The Stress Response Regulator AflSkn7 Influences Morphological Development, Stress Response, and Pathogenicity in the Fungus Aspergillus flavus

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Abstract: This study focused on AflSkn7, which is a stress response regulator in the aflatoxin-producing Aspergillus flavus. The AflAflSkn7 mutants exhibited partially defective conidial formation and a complete inability to generate sclerotia, indicating AflSkn7 affects A. flavus asexual and sexual development. The mutants tolerated osmotic stress but were partially susceptible to the effects of cell wall stress. Additionally, the AflAflSkn7 mutants were especially sensitive to oxidative stress. These observations confirmed that AflSkn7 influences oxidative stress responses rather than osmotic stress responses. Additionally, AflSkn7 was observed to increase aflatoxin biosynthesis and seed infection rates. These results indicate AflSkn7 affects A. flavus morphological development, stress response, aflatoxin production, and pathogenicity. The results of this study may facilitate the development of new methods to manage A. flavus infections.

Keywords: Aspergillus flavus; aflatoxins; development; stress response; pathogenicity

1. Introduction

Aspergillus flavus, a saprophytic soil fungus, causes preharvest and postharvest diseases in seed crops such as maize, peanut, and cottonseed [1]. This fungus produces the mycotoxin aflatoxin, which can affect the structure of DNA [2]. If animals or humans ingest aflatoxin-contaminated seeds, they may develop liver tumors [1]. Consequently, contamination of agricultural commodities by aflatoxin may result in considerable economic losses and serious health problems, especially in developing countries [2]. Aflatoxins are a group of toxic secondary metabolites that are mainly produced by two Aspergillus species (Section Flavi), namely A. flavus and A. parasiticus [3]. Aspergillus flavus can produce two aflatoxins, B₁ and B₂. Aflatoxin B₁ (AFB₁) is considered the most toxic aflatoxin, and its AFB₁ epoxides and AFB₁-exo-epoxides are often detected in hepatocellular carcinoma patients [1].

Previous studies on A. flavus revealed that approximately 30 genes clustered within a 70-kb genomic region near the telomere of chromosome 3 are involved in aflatoxin biosynthesis [1,4]. Most of these genes encode enzymes catalyzing one or multiple reactions in the aflatoxin biosynthetic pathway, while other genes encode regulator proteins. For example, aflR encodes a global regulatory protein required for transcriptional activation of most structural genes [5]. Additionally, many environmental factors regulate aflatoxin biosynthesis, including temperature, water activity, and oxidative stress [6–9]. Despite the crucial roles of environmental factors during aflatoxin biosynthesis,
how aflatoxin producers sense and cope with various environmental stresses is currently unknown. Therefore, characterizing the underlying mechanism may enable the development of improved methods to control *A. flavus* infections.

Histidine kinase-mediated phosphorelays, also known as two-component signaling systems, are common fungal pathways for sensing and adapting to environmental changes. The Sln1 pathway in *Saccharomyces cerevisiae* is one of the best-studied two-component pathways, and it primarily responds to osmotic stress [10]. Skn7 (Suppressor of kre nine 7) is responsible for another branch of the Sln1 pathway. In *S. cerevisiae*, Skn7 functions in response to oxidative and cell wall stresses using different mechanisms [11]. Following exposure to cell wall stress, the conserved aspartate residue of Skn7 is phosphorylated via Ypd1. In contrast, Skn7 is phosphorylated at a conserved threonine residue in the receiver domain in response to oxidative stress [12]. However, one study revealed that Skn7 is also involved in responses to osmotic stress [13]. Furthermore, there is considerable evidence linking Skn7 function to thermal stress adaptation, fungicide sensitivity, sexual mating, autolysis, sporulation, and virulence in other fungi [10,14].

Therefore, the main objective of this study was to characterize *A. flavus* Skn7 functions regarding morphological development, stress responses, and pathogenicity. A thorough characterization of Skn7 functions may facilitate the development of new methods to control *A. flavus* infections. ∆*AflSkn7* deletion mutants were generated, and the mutant phenotypes related to stress and virulence were analyzed.

2. Results

2.1. *Aspergillus Flavus* Skn7 Sequence Analysis

The sequence of the *A. flavus* putative response regulator Skn7 (AFL2G_02624) was retrieved from the Broad Institute database. Based on the annotation, the *AflSkn7* open reading frame (ORF) consists of 2851 bp and is interrupted by four introns (76, 60, 62, and 59 bp). The encoded amino acid sequence is highly similar to orthologous sequences from other fungi, and is most similar to *Aspergillus oryzae* Skn7 (Figure 1A). Sequence analysis revealed that *AflSkn7* contains two conserved domains: an N-terminal heat-shock transcription factor-like DNA-binding domain and a C-terminal CheY-like receiver domain (Figure 1B). These two domains are common among Skn7-like proteins from other fungi, suggesting *AflSkn7* regulates stress responses in *A. flavus*.
2.2. Construction of ΔAflSkn7 Deletion Mutants

To confirm the biological function of Skn7 in *A. flavus*, ΔAflSkn7 deletion mutants were constructed using a homologous recombination strategy (Figure 2A). Knockout mutants were screened from uridine autotrophy transformants by polymerase chain reaction (PCR) analysis. Two deletion mutants (i.e., *skn7*-5 and *skn7*-11) were identified with two PCR primer pairs, which amplified fragments that corresponded to the ORF and 5' flanking regions of *AflSkn7*, respectively (Figure 2B). To avoid false positives, PCR primers (i.e., AflSkn7UUF/AflSkn7DDR) matched to the upstream and downstream regions of fusion fragment were used to verify the transformants. When PCR products were sequenced, the alignment results indicated 100% identity between the sequenced products and the expected sequences (Figure S1). The PCR and sequencing results indicated that *AflSkn7* was exactly replaced by *pryG* in knockout mutants. Additionally, reverse transcription (RT)-PCR using primers specific for the ORF region confirmed *AflSkn7* was not expressed in deletion mutants (Figure 2C), suggesting *AflSkn7* gene was disrupted in *skn7*-5 and *skn7*-11 strains. Furthermore, the stable and identical phenotype of the two mutants as following suggested that changes in these mutants were caused by the inactivation of *AflSkn7* in *A. flavus*.

**Figure 1.** Sequence and domain analysis of Skn7 proteins from different fungi. (A) Phylogenetic analysis of Skn7 proteins from 28 fungal species. Bootstrap values were calculated using the neighbor-joining method with 1000 replicates and are indicated at each node. All clades with *Aspergillus* species are highlighted in red, and *Aspergillus flavus* is indicated in blue point. (B) Schematic diagrams of Skn7 proteins from different fungi. Gray parts correspond to non-conserved regions. Pink parts indicate the heat-shock transcription factor-like DNA-binding domain. Green parts represent the CheY-like receptor domain. The scale corresponds to protein length.

**Figure 2.** Gene deletion method and confirmation. (A) Schematic illustration of the gene deletion method. The numbers indicate the length of the corresponding gene. (B) Polymerase chain reaction (PCR) confirmation of ΔAflSkn7 gene deletion. ORF, *AflSkn7* coding region amplified by PCR; UA, 5' region of *AflSkn7* amplified by PCR. (C) Reverse transcription-PCR confirmation of ΔAflSkn7 gene deletion.
2.3. Involvement of AflSkn7 during Asexual Development

The growth rates of the wild-type fungus and ∆AflSkn7 deletion mutants were similar on potato dextrose agar (PDA) medium at 28 °C (Figure 3A). In contrast, there were considerable differences in the production of conidia on PDA medium (Figure 3A), with the wild-type fungus producing approximately five-fold more conidia than the mutants (Figure 3B). Microscopic examinations indicated ∆AflSkn7 mutants produced fewer conidiophores and less aerial hyphae than the wild-type fungus (Figure 3C). Additionally, the mutant conidiophores were smaller than those of the wild-type fungus, suggesting the loss of Skn7 might delay conidiation. These findings suggested AflSkn7 was involved in A. flavus asexual development.

![Figure 3](image)

**Figure 3.** Effects of AflSkn7 on conidial formation: (A) colony morphology of strains during conidiation; (B) comparison of conidial production among wild-type fungus and two mutants; and (C) microscopic analysis of conidial structures (magnification: ×200). Conidiophores for each strain are indicated by white arrows. All strains were cultured on potato dextrose agar medium at 28 °C for seven days. Asterisks indicate significant differences from the results of the wild-type fungus (*** p ≤ 0.001).

2.4. Requirement of AflSkn7 during Sexual Development

Considering it affects asexual development, the potential roles of AflSkn7 during sexual development were investigated. After culturing fungal strains on yeast extract-sucrose (YES) medium at 37 °C for seven days, all strains were washed with 75% ethanol. There were almost no morphological differences between the wild-type and mutant strains prior to the ethanol wash (Figure 4A, left). After the wash, the wild-type fungus was able to produce sclerotia on YES medium (Figure 4A, top), while both ∆AflSkn7 mutants exhibited severely limited sclerotial production (Figure 4A, middle and bottom). A quantitative analysis revealed that the ∆AflSkn7 mutants produced far fewer than the wild-type strain (Figure 4B). A quantitative RT-PCR (qRT-PCR) method was used to analyze the expression of nsdC and nsdD, which influence sclerotial development. The nsdC expression level in the wild-type fungus was at least two-fold higher than that of the ∆AflSkn7-5 mutant (Figure 4C, left). Additionally, the nsdD expression level was at least four-fold higher in the wild-type strain than in the ∆AflSkn7-5 mutant (Figure 4C, right). These data indicated that AflSkn7 had a considerable effect on A. flavus sexual development.
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were significantly different between the wild-type fungus and the mutants, suggesting the cell wall

2.5. Effects of the Deletion of AflSkn7 on Sensitivity to Osmotic and Cell Wall Stresses

∆ components of the Congo red (Figure 5B). The expression levels of fsk1 mutants. Furthermore, the adverse effects caused by calcofluor white were greater than those of the wild-type fungus (Figure 5B). In contrast, both agents strongly inhibited the growth of ∆AflSkn7 mutants. Additional information provided on the right side. (A) Colony morphology of strains with sclerotia. Enlarged image of each plate is provided on the right side. (B) Abundance of sclerotia produced by the wild-type fungus and two mutants. (C) Expression analysis of two genes related to sclerotial formation in the wild-type fungus and two mutants. All strains were cultured on YES medium at 37 °C. After incubating for seven days, cultures were washed with 75% ethanol. The 2^−ΔΔCt method was used to evaluate of target gene expression levels, which were relative to that of the β-actin reference gene. Asterisks indicate significant differences from the results of the wild-type fungus (* p ≤ 0.05; ** p ≤ 0.01; *** p ≤ 0.001).

Figure 4. Effects of AflSkn7 on sclerotial formation. (A) Colony morphology of strains with sclerotia. Enlarged image of each plate is provided on the right side. (B) Abundance of sclerotia produced by the wild-type fungus and two mutants. (C) Expression analysis of two genes related to sclerotial formation in the wild-type fungus and two mutants. All strains were cultured on YES medium at 37 °C. After incubating for seven days, cultures were washed with 75% ethanol. The 2^−ΔΔCt method was used to evaluate of target gene expression levels, which were relative to that of the β-actin reference gene. Asterisks indicate significant differences from the results of the wild-type fungus (* p ≤ 0.05; ** p ≤ 0.01; *** p ≤ 0.001).

2.5. Effects of the Deletion of AflSkn7 on Sensitivity to Osmotic and Cell Wall Stresses

Skn7 responds to osmotic and cell wall stresses in many fungi. In this study, the sensitivity of A. flavus to different osmoticants and cell wall-damaging agents was examined. Under osmotic stress conditions, the mycelial growth rate of all A. flavus strains decreased. However, disruption of the AflSkn7 gene did not result in increased sensitivity to NaCl and KCl (Figure 5A). To analyze the sensitivity of the ΔAflSkn7 mutants to cell wall stress, all strains were grown on YES medium supplemented with Congo red and calcofluor white. Both agents subtly affected the radial growth of the wild-type fungus (Figure 5B). In contrast, both agents strongly inhibited the growth of ΔAflSkn7 mutants. Furthermore, the adverse effects caused by calcofluor white were greater than those of Congo red (Figure 5B). The expression levels of fsk1 and och1, which are cell wall synthesis genes, were significantly different between the wild-type fungus and the mutants, suggesting the cell wall components of the ΔAflSkn7 mutants were disordered (Figure 5C). Therefore, AflSkn7 responds to cell wall stress, but not to osmotic stress in A. flavus.
wild-type and mutant strains were greater for the superoxide dismutase gene than for the glutathione reductase gene. These observations implied that AflSkn7 has a critical role during responses to oxidative stress.

2.6. Effects of the Deletion of AflSkn7 on Sensitivity to Oxidative Stress

Oxidative stress is another important environmental stimulus for fungi. There is abundant evidence indicating the secondary metabolism and pathogenicity of filamentous fungi are affected by oxidative stress [15]. In this study, A. flavus strains lacking AflSkn7 were hypersensitive to hydrogen peroxide (H₂O₂) and tert-butyl-hydrogen peroxide (t-BOOH) (Figure 6A). Mutant growth was completely inhibited by 6 mM H₂O₂. In contrast, growth of the wild-type fungus was unaffected even at 6 mM H₂O₂ (Figure 6B). The expression levels of two genes encoding antioxidants (i.e., Cu, Zn-superoxide dismutase: AFLG_10810 and glutathione reductase: AFLG_01044) were higher in the wild-type fungus than in the two mutants (Figure 6C). The expression level differences between the wild-type and mutant strains were greater for the superoxide dismutase gene than for the glutathione reductase gene. These observations implied that AflSkn7 has a critical role during responses to oxidative stress.
Toxins with the results of strains cultured in potato dextrose broth (PDB) broth (Figure S2). Quantitative wild-type All evidence indicated that AflSkn7 increased aflatoxin biosynthesis in (Figure 7B). To confirm this observation, the expression levels of two aflatoxin biosynthesis genes (genes in the afl and analyses of both aflatoxins. Therefore, the role of AflSkn7 for aflatoxin biosynthesis was investigated. Only the

2.7. Role of AflSkn7 for Aflatoxin Biosynthesis

As previously mentioned, the virulence of A. flavus is partially based on the production of aflatoxins. Therefore, the role of AflSkn7 for aflatoxin biosynthesis was investigated. Only the wild-type A. flavus strain produced aflatoxins in YES broth at 28 °C (Figure 7A). This was consistent with the results of strains cultured in potato dextrose broth (PDB) broth (Figure S2). Quantitative analyses of both ΔAflSkn7 mutants indicated they were unable to produce aflatoxins in YES broth (Figure 7B). To confirm this observation, the expression levels of two aflatoxin biosynthesis genes (aflJ and aflQ) were analyzed in all strains. As expected, the transcription levels of the aflatoxin biosynthesis genes in the ΔAflSkn7 mutants were considerably lower than that in the wild-type strain (Figure 7C). All evidence indicated that AflSkn7 increased aflatoxin biosynthesis in A. flavus.

Figure 6. Effects of oxidative stress on wild-type (WT) fungus and two mutants (ΔSkn7-5 and ΔSkn7-11): (A) colony morphology of strains exposed to different oxidative stresses; and (B) quantitative reverse transcription polymerase chain reaction analysis of antioxidant gene expression. The 2^−ΔΔCt method was used to evaluate of target gene expression levels, which were relative to that of the β-actin reference gene. Asterisks indicate significant differences from the results of the wild-type fungus (** p ≤ 0.01; *** p ≤ 0.001).

Figure 7. Cont.
2.8. Contribution of AflSkn7 to Pathogenicity

Whether AflSkn7 affects the ability of *A. flavus* to colonize seeds and produce aflatoxin was investigated. The wild-type and mutant strains were still able to infect maize kernels seven days after inoculation. However, the wild-type fungus grew better on kernels than the ΔAflSkn7 mutants, and also produced more conidia (Figure 8A,B). To examine the consequences of fungal infections of seeds, aflatoxin levels on contaminated seeds were determined. Similar to the results observed for the fungal strains grown in YES broth, the wild-type fungal strain produced more aflatoxin on infected kernels than the ΔAflSkn7 mutants (Figure 8C). These results indicated AflSkn7 affected not only *A. flavus* colonization, but also aflatoxin biosynthesis. Thus, AflSkn7 likely influences *A. flavus* pathogenicity.

![Figure 8](image_url)

**Figure 8.** Effects of deletion of AflSkn7 on fungal pathogenicity on maize: (A) maize kernels were treated with water (control), wild-type fungus (WT), or two AflSkn7 deletion mutants (Δskn7-5 and Δskn7-11) and cultured at 28 °C for seven days; (B) quantitative analysis of conidia collected from infected maize kernels; and (C) thin-layer chromatography analysis of aflatoxin production in infected maize kernels. Asterisks indicate significant differences from the results of the wild-type fungus (*** p ≤ 0.001).
3. Discussion

Skn7 and its orthologs are widely distributed in fungi, suggesting they have important roles influencing fungal survival. The results presented herein are consistent with those of a previous study, which reported that an *Aspergillus nidulans* Skn7 mutant (i.e., ΔSrrA) exhibits partly defective conidiation [16]. Another study confirmed that SrrA considerably affects the expression of brlA during conidial formation [17]. In *Alternaria alternata*, the defective conidia observed in Skn7 mutants may be the result of inhibited cell wall biosynthesis [18]. In contrast to the results of the current study, the *Metarhizium robertsi* Skn7 mutant is completely unable to produce conidia [14], while *A. fumigatus* Skn7 is likely not involved in regulating the production of conidia [19]. These results indicate that Skn7 regulates conidial formation in a species-specific manner.

In addition to the effects of Skn7 on the production of conidia, this study determined that AflSkn7 positively influences the formation of sclerotia, suggesting Skn7 facilitates *A. flavus* sexual development. However, Skn7 did not affect sexual development in *Fusarium graminearum* [20], while *Cryptococcus neoformans* Skn7 inhibited sexual reproduction [21]. Interestingly, a *Botrytis cinerea* Skn7 mutant produced more sclerotia than the wild-type fungus, although they were smaller [22]. Additionally, similar to the results of this study, the *Schizosaccharomyces pombe* Skn7 (i.e., Prr1) positively affected sexual development [23]. In *S. pombe*, Prr1 induces Ste11 expression, which is essential for sexual development [23]. In *A. nidulans*, the SteC gene, which is a homolog of *S. pombe* Ste11, is important for sexual development [24]. There is currently no conclusive evidence that Skn7 regulates sexual development in the model filamentous fungus *A. nidulans*.

An important function of Skn7 in *A. flavus* and other fungi is its response to oxidative stress [17]. Wild-type *A. nidulans* can tolerate up to 2 mM H$_2$O$_2$, while the ΔsrrA mutants are unable to grow in the presence of 1 mM H$_2$O$_2$ [16]. Similarly, the *A. fumigatus* Skn7 mutant is hypersensitive to oxidative stress mediated by H$_2$O$_2$ [19]. Skn7 is believed to induce the expression of several antioxidant genes that provide resistance to H$_2$O$_2$, including genes for catalase, superoxide dismutase, and thioredoxin [16, 25]. These results are consistent with the observations described herein, which indicated that AflSkn7 mutants are hypersensitive to oxidative stress and exhibit decreased antioxidant gene expression levels. In contrast with the findings of the studies mentioned above, in the insect-pathogenic fungus *Metarhizium robertsii*, Skn7 did not influence responses to oxidative stress [14]. Likewise, in the rice pathogen *Magnaporthe oryzae*, sensitivity to oxidative stress was not altered in Skn7 mutants, suggesting other signal transduction systems may be involved in responses to oxidative stress [26].

A role for Skn7 in responses to cell wall stress was also noted. The AflSkn7 mutants were more sensitive to cell wall damaging agents than the wild-type fungus. Congo red and calcofluor white decrease β-glucan levels and chitin accumulation in cell walls, respectively [17]. In ΔMrSkn7 mutants, the accumulation of glucan and chitin components is inhibited. Additionally, the expression levels of genes associated with cell wall biosynthesis are significantly lower in the mutants than in the wild-type fungus [14]. Genes that regulate cell wall biosynthesis usually play a role in cell wall integration (CWI) signaling. The fact that Skn7 interacts with guanosine triphosphate (GTP)-bound Rho1, which is considered to be the main regulator of CWI signaling in yeast, strongly suggests Skn7 has a role in CWI signaling [27]. Therefore, the Rho1-mediated Slt2 mitogen-activated protein kinase pathway may influence AflSkn7 responses to cell wall stress. Additionally, the ChaAP1/Skn7 *Cochliobolus heterostrophus* double mutant is more sensitive to cell wall stress than either of the single mutants [11]. Thus, AflSkn7 mutants may become more sensitive to wall stress following the deletion of AflAP1. The current study revealed that *A. flavus* Skn7 mutants were insensitive to osmotic stress. However, the radial growth of the *A. nidulans* ΔsrrA mutant decreased slightly following exposure to osmotic stress conditions [16]. It is possible that the stress response regulators Skn7 and Ssk1 may periodically have overlapping roles [10]. The overlapping functions of Skn7 and Ssk1 have been observed in *M. oryzae* [26]. Additional research is required to determine the exact relationship between Skn7 and Ssk1 in *A. flavus*. 
Aflatoxin biosynthesis in filamentous fungi is induced and intensified by intracellular oxidative stress [15]. However, Roze and co-workers reported that aflatoxin biosynthesis resulted in the production of secondary reactive oxygen species (ROS) potentially able to provide resistance to oxidative stress [28]. In *A. parasiticus*, *ApyapA* (a yeast *yap1* homolog) controls the expression levels of several oxidative stress response genes involved in scavenging ROS [29]. The disruption of *ApyapA* led to a precocious ROS formation and aflatoxin biosynthesis [29]. Similar results were observed in *A. ochracus*, in which ochratoxin A biosynthesis is regulated by the *yap1*-homolog *Ao yap1* under oxidative stress condition [30]. Additionally, *Skn7* is associated with *Yap1* in yeast responses oxidative stress [12], and Hagiwara and co-workers proposed that *SrrA* and *NapA* (*yap1* homolog) formed a complex to activate antioxidant genes in *A. nidulans* [17]. A study on the promoters of aflatoxin genes and stress–response genes revealed that the *SrrA* and *Yap1* binding-sites were located close together in the promoters [31]. Therefore, *SrrA* may interact indirectly with *Apyap1* to activate the expression of aflatoxin genes. The current study has for the first time provided direct evidence that *Skn7* is very important for regulating aflatoxin biosynthesis. As complementary evidence, deoxynivalenol production by *F. graminearum* is lower in *FgSkn7* mutant strains than in wild-type fungus [20]. Furthermore, deoxynivalenol biosynthesis is upregulated when *F. graminearum* is exposed to oxidative stress [20].

The effect of *Skn7* on pathogenicity is highly variable among plant pathogens. The pathogenicity of an *M. oryzae Skn7* mutant on rice is similar to that of the wild-type strain [26], and the *FgSkn7* mutant maintains full virulence on wheat [20]. However, deleting *Skn7* from *A. alternata* significantly influences virulence, and the contribution of *AaSkn7* on pathogenicity is closely linked to fungal antioxidant activities [18]. The ability of pathogens to overcome oxidative stress conditions created by a plant host is critical for successful colonization [32]. This is supported by the fact the ∆*AflSkn7* mutants analyzed in this study simultaneously inhibited antioxidant synthesis and colonization of maize (Figures 6B and 8A). The efficiency of antioxidative activities is unexpectedly significantly decreased in a *B. cinerea Skn7* mutant, but the pathogen is just as virulent on tomato and cucumber leaves as the wild-type strain [13]. However, pathogenic fungi must overcome several obstacles, including osmotic stress and wall stress, to colonize in plant hosts. For example, *M. oryzae* must adapt to osmotic stress during infection [26]. Thus, the inhibited ability of ∆*AflSkn7* mutants to adapt to cell wall stress may restrict their ability to infect plants. The ∆*AflSkn7* mutants produced little aflatoxin and were weakly pathogenic on maize kernels (Figure 8A,C). A low aflatoxin producer usually exhibits a low germination rate, which may also influence the pathogenicity of a mutant [28]. In summary, the roles of *Skn7* during development, aflatoxin production, adaptation to environmental stresses, and resistance to host defense mechanisms may affect pathogen virulence [1].

4. Materials and Methods

4.1. Fungal Strains and Growth Conditions

The *Aspergillus flavus* CA14 wild-type (∆*ku70*, ∆*niaD*) and recipient (∆*ku70*, ∆*pyrG*, ∆*niaD*) strains were obtained from Dr. Ana M. Calvo, Northern Illinois University, DeKalb, IL, USA. The fungi were cultured on YES medium (20 g yeast extract, 150 g sucrose, 1 g MgSO₄·7H₂O, and 1 L water) at 28 °C. Fungal strains were grown on PDA (Difco) under light for 5 days to produce conidia. Fungi were grown in yeast glucose trace element (YGT) broth (5 g yeast extract, 20 g glucose, 1 mL trace elements, and 1 L water) at 37 °C on a rotary shaker for mycelial growth, and strains were grown on potato dextrose agar (YPD) medium (10 g yeast extract, 20 g peptone, 20 g glucose, 20 g agar, and 1 L water) to produce sclerotia. Czapek-Dox medium (Difco) supplemented with 3% sucrose was used for mutant selection.
4.2. Sequence Analysis

The AflSkn7 (AFL2G_02624) sequence was retrieved from the Broad Institute database [33]. Sequences of other Skn7 orthologs were downloaded from the National Center for Biotechnology Information database using the BLASTP algorithm with the AflSkn7 protein sequence as a query. Domains were predicted using the Simple Modular Architecture Research Tool [34] and drawn with the Illustrator for Biological Sequences program [35]. Sequences were aligned using ClustalW, and a phylogenetic tree was generated with MEGA5.1 (Center for Evolutionary Medicine and Informatics, Tempe, AZ, USA, 2012).

4.3. Gene Deletion

Gene knockout mutants were generated using homologous recombination [36]. For the homologous fragments, the 5' and 3' regions of AflSkn7 (921 and 1043 bp, respectively) were amplified with the primer pairs AflSkn7UF/AflSkn7UR and AflSkn7DF/AflSkn7DR (Table S1), which contain sequences that overlap the marker gene. For the marker gene, pyrG was amplified with the primer pair pyrGF/pyrGR (Table S1). The resulting PCR products were purified and linked by overlapping PCR with the primer pair AflSkn7UF/AflSkn7DR. The fusion PCR products were purified and inserted into recipient strain protoplasts using a published procedure [33]. Putative transformants were verified by diagnostic PCR (using primer pairs AflSkn7UF/pyrGR and AflSkn7F/AflSkn7R) and RT-PCR (using primer pairs AflSkn7RTF and AflSkn7RTR) (Table S1). Putative mutants were further verified with one PCR primer pair (AflSkn7UUF/AflSkn7DDR), and PCR products were sequenced.

4.4. Molecular Manipulations

Fungal genomic DNA was extracted from mycelia grown in YGT broth at 37 °C for 24 h with shaking (180 rpm). Genomic DNA was isolated using lysis buffer (400 mM Tris-HCl, 60 mM EDTA, 150 mM NaCl, and 1% SDS). Briefly, mycelia were harvested, mixed with 500 µL lysis buffer, and incubated at 60 °C for 20 min. Potassium acetate (200 µL) was added to the mixture. After gentle mixing, the sample was centrifuged at 12,000 × g for 10 min. The supernatant was gently mixed with 500 µL isopropanol (i.e., slow inversions). The resulting DNA pellet was collected by centrifugation at 12,000 × g for 2 min. The pelleted DNA was washed with 1 mL 70% ethanol, air-dried for 5 min, and suspended in 30 µL TE buffer (400 mM Tris-HCl and 60 mM EDTA). TRIzol reagent (Bioteck, Beijing, China) was used to isolate RNA according to the manufacturer’s instructions.

4.5. Stress Sensitivity Assay

Conidia from each fungal strain were cultured on PDA under light at 37 °C for 5 days. Conidia were harvested by washing the PDA surface with sterile distilled water containing 0.05% Tween 20 and 7% DMSO (Sinopharm Chemical Reagent, Shanghai, China), and then counted using a hemocytometer. A conidial suspension (10^4) was used to inoculate YES medium supplemented with 100 µg/mL calcofluor white (Sigma, St. Louis, MO, USA), 100 µg/mL Congo red (Sigma, St. Louis, MO, USA), 1 M NaCl, or 1 M KCl. Different concentrations of H₂O₂ were added to the YES medium for the oxidative stress assay. All plates were incubated in darkness at 37 °C for 5 days, and colonies were photographed.

4.6. Aflatoxin Analysis

Approximately 10^6 conidia of each strain were used to inoculate 25 mL YES broth or PDB (supplemented with zinc and molybdenum ions) in 100-mL flasks, which were then incubated at 28 °C with shaking at 180 rpm. After 6 days, 25 mL chloroform was added to each flask, and cultures were incubated for an additional 1 h. A 10-mL aliquot from the chloroform layer was transferred to a clean tube and evaporated to dryness at 70 °C. The solid residue was resuspended in 100 µL chloroform, and a 20-µL sample was spotted onto a thin-layer chromatography plate (Si250F; Jindao, Qingdao,
China). The plate was then treated with a chloroform:acetone (95:5, v/v) solution and viewed under UV light. Aflatoxins were quantified using a high-performance liquid chromatography system with a UV light detector. A solid residue sample was resuspended in methanol, and a 15-µL suspension was injected into a reversed-phase (C-18 column) high-performance liquid chromatography system (Waters, Milford, MA, USA). The solvent consisted of water:methanol:acetonitrile (56:22:22, v/v/v) at a flow rate of 1 mL/min.

4.7. Quantitative Reverse Transcription Polymerase Chain Reaction Analysis

Extracted RNA (2 µg) was treated with DNase I (Thermo Fisher Scientific, Waltham, MA, USA) and then reverse transcribed to cDNA using the Revert Aid First-strand cDNA Synthesis kit (Thermo Fisher Scientific, Waltham, MA, USA). The qRT-PCR was conducted using the SYBR Green Premix kit (Takara, Dalian, China) and the PikoReal 96 Real-Time PCR system (Thermo Scientific, Vantaa, Finland) following the manufacturer’s instructions. Target gene expression levels were evaluated using the $2^{-\Delta\Delta C_{\text{T}}}$ method [37]. The primers used to amplify target genes are listed in Table S1. The β-actin gene was selected as the reference gene.

4.8. Pathogenicity Assay

Pathogenicity tests were completed using maize seeds [38]. Embryos were removed to prevent germination, and endosperms were wounded by toothpicks to provide an infection site. Wounded kernels were surface-sterilized with 0.05% sodium hypochlorite for 3 min and 75% ethanol for 1 min. Sterilized seeds were rinsed three times with sterile water, and then added to 50-mL sterilized flasks. The seeds were inoculated with $10^7$ conidia, which were suspended in 15 mL sterile distilled water. After shaking at 180 rpm at 28 °C for 30 min, inoculated kernels were transferred to moistened filter paper in Petri dishes, and incubated at 28 °C. Kernels were harvested after 7 days to count spores and quantify aflatoxins.

Supplementary Materials: The following are available online at www.mdpi.com/2072-6651/8/7/202/s1, Figure S1: The alignment between the sequenced products and the expected sequences. The skn7-5 and skn7-11 indicated the sequenced products, while the pyrG indicated the expected sequences. Sequences marked with red and yellow letters were two primer pairs (i.e., AflSkn7DR and AflSkn7UF, respectively), which were used for amplifying fusion fragments. Figure S2: Thin-layer chromatography analysis for aflatoxin production in the indicated strains. Each strain was inoculated into 25 mL of PDB broth (supplemented with Zinc and Molybdenum ions) in a 100 mL flask and incubated with shaking at 180 r/min at 28 °C. After 6 days, the aflatoxin was extracted using a chloroform method. ST, aflatoxin B1; WT, the wild type; ΔSkn7-5 and ΔSkn7-11, two AflSkn7 deletion mutants. Table S1: Primers used in this study.

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