additional Information

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