The Mitogen-Activated Protein Kinase CgMK1 Governs Appressorium Formation, Melanin Synthesis, and Plant Infection of Colletotrichum gloeosporioides

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The fungus Colletotrichum gloeosporioides infects plant hosts with a specialized cell called an appressorium, which is melanized and required for plant cell wall penetration. Here, we show that the mitogen-activated protein kinase CgMK1 governs appressorium formation and virulence in the poplar anthracnose fungus C. gloeosporioides. Deletion of CgMK1 impairs aerial hyphal growth and biomass accumulation, and CgMK1 is responsible for the expression of melanin biosynthesis-associated genes. CgMK1 deletion mutants are unable to form appressorium and lose the capacity to colonize either wounded or unwounded poplar leaves, leading to loss of virulence. We demonstrate that the exogenous application of cAMP fails to restore defective appressorium formation in the CgMK1 deletion mutants, suggesting that CgMK1 may function downstream or independent of a cAMP-dependent signal for appressorium formation. Moreover, CgMK1 mutants were sensitive to high osmosis, indicating that CgMK1 plays an important role in stress response. We conclude that CgMK1 plays a vital role in regulating appressorium formation, melanin biosynthesis, and virulence in C. gloeosporioides.

Keywords: Colletotrichum gloeosporioides, MAPK, appressorium formation, virulence, penetration

INTRODUCTION

Various genes in phytopathogenic fungi are involved in certain signaling pathways associated with the recognition of chemical substances from the host and in response to other extracellular stimuli. The mitogen-activated protein kinase (MAPK) signaling pathway is a downstream cellular signaling event that is remarkably conserved and mainly consists of MAPK cascades, including three-tiered protein kinase modules by which phosphorylation sites of tyrosine and threonine deliver signals, ultimately activating the downstream components of the pathway (Marshall, 1994; Seger and Krebs, 1995; Hamel et al., 2012). MAPK signaling pathways are divided into five categories that are classified according to the different functions of Saccharomyces cerevisiae, including conditioning the response of mating pheromones, controlling homeostasis under high osmolarity stress, and maintaining cell wall integrity (Herskowitz, 1995). The interplay of these signaling pathways in certain external stress conditions has previously been illustrated (Baltanás et al., 2013). Among the five types of MAPK signaling pathways, the functions of a Fus3/Kss1-related gene in phytopathogenic fungi include, in addition to filamentous growth, a vital role in...
virulence and in host surface colonization and the formation of infection structures, as well as penetration and development into the cell tissues of the host.

Fus3/Kss1 orthologs are denominated as "pathogenic MAPKs," which are vital genes that are significant for infection structure differentiation and various processes in pathogenic fungi. Increasing evidence has indicated that the sensor genes orchestrate virulence via Fus3/Kss1 orthologs, such as SHO1 and MSB2 (Leroch et al., 2015). Various components that converge downstream of fus3/Kss1 have also been identified, such as STE12 (Park et al., 2002) and SLF1 (Li et al., 2011). Some genes associated with other signaling pathways are also implicated in fus3/kss1-dependent pathways such as the inactivation of the TOR pathway (Marroquin-Guzman and Wilson, 2015), as well as the thioredoxin-associated gene, TRX2 (Wang et al., 2016). Previous studies have shown that the activation of fus3/kss1-associated genes is necessary for the development of a unique infection structure for endangering hosts in phytopathogenic fungi. For instance, in Magnaporthe oryzae, deletion of PMK1 (Fus3/Kss1 ortholog) disrupts the capacity of appressoria differentiation, thereby resulting in non-pathogenicity (Xu and Hamer, 1996). Additionally, strains lacking PMK1 are also prevented from attacking the host via wounded tissues, suggesting that PMK1 is vital for various invasive stages of fungal attack toward the host. Subsequently, ortholog analysis has demonstrated that Fus3/Kss1 is essential for pathogenicity in phytopathogens, not only in appressorium-forming fungi, such as CMK1 in Colletotrichum orbiculare (Takano et al., 2000) and ChMK1 in C. higginsianum (Wei et al., 2016), but also in non-classical appressorium-forming fungi such as VMK1 in Verticillium dahliae (Rauyaree et al., 2005) and CsFUS3 in Bipolaris sorokiniana (Leng and Zhong, 2015). Indeed, these MAPKs have been verified to be closely associated with the infection morphology and pathogenicity of fungi.

Anthracnose, a typical poplar foliage disease generated by the hemibiotrophic ascomycete fungus Colletotrichum gloeosporioides, is highly prevalent in northeast China and results in substantial economic losses. During pathogenesis process, an infectious structure characterized by the appearance of dome-like structures called forming at the hyphal apex (Kleemann et al., 2008). Previous studies have explored the significance of MAPK components and the cAMP/PKA pathway in appressorium formation in C. gloeosporioides. The functions of certain MAPK modules have been reported, such as the deletion of Cgl-SLT2 results in deficient sporulation, disruption of appressorium formation, and a reduction in pathogenicity in mango (Yong et al., 2013). CgMEK1 deletion mutants exhibit defective in appressoria development and the loss of virulence (Kim et al., 2000). Moreover, CgPKC, a cAMP pathway-related protein kinase in C. gloeosporioides, is an imperative gene in virulence and appressorium formation (Priyatno et al., 2012). Together, these findings could facilitate the elucidation of the molecular mechanism underlying appressorium formation. However, details investigations on the signaling cascades in C. gloeosporioides are not well-studied to date.

We therefore aimed to explore the multiple functions of CgMK1. Here, we demonstrated that CgMK1 (an ortholog of Fus3/Kss1) in C. gloeosporioides plays an important role in appressorium formation and pathogenesis. The results also indicated that CgMK1 affects aerial hyphal growth and governs the expression of melanin production-related genes. Furthermore, based on the exogenous application of cAMP, we reveal that CgMK1 may function downstream or independent of a cAMP-dependent signal in appressorium formation.

MATERIALS AND METHODS

Strains and Growth Tests

The wild-type of C. gloeosporioides is CFCC80308, isolated from Populus × beijingensis. The wild-type and its derivatives were cultured in complete medium (CM) or on potato dextrose agar (PDA) plates at 25°C with day/night cycles as previously described (Xu et al., 2016). To assay for defects in stress responses, the size and morphology of the colonies were measured with cultures grown on CM with 1.2 M NaCl, 1 M sorbitol, Congo Red (100 µg/ml), or Calcofluor (60 µg/ml). Statistical analyses of all data were performed by using the Duncan’s range test of SPSS20.0. The P-value < 0.05 was considered as statistically significant. All assays were performed in triplicate, in three independent experiments.

Bioinformatics Analysis of CgMK1

The sequence of CgMK1 was retrieved from the C. gloeosporioides genome database1. A BLASTp search on the National Center for Biotechnology Information (NCBI) website was used to find all annotated Fus3 genes from other fungal genomes. The InterPro2 was used to predict domain of CgMK1. Amino-acid sequences of CgMK1 and Fus3/Kss1 homologs from other fungi were aligned by the ClustalX 2.1. For homology analysis, we constructed a phylogenetic tree in MEGA 7.0 with 1000 bootstrap replicates and the neighbor-joining method as previously described (Sun et al., 2016).

Targeted Gene Knockout and Complementation

The CgMK1 gene replacement constructs were established using the split-marker gene replacement strategy. To construct the upstream (~1.2 kb) and downstream (~1.4 kb) flanking sequences of CgMK1, the primers CgMK1-5For, CgMK1-5Frev, CgMK1-3Ffor, and CgMK1-3Frev were used for the 5’-flanking sequence and the 3’-flanking sequence, respectively. The next step required fusing the fragments and two-thirds of the hygromycin cassette by overlapping PCR using the primers CgMK1-5Ffor/HY-R and YG-F/CgMK1-3Frev, respectively. The CgMK1 gene replacement and transformation methods were performed as described in a previous study (Xu et al., 2016). The PCR assays were used to confirm whether the CgMK1 gene had been replaced using the primers External-CgMK1for/External-CgMK1rev and RT-CgMK1for/RT-CgMK1rev. To analyze

1http://genome.jgi.doe.gov/Gloci1/Gloci1.home.html
2http://www.ebi.ac.uk/interpro/
homologous recombination events in the transformants, southern blotting was performed to confirm the deletion of CgMK1 with the DIG High PrimeDNALabeling and Detection Starter Kit I in conformity to the manufacturers' protocol (Roche, Germany). ScaI was used to digest the genomic DNA of wild-type and transformants. The probe amplified with primers ProbePHfor, ProbePHrev from HPH and the primers ProbeCgMK1for, ProbeCgMK1rev from the C. gloeosporioides CFCC80308 and labeled with DIG primer (Table 1).

The fragment for complementing the CgMK1 deletion strains, which contains the entire CgMK1 coding region, its native promoter, and terminator regions, was PCR-amplified using primers CgMK1-Compfor/CgMK1-Comprev, and inserted into a modified pBC-phleo vector conferring phleomycin resistance (provided by FGSC). After harvesting the transformants, we selected the strain from the TB3 medium (with 100 µg/mL phleomycin) and confirmed the results using RT-PCR.

**Generation of the CgMK1/CgMK1::eGFP Fusion Construct**

To explore the subcellular location of CgMK1, the sequence was amplified from the wild-type strain using the primers CgMK1-5Ffor/promoter + CgMK1rev, CgMK1-5Ffor/promoter + CgMK1 + eGFPrev and a pKD6-GFP plasmid. The resulting DNA product was then transferred into the protoplasts of the ΔCgMK1-13 mutant. Upon harvesting the transformants, we selected the strain from TB3 medium (with 100 µg/mL phleomycin), examined the eGFP signals, and confirmed using qPCR. These tests were conducted as previously described (Xu et al., 2016).

**Quantitative RT-PCR Assays**

To obtain whole RNA of the vegetative mycelia, TRIzol reagent (Invitrogen) was used to isolate RNA from the conidia of the wild-type and transformants. The RNA was purified with the PureLink RNA Mini Kit (Ambion). The RNA products were incubated with DNase I (TaKaRa) to reduce DNA contamination. For the quantitative real-time PCR (qRT-PCR) assays, Oligo-DT and SuperScript III reverse transcriptase (Invitrogen) were used to transcribe the mRNA into cDNA. The qRT-PCR reactions were conducted to analyze the transcript levels. The Cg18S gene served as an internal reference in this study. The 2−ΔΔCT method was used to analyze gene expression levels of the genes (Livak and Schmittgen, 2001). These tests were carried out as previously described (Xu et al., 2016). Quantitative real-time PCR was performed in triplicate with three independent biological experiments. All primers used in this study are listed in Table 1.

**Conidiation and Appressorium Formation Tests**

For the conidiation assays, we selected 5 mm diameter mycelial plugs from the edge of a 3-days-old colony that had been placed onto PDA medium and grown at 25°C for 7 days. Conidial suspensions from each strain were selected from the PDA plates after applying 5 mL of sterile water. Conidial suspensions were placed at 25°C for 12 h on a GelBond membrane to allow germination and growth. The suspensions were monitored, and the colony-forming efficiency was assessed under the microscope.

**Table 1 | PCR primers used in this study.**

| Primer name | Sequence | Use in this study |
|-------------|----------|------------------|
| CgMK1-5Ffor | CTTCACCTGCGCCTGTCGTTT | 5F flanking sequence |
| CgMK1-5Frev | GATACATTCCGCGCCAGAAC | 3F flanking sequence |
| CgMK1-3Ffor | TCCCTACCTGGAGAAGTTCT | External sequence used for validation of mutant |
| CgMK1-3Frev | CGCTTCAAAGCCTTCCCCTA | Complementary sequence |
| External-CgMK1for | CACCGACCAAGGCAATTGTG | Complementary sequence without stop codon |
| External-CgMK1rev | GCATTGGTGTGCCGTTCTTT | Complementary and sublocation sequence cassette |
| CgMK1-Compfor | CGTCCCTTTTGGGCAAACCTTAC | qRT-PCR of CgMK1 |
| CgMK1-Comprev | TAAACCTGAAAGACCCTTGG | qRT-PCR of Cg18S |
| Promoter + CgMK1rev | CGCGATAATCTCTCGCTAAA | qRT-PCR for hybridization |
| Promoter + CgMK1 + eGFPrev | AC GGCTTCTTTTCTCTTAGTGGTA | Probe PH for hybridization |
| RT-CgMK1for | TGCTAGTGGAAAGGCCCTTG | Probe CgMK1 sequence used for hybridization |
| RT-CgMK1rev | AGGAAAGTACTTCTCTGCGC | The 2/3rd portion of the hygromycin cassette |
| Cg18S-up | GTGAGGCCCTCAAAGGTAGTGG | Hygromycin cassette |
| Cg18S-down | GGATCCCAGTGCGAGACGT | Hygromycin cassette |
| ProbePHfor | AAGAACGGCAGACCAATGC | Hygromycin cassette |
| ProbePHrev | ATGAGAGAAGAACAGGCGC | Hygromycin cassette |
| ProbeCgMK1for | GCGAAGATCTCTGCTCCTC | Hygromycin cassette |
| ProbeCgMK1rev | GATGTTGCGGCAAACCTTATT | Hygromycin cassette |
Equal numbers of spores were placed on the hydrophobic surface of the GelBond film (Lonza) at 28°C for 12 h to examine appressoria formation. The appressoria from germinated spores were counted. At least 200 conidia were surveyed in experiments. The rates of conidiation and appressorium formation assays were performed in triplicate, in three independent experiments.

Pathogenicity Assays
To test the virulence, approximately 3-week-old detached leaves of *Populus × beijingensis* were incubated with 20 μL conidium suspension (10^5 conidia/mL) at 25°C. Sufficient moisture was maintained, and the morphology and size of the lesions were observed and measured at 4–5 dpi. For the wound-inoculation assays, the conidia of the strain were placed on the wounded sites of the poplar leaves, and the morphology and size of the lesions were observed and measured at 2–3 dpi. At least three virulence experiments were performed, with seven replicates each.

RESULTS

Deletion of *CgMK1* Impairs Aerial Hyphae Growth

We explored a gene in the *C. gloeosporioides* genome database (JGI) by BLASTp search that displayed high identity with PMK1 of *M. oryzae*, thus designated as *CgMK1*. *CgMK1* contained the protein kinase domain with a threonine and tyrosine phosphorylation site, and displayed high identity with Fus3/Kss1 from *C. orbiculare* (99.4%), *V. dahliae* (98.0%), *Botrytis cinerea* (94.6%), and *Fusarium graminearum* (98.6%) (*Supplementary Figure S1A*). Furthermore, phylogenetic analysis demonstrated that *CgMK1* is homologous to Fus3/Kss1 MAPKs of other fungi and the Fus3/Kss1 is highly conserved among fungi (*Supplementary Figure S1B*).

To investigate the role of *CgMK1* in *C. gloeosporioides*, *CgMK1* deletion mutants (Δ*CgMK1*-8, Δ*CgMK1*-13) were obtained by replacing the wild-type *CgMK1* gene with a hygromycin cassette (*Supplementary Figure S2*). For mutant complementation, we reintroduced the wild-type *CgMK1* into the Δ*CgMK1*-13 mutant to produce the complementation strain (Δ*CgMK1*/CgMK1; *Supplementary Figure S2E*).

While the Δ*CgMK1* mutants showed no obvious defects of growth rate on potato dextrose agar (PDA), aerial hyphal growth was remarkably reduced (*Figures 1A,B*). Likewise, radial growth of the Δ*CgMK1* mutants on solid media showed no difference to the wild-type, but biomass production in the liquid was apparently reduced (*Figures 1C,D*). These results imply that *CgMK1* is essential to aerial hyphal growth.
Deletion of *CgMK1* Causes the Repression of Melanin Biosynthesis Genes

Because the deletion of *CgMK1* clearly affected melanin deposition in the vegetative hyphae, we then investigated whether the melanin defect of the Δ*CgMK1* mutants could result from the downregulation of melanin biosynthesis genes. To verify the hypothesis, we identified key genes participated in melanin biosynthesis in *C. gloeosporioides* using comparative genomics. Here, three genes associated with melanin production via the dihydroxy naphthalene (DHN) pathway were identified as follows, β-ketoacyl synthase (PKS1), scytalone dehydratase (SCD1), and trihydroxy naphthalene reductase (THR1). The transcript levels of these three genes were measured through quantitative real-time PCR method to compare their expression between wild-type and Δ*CgMK1* mutant. The $2^{-\Delta\Delta CT}$ method of data statistics showed that the transcription levels of the melanin biosynthesis genes were strongly repressed in the Δ*CgMK1* mutants. The relative expression levels of these genes in the Δ*CgMK1* mutants were at least 50 fold lower gene transcript abundance compared to that in the wild-type strain (Figure 2). These results thus indicate that *CgMK1* plays a key role in controlling melanin production.
**CgMK1 Is Necessary for Appressorium Formation**

We also examined the role of CgMK1 in appressorium formation and conidia germination using a GelBond membrane. Firstly, we determined that the ΔCgMK1 mutant and the wild-type exhibit a similar conidia germination phenotypes (data not shown), though the germ tube of the ΔCgMK1 mutants continued to lengthen instead of forming appressoria at 12 hpi (Figure 3A). Based on the data statistics, about 70% of the appressoria were produced in the wild-type strain and ΔCgMK1/ΔCgMK1; however, none were produced by the ΔCgMK1 mutants (Figure 3B). These findings demonstrate that CgMK1 is required for appressorium development.

**Exogenous cAMP Does Not Restore the Appressorium Defective of ΔCgMK1 Mutants**

Previous studies have shown that the extracellular application of cAMP induces appressorium formation on hydrophilic surfaces, indicating that the cAMP-dependent signal is important for appressorium development. Here, the application of extracellular cAMP to the ΔCgMK1 mutants induced conidia germination in a similar manner as in the wild-type, but exogenous cAMP was unable to recover the appressorium formation defect in the ΔCgMK1 mutants (Figure 4). This suggests that CgMK1 controls the formation of appressoria in a CAMP-independent manner or functions downstream of cAMP signaling.

**Disruption of CgMK1 Results in a Loss of Virulence**

The ΔCgMK1 mutants lost the capability to form appressoria on artificial hydrophobic surfaces. To determine whether this results in defective pathogenicity, the strains were inoculated on detached but intact poplar (Populus x beijingensis) leaves. At 7 dpi, the ΔCgMK1 mutants appeared to completely lose virulence compared to the distinct lesion symptoms of the leaves placed with the ΔCgMK1/ΔCgMK1 and wild-type (Figures 5A,C). Since the ΔCgMK1 mutants could not form appressoria, the wounded poplar leaves were inoculated with the above strain as well. Similar to the symptoms of the intact leaves, the wild-type strain and ΔCgMK1/ΔCgMK1 caused typical disease...
The CgMK1 deletion mutants lose pathogenicity on poplar leaves, indicating that CgMK1 governs virulence. (A) Spores (10⁵/mL) from WT, ΔCgMK1 mutants and the ΔCgMK1/CgMK1 were inoculated on the detached leaves of 1-year-old poplar seedlings. Ten healthy intact poplar leaves were used in each independent experiment with three replicates. The images were captured at 7 dpi. (B) Spores (10⁵/mL) of WT, ΔCgMK1 mutants and the ΔCgMK1/CgMK1 were inoculated on detached leaves of 1-year-old poplar seedlings. Ten poplar leaves were wounded and were used in each independent experiment with three replicates. The images were captured at 7 dpi. (C) Quantification of lesion sizes caused by the strains in each independent experiment. Values represent the mean of three repetitions. The data were performed by using the Duncan’s range test. Asterisks mean statistically significant differences at P = 0.05. (D) Infection hyphae development of the strains on the detached poplar leaves at 48 hpi. Invasive hyphae were treated with lactophenol aniline blue. A: appressorium; C: conidium; IH: invasive hyphae. Scale bar: 10 μm.

symptoms; in contrast, the ΔCgMK1 mutants failed to infect the wounded leaves (Figures 5B,C). Although the wild-type and ΔCgMK1/CgMK1 inoculated on the leaf surfaces produced masses of invasive hyphae by 2 dpi, the ΔCgMK1 mutants did not (Figure 5D). Microscopic observations showed that the ΔCgMK1 mutants extended the germ tube but hardly penetrated into the plant cell. Conversely, in the wild-type, the majority of the conidia developed melanized appressoria, which invaded the epidermal cells and then formed swollen invading hyphae (Figure 5D).
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FIGURE 6 | Assays for penetration and colonization defects in the CgMK1 mutants. (A) Onion epidermal cell penetration assay of the ΔCgMK1 mutants. The assay was performed by inoculating conidial suspensions from the indicated strains. Examination under a light microscope was performed after aniline-blue staining. A: appressorium; C: conidium; IH: invasive hyphae. Scale bar: 10 µm. (B) Cellophane membranes penetration assay was performed by the conidial suspensions from strains. Colonies of the indicated strains grown on PDA plates covered with a cellophane membrane (before). Then the medium was removed the cellophane membrane and placed for additional days (after). Photos in the first row were taken at 3 dpi, the second row were taken at 5 dpi that showed growth of strains after penetration from cellophane membrane.

We also performed a penetration assay on onion epidermal cells and cellophane membranes. The results showed that the ΔCgMK1 mutants produced massive amounts of hyphae on the onion epidermis, but very few hyphae could penetrate into the onion epidermal cells. Nevertheless, the wild-type and ΔCgMK1/CgMK1 strains infected the epidermal cells and expanded into the epidermal tissues at 24 hpi (Figure 6A). We also found that, in contrast to the ΔCgMK1/CgMK1 and wild-type, the ΔCgMK1 mutants failed to penetrate the cellophane membranes and were then unable to grow (Figure 6B). Collectively, the ΔCgMK1 mutants lost virulence due to their failure to penetrate into the plant tissues.

Subcellular Localization of the CgMK1
To investigate the cellular localization of CgMK1 in C. gloeosporioides, we generated a fragment containing the CgMK1 and enhanced green fluorescent protein (eGFP), hereby designated as CgMK1_C:eGFP. The CgMK1_C:eGFP and phleomycin resistance cassette were introduced into ΔCgMK1-13 protoplasts by co-transformation. GFP signals were detected under the active expression of the native promoter and whole CgMK1 sequence. In our study, the fluorescence was detected at different stages of conidium, germinating conidia, germ tube, and appressorium formation (Figure 7) Moreover, the expression was relatively higher in the mature appressorium than the other stages. These results indicate that CgMK1 is induced during appressorium formation.

CgMK1 Is Involved in the Response to High Osmotic Stress
To determine the function of CgMK1 in stress responses, we observed the wild-type, ΔCgMK1 and ΔCgMK1/CgMK1 strains on complete medium (CM) with NaCl, sorbitol, calcofluor white (CFW) and Congo red (CR). The ΔCgMK1 mutants showed sensitivity to 1.2 M NaCl, 1 M Sorbitol (Figure 8A). We also tested the growth rate of the strains on plates containing NaCl and sorbitol. The rate of growth inhibition of ΔCgMK1 mutants when grown on CM containing NaCl and sorbitol was higher than that of wild-type and ΔCgMK1/CgMK1 strain, respectively (Figure 8B). On the other hand, the ΔCgMK1 mutants showed no obvious difference from wild-type and ΔCgMK1/CgMK1 strain in growth in response to CFW and CR (Supplementary Figure S3). These results suggest that CgMK1 is involved in the response to high osmotic stress, but not important in the maintenance of cell wall integrity.

DISCUSSION
Here, we examined the functions of CgMK1 in appressorium formation and virulence of poplar anthracnose fungus, C. gloeosporioides. The results demonstrated that CgMK1 plays key roles in appressorium formation, melanin production, and plant infection.
Fungal MAPK cascades play key roles in governing fungal development and signaling fungal responses to a variety of stress stimuli. In search of the component of MAPK, Fus3/Kss1-type MAPK was found to be required for fungi to attack plant tissues. Studies on the Fus3/Kss1-type MAPK in pathogenic fungi has revealed the importance in the production of infection structures (Zhao and Xu, 2007; Zhao et al., 2007). In pathogenic fungi, appressoria are special infection structures that impart high turgor pressure and excrete plant cell degradation enzymes that aid the penetration of the infection peg into plant cells (Mendgen et al., 1996; van Kan, 2006). Previous studies have demonstrated that many conserved signal modules are involved in appressorium formation and pathogenicity, at least including small GTPases (Fukada and Kubo, 2015; Xu et al., 2016) and the MAPK pathway (Hamel et al., 2012). In C. gloeosporioides, loss of CgRhoB resulted in shorter germ tubes and enhanced appressorium formation after germination on the hydrophobic surface. However, the present study demonstrated that CgMK1 is crucial for appressorium formation rather than conidia germination in C. gloeosporioides. The ΔCgMK1 mutants exhibited no obviously defective in conidia germination, indicating that CgMK1 does not contribute to conidia germination, which differs from C. orbiculare (Takano et al., 2000; Xiong et al., 2015). Subcellular localization of the CgMK1 showed that CgMK1 is induced during appressorium formation.

The C. gloeosporioides strain lacking the CgPKAC gene displayed delayed appressorium formation (Priyatno et al., 2012). The CMK1 cooperated with cAMP to control appressorium formation and germination in C. orbiculare (Takano et al., 2000). The PMK1 deletion mutants failed to develop appressoria on Teflon membranes in M. oryzae, and formed swollen germ tubes when exogenous cAMP was applied to the hydrophilic surfaces (Xu and Hamer, 1996). The results indicated that cAMP is necessary for appressorium formation and functions upstream of MAPK in appressoria-forming fungi. To explore the reason why the ΔCgMK1 mutants failed to form the appressorium, we also revealed the function of exogenous cAMP in ΔCgMK1 mutants. The results showed that exogenous cAMP application does not restore the defect of appressorium formation in the ΔCgMK1 mutants. This suggested that CgMK1 controls the formation of appressoria in a cAMP-independent manner or functions downstream of CAMP signaling.

Appressorium melanization is vital for the high turgor pressure necessary for the invasion and growth of appressoria-forming pathogens (Howard and Ferrari, 1989). In C. orbiculare (Takano et al., 2000), several genes involved in melanin biosynthesis have been isolated, such as PKS1, THR1, and SCD1. CgPKS1, a melanin biosynthesis gene in C. graminicola, is not required for appressoria turgor generation, but is important for penetration and melanin synthesis (Ludwig et al., 2014). In

![FIGURE 7](image_url) Subcellular localization of CgMK1 in appressorium formation. Fluorescence of CgMK1 was detected by C-terminal eGFP fusion under its promoter. Green fluorescence was detected during the stages of germination and appressorium formation on the hydrophobic surface of the GelBond membrane. Scale bar: 10 µm.
this study, the ΔCgMK1 mutants exhibited slightly darkened colonies. The expression of three melanin biosynthesis genes including PKS1, THR1, and SCD1 were significantly down-regulated in the ΔCgMK1 mutants.

Previous studies have demonstrated that deletion of the Fus3/Kss1-related pathway results in a loss of pathogenesis owing to an inability of several pathogenic fungi to colonize host tissues or form infection structures, for example VMK1 in V. dahliae, and PMK1 in M. oryzae (Xu and Hamer, 1996). Knockout of CgMK1 resulted in mutants that were non-pathogenic and unable to form lesions on poplar leaves. On the poplar epidermis, ΔCgMK1 mutants formed elongated germ tubes, but were unable to infect the epidermal cells. In addition, penetration assays with cellophane membranes and onion epidermal cells suggested that the ΔCgMK1 mutants were unable to penetrate the cellophane membranes. Furthermore, we discovered that the hyphae of the ΔCgMK1 mutants could only marginally infect onion epidermal cells. This result differs from that displayed in other fungi such as C. orbiculare (Takano et al., 2000), where the Fus3/Kss1 homologs in other fungi were aligned by the ClustalX 2.1. The conserved TEY sequence needed for kinase activation is marked by asterisks. aa, amino acid.

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2017.02216/full#supplementary-material

**AUTHOR CONTRIBUTIONS**

YW, CT, and PH designed the experiments. PH, XW, and XZ performed the experiments and the data analyses. YW and PH prepared the figures and wrote the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2017.02216/full#supplementary-material
with full-length protein sequences and neighbor-joining with 1000 bootstrap replicates.

**FIGURE S2 |** Disruption and complementation of CgMK1 in C. gloeosporioides. *(A,B)* Replacement method for the deletion of CgMK1. The 1.0 kb sequence of CgMK1 was substituted for a sequence including the hph cassette (Probe 3). Filtration of the ΔCgMK1 mutants with primers External-CgMK1for and External-CgMK1rev *(A)* and primers RT-CgMK1for and RT-CgMK1rev *(B)*. *(C)* Confirmation of the ΔCgMK1 mutants and complementation of CgMK1 by semiquantitative reverse transcription PCR. *(D,E)* Southern blot of wild-type strain and two independent ΔCgMK1 mutants (ΔCgMK1-8, ΔCgMK1-13), DNA specimens of transformants were digested by Scal. The enzyme-digested product was probed with a DNA sequence from a hph gene (Probe 3) of the HPH *(D)* and CgMK1 (1.493 bp fragment of the 3′-flanking sequence of CgMK1) *(E).*

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