Article

CgSCD1 Is Essential for Melanin Biosynthesis and Pathogenicity of Colletotrichum gloeosporioides

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Received: 17 January 2020; Accepted: 19 February 2020; Published: 20 February 2020

Abstract: Colletotrichum gloeosporioides, an important phytopathogenic fungus, mainly infects tropical fruits and results in serious anthracnose. Previous studies have shown that melanin biosynthesis inhibitor can inhibit the melanization of the appressoria of Magnaporthe grisea and Colletotrichum orbiculare, resulting in limited infection of the hosts. In this study, we identified and characterized a scytalone dehydratase gene (CgSCD1) from C. gloeosporioides which is involved in melanin synthesis. The CgSCD1 gene deletion mutant ΔCgscd1 was obtained using homologous recombination. The ΔCgscd1 mutant showed no melanin accumulation on appressoria formation and vegetative hyphae. Furthermore, the virulence of ΔCgscd1 was significantly reduced in comparison with the wild-type (WT) strain. Further investigations showed that the growth rate as well as germination and appressorium formation of ΔCgscd1 displayed no difference compared to the wild-type and complemented transformant Cgscd1com strains. Furthermore, we found that the appressorial turgor pressure in the ΔCgscd1 mutant showed no difference compared to that in the WT and Cgscd1com strains in the incipient cytorrhysis experiment. However, fewer infectious hyphae of ΔCgscd1 were observed in the penetration experiments, suggesting that the penetration ability of nonpigmented appressoria was partially impaired. In conclusion, we identified the CgSCD1 gene, which is involved in melanin synthesis and pathogenicity, and found that the melanization defect did not affect appressorial turgor pressure in C. gloeosporioides.

Keywords: Colletotrichum gloeosporioides; scytalone dehydratase; appressoria; virulence

1. Introduction

Colletotrichum gloeosporioides is a fungal phytopathogen that causes devastation of crop and ornamental plants worldwide, resulting in substantial economic losses. During the infection process, C. gloeosporioides produces a series of specialized infection structures, including germ tubes, appressoria, infectious hyphae, and necrotrophic hyphae [1]. The host infection process can be roughly divided into two stages, biotrophy and necrotrophy, based on differences in the infection strategy [2]. Melanized appressoria are firstly produced at the top of the germ tubes, generated by conidia-penetrated host surfaces through a combination of mechanical forces and enzymatic degradation; the spherical biotrophic hyphae in living host cells (biotrophy stage) are then transformed and differentiated into thin, fast-growing secondary hyphae, thus causing the host tissue to be destroyed rapidly (necrotrophy stage) [1].

Most filamentous fungi have the ability to produce dark-pigmented melanin, which is used to protect organisms against destructive environmental stresses such as UV radiation, extreme temperatures, and strong oxidants [3–5]. As an amorphous polymer, the production of melanin can be divided into two common synthesis strategies, which are L-3,4-dihydroxyphenylalanine (L-dopa)
melanin formed by the polymerization of phenolic compounds and 1,8-dihydroxynaphthalene (DHN) melanin formed by a polyketide synthase and subsequent polymerization [6].

There are plenty of related studies on the DHN melanin synthesis pathway in Colletotrichum orbiculare, which is a pathogen that causes anthracnose disease of cucurbit plants, particularly cantaloupe and cucumber [7]. The process of melanogenesis starts with pentaketide 1,3,6,8-tetrahydroxynaphthalene (T4HN), directly formed by polyketide synthase (transcribed by PKS1), and proceeds to the synthesis of scytalone by T4HN reductase (transcribed by T4HR1) and 1,3,8-trihydroxynaphthalene (T3HN) reductase (transcribed by THR1). From scytalone, the synthesis process needs to go through two dehydration steps by SCD1 and one reduction step by THR1, including dehydration of scytalone to T3HN, reduction of T3HN to vermelone, and dehydration of vermelone to 1,8-DHN, which are polymerized and oxidized to obtain melanin [8]. The crucial enzymes of melanin synthesis, transcribed by PKS1, T4HR1, THR1, SCD1, and a transcription factor of melanin synthesis (CMR1), have been systematically studied as important biosynthetic enzymes as well as pathogenic factors of C. orbiculare [9–12].

The biological functions of DHN melanin in various fungi species have been explored through albino mutants which include random mutations and targeted knockout mutants, suggesting that DHN melanin plays an important role in fungal life cycles. Melanin promotes virulence by increasing resistance to hydrogen peroxide in human pathogen Talaromyces marneffei [13]. In the filamentous ascomycete Sordaria macrospora, melanin biosynthesis is related to fruiting body development and is controlled by specific regulatory genes involved in sexual differentiation [14].

Previous studies of rice blast pathogen Magnaporthe grisea have shown that melanized appressoria play an important role in mechanical penetration. Melanin-deficient mutants are unable to infect intact plants. The mechanical pressures caused by the increased osmotic pressure inside melanized appressoria and strong adhesives may be essential to the penetration process [15–18]. In Colletotrichum kahawae, the pathogen of coffee berry disease, unmelanized appressoria induced by tricyclazole show turgor pressure as low as one-quarter that of melanized appressoria, accompanied by a decrease in infection [19]. The albino mutant of C. graminicola is unable to penetrate intact leaves, resulting in decreased virulence. However, the accumulation of osmolytes and the generation of turgor in appressoria are independent of melanin [20]. Related studies in Phakopsora pachyrhizi have shown similar results, with turgor pressure accumulation in appressoria also found to be independent of melanin [21]. The causes of this phenomenon, probably due to the diversity of species and the functions of appressoria, are not completely consistent during the infection processes.

To investigate the synthesis pathway and function of melanin in the appressorial formation and penetration process in C. gloeosporioides, we identified and characterized the CgSCD1 gene, which encodes for scytalone dehydratase in the melanin biosynthesis pathway. We then established a CgSCD1 deletion mutant, ΔCgscd1, which showed no melanization and decreased virulence. In a further study, we found that the decreased virulence of the ΔCgscd1 mutant was due to the damaged penetration ability of appressoria but that this was not related to turgor pressure, which was unchanged. We also found that CgSCD1 on the cytoplasm of melanized appressoria, on the germ tubes, and on the germinated conidia.

2. Results

2.1. Cloning of the Scytalone Dehydratase Gene from C. gloeosporioides

The EX2016-02 strain of wild-type C. gloeosporioides was isolated from the postharvest fruit of key lime (Citrus aurantifolia) in China [22]. Sequencing results of gDNA and cDNA PCR amplification products showed that the CgSCD1 gene was 784 bp in length (GenBank accession number: MN539625), containing two introns and encoding a homolog of scytalone dehydratases. The results of multiple sequence alignment (Figure 1A) indicated that CgSCD1 had many conserved sites identical to the sequences of other fungal species [23,24]. A phylogenetic tree was constructed to update the new
functional clades using MEGA-X software. CgSCD1 displayed high identity with proteins of *C. orbiculare* which contain a conserved nuclear transport factor 2 (NTF2-like) superfamily domain. All of the above bioinformatics analyses suggest that the predicted product of CgSCD1 may have the same function as the fungal scytalone dehydratase reported in other species. Another homologous protein EQB45228, annotated scytalone dehydratase, was also found from the reference genome data of *C. gloeosporioides* strain Cg14. However, phylogenetic tree analysis showed that it did not aggregate with the CgSCD1 sequences of other fungal species (Figure 1B).

**Figure 1.** Bioinformatics analysis of CgSCD1. (A) Comparison of the amino acid sequence of the *Colletotrichum gloeosporioides* CgSCD1 with those of *Alternaria alternata*, *Aspergillus nidulans*, *Botrytis cinerea*, *Magnaporthe grisea*, *Neurospora crassa*, and *Verticillium dahliae*. Identical amino acids are highlighted with a black background. (B) Phylogenetic analysis of CgSCD1 and homologous proteins from other fungi. The protein sequence (MN566451) shown in red letters was from this experiment.

### 2.2. Generation of the ΔCgscd1 Mutant and Complementation

To investigate the function of CgSCD1, a ΔCgscd1 mutant and the complementation Cgscd1com strain were generated by double-crossover homologous recombination and T-DNA insertion, respectively. In the ΔCgscd1 mutant, the original CgSCD1 sequence was replaced by NrsR cassettes, and no CgSCD1 gene expression was detected by RT-PCR (Figure 2C,D). Fungal transformants were obtained after selection by resistance to nourseothricin. Finally, the ΔCgscd1 strain was confirmed to be an invalid mutant by sequencing and RT-PCR. The existence of a NrsR cassette was confirmed by PCR using primers N-F and N-R, with an expected product size of about 2.1 kb (Figure 2C). In addition, successful deletion in ΔCgscd1 was verified by RT-PCR, and no expression of CgSCD1 was detected at the transcription level (Figure 2D). In the Cgscd1com strain, the hygromycin resistance cassette and the CgSCD1 gene were amplified (Figure 2C,D).
2.3. Characterization of ΔCgscd1 and Cgscd1com

The phenotypes of the wild-type (WT), ΔCgscd1 mutant, and Cgscd1com strains were characterized by the color and growth rate of the colony, conidium germination, appressorium formation, and melanization. The ΔCgscd1 mutant only formed colorless appressoria (Figure 3A). The Cgscd1com strain showed a melanized colony and appressoria, as did the WT strain. In the ΔCgscd1 mutant, the colonies were white, and no melanin accumulated during the initial culture. After about 7 days of incubation, the colonies and the potato dextrose agar (PDA) medium turned brown-red. This change in color may be caused by the accumulation of the scytalone, an intermediate metabolite of melanin synthesis, which was secreted into the culture medium (Figure 4) [25]. This indicates that the deletion of CgSCD1 resulted in the albino mutant.
The growth rate of the ΔCgscd1 strain on PDA, as determined through the calculation of colony diameter, displayed no difference compared to the WT and Cgscd1com strains (Figure 3B). However, the dry weight of mycelium in the ΔCgscd1 mutant was considerably reduced after 7 days at 25 °C compared to that in WT, and was restored in the Cgscd1com strain (Figure 3C). Furthermore, in the WT, ΔCgscd1, and Cgscd1com strains, the germination capabilities as well as appressorium formation rate were almost 90%, with no differences in germination and appressorium formation (Figure 3D). However, almost 50% of the appressoria of the ΔCgscd1 mutant developed a secondary or tertiary appressorium. Less than 20% of the appressoria of the WT and Cgscd1com strains differentiated into a secondary appressorium (Figure 3A). This suggests that the deletion of CgSCD1 in the ΔCgscd1 mutant affected the biomass and appressorium lateral germination.

Figure 3. Phenotypic characteristics of the wild-type (WT), ΔCgscd1, and Cgscd1com strains of C. gloeosporioides. (A) Colonies grown in potato dextrose agar (PDA) for 7 days, at 25 °C (images on the two columns on the left). Germinated conidia, from cultures grown for 7 days, at 25 °C (images on the two columns on the right). Arrows point to mature appressoria. Ap, appressoria; C, conidia. Scale bar (for (A)) = 10 µm. (B) Periodic (24 h) measurement of colony growth diameter. (C) Dry weight of mycelia grown on cellophane-covered PDA for 7 days. (D) Germination and appressorium formation of each strain. (E) Quantitative differences in the WT, ΔCgscd1, and Cgscd1com strains in lateral germination events of appressoria (1, 2, 3 represent the number of appressoria produced by a single conidium; n = 100). For (C,E): p < 0.05 (*).
The growth rate of the ΔCgscd1 strain on PDA, as determined through the calculation of colony diameter, displayed no difference compared to the WT and Cgscd1com strains (Figure 3B). However, the dry weight of mycelium in the ΔCgscd1 mutant was considerably reduced after 7 days at 25 °C compared to that in WT, and was restored in the Cgscd1com strain (Figure 3C). Furthermore, in the WT, ΔCgscd1, and Cgscd1com strains, the germination capabilities as well as appressorium formation rate were almost 90%, with no differences in germination and appressorium formation (Figure 3D). However, almost 50% of the appressoria of the ΔCgscd1 mutant developed a secondary or tertiary appressorium. Less than 20% of the appressoria of the WT and Cgscd1com strains differentiated into a secondary appressorium (Figure 3A). This suggests that the deletion of CgSCD1 in the ΔCgscd1 mutant affected the biomass and appressorium lateral germination.

2.4. In Vitro Scytalone Release from ΔCgscd1 is Able to Restore Sclerotia Melanization in ΔBcpks12

The melanin biosynthesis pathway in Botrytis cinerea started with the synthesis of pentaketide by Bcpks12, proceeded to the synthesis of scytalone, and then the scytalone went through two dehydration steps by Bscsd1 [26]. The sclerotia of ΔBcpks12 were light yellow from the inside core to the surface [27]. Here, we predicted that scytalone would be accumulated in the ΔCgscd1 mutant and it may be used in the ΔBcpks12 mutant to restore sclerotia melanization in the ΔBcpks12 mutant. To confirm our hypothesis, a cooperation experiment was performed using ΔCgscd1 and ΔBcpks12. In the test of the cooperation culture, we observed that the sclerotia formed by the ΔBcpks12 mutant were initially white, and then turned light yellow. However, the sclerotia of the ΔBcpks12 mutant on the side close to the ΔCgscd1 colonies turned black after 14 days (Figure 4), indicating that scytalone from the ΔCgscd1 mutant could be secreted into the culture medium and absorbed by ΔBcpks12, and could then continue to synthesize melanin in vivo to restore sclerotia melanization.

2.5. Analysis of ΔCgscd1 Mutant Virulence

To investigate whether the CgSCD1 gene is involved in pathogenicity, virulence analysis was performed on tomato and lime fruits, as well as tomato leaf. We found that the ΔCgscd1 mutant failed to cause obvious sunken lesions on the fruit surface compared with the WT and Cgscd1com strains (Figure 5A). Additionally, the size of the lesion area for the ΔCgscd1 mutant was significantly reduced compared with the WT and Cgscd1com strains (Figure 5B). This indicates that the deletion of CgSCD1 affected the virulence of the ΔCgscd1 mutant. To further study the infection ability of the ΔCgscd1 mutant, 3,3’-diaminobenzidine (DAB) staining was used to test hydrogen peroxide accumulation on the surface [27]. Here, we predicted that scytalone would be accumulated in the dehydration steps by ΔBcpks12, proceeded to the synthesis of scytalone, and then the scytalone went through two steps by ΔCgscd1, and then turned light yellow. However, the sclerotia of the ΔCgscd1 mutant showed almost no accumulation of hydrogen peroxide at 36 hours post-inoculation (hpi), indicating that the early infection process was also limited (Figure 5C). The reason for this could then continue to synthesize melanin in vivo to restore sclerotia melanization.
mutant, 3,3′-diaminobenzidine (DAB) staining was used to test hydrogen peroxide accumulation on the leaf surface during infection.

![Image](https://example.com/image.png)

**Figure 5.** Plant infection and the virulence analysis of ΔCgscd1 mutant. (A) Inoculation of tomato and lime fruits with wounds. (B) Analysis of the lesion area (mm²). (C) The conidial suspension was inoculated on the surface of tomato leaves without injury. The ΔCgscd1 mutant showed almost no accumulation of hydrogen peroxide at 36 hours post-inoculation (hpi). Scale bars (for (A,C)) = 5 mm. For (B): p < 0.05 (*).

The ΔCgscd1 mutant showed almost no accumulation of hydrogen peroxide at 36 h post-inoculation (hpi), indicating that the early infection process was also limited (Figure 5C). The reason for this could be that the albino appressoria of the ΔCgscd1 mutant may not have penetrated the host surface to form infectious hyphae, since the inoculation was performed on leaves without injury.

### 2.6. Loss of Partial Appressorium Function due to Deletion of the CgSCD1 Gene

It has been reported that the appressoria of rice blast fungus (*M. grisea*) use glycerol to generate pressure, which ruptures plant cuticles. Melanin functions as an impermeable barrier to osmolytes, which allows appressoria to accumulate high turgor pressure [18]. Since the ΔCgscd1 mutant had albino appressoria and showed reduced virulence, we predicted that the reduced virulence in the ΔCgscd1 mutant was caused by a loss of appressoria function.

Firstly, the turgor pressure of appressoria was tested in the WT, ΔCgscd1, and Cgscd1com strains according to the method described by Howard [28]. The appressorial turgor pressure was determined by using a serial concentration of PEG6000 solution, which can lead to the collapse of appressoria when the mimic pressure is higher than the appressorial pressure. The conidia and germ tube in the WT, ΔCgscd1, and Cgscd1com strains were shrunken, distorted, and deformed under an osmotic pressure of 5 MPa. Meanwhile, a longitudinal cavity was observed at the appressoria, indicating that the appressoria had collapsed under external osmotic pressure (Figure 6A). With the increase in osmotic pressure from 1 to 5 MPa, the trends in the appressoria collapse rates of the three tested strains were consistent, indicating that the appressorial turgor pressure was almost the same (Figure 6B). This demonstrates that deletion of the CgSCD1 gene did not affect the appressorial internal turgor pressure of the ΔCgscd1 mutant, although the ΔCgscd1 mutant formed albino appressoria in which no melanin accumulated on the cell wall.
Figure 6. Effect of CgSCD1 on appressorium function. (A) Appressoria treated with 5 MPa (osmotic pressure) PEG6000 solution and formed a longitudinal cavity. The arrows point to the longitudinal cavity. Treatment with (a) water (control); (b) PEG6000 solution. (B) The appressorium collapse rates of the three test strains at various osmotic pressures. (C) Onion epidermal infection test. (D) The germination tube and infection hyphae of the WT, ΔCgscd1, and Cgscd1com strains were observed. (E) The effect of lysing enzymes on the integrity of appressorial cell walls. Samples were stained with Nile red. (a) Before enzymolysis. (b) After enzymolysis. Ap, appressorium; C, conidium; IH, infectious hypha; GT, germ tube. Scale bars (for (A,C,E)) = 10 µm. For (D): p < 0.05 (*).

Secondly, the penetration ability in ΔCgscd1 was tested using onion epidermal infection. In the onion epidermal infection experiment, the germ tubes and appressoria are stained by lactophenol cotton blue as they are present on the surface, while the infectious hyphae are not. The infectious hyphae can be easily identified due to them being colorless in the cotton blue staining and on the basis of their distinct morphology as vesicular structures greater in width than the germ tube (Figure 6C). The ΔCgscd1 mutant showed no significant difference in the germination tube formation rate compared with the WT and Cgscd1com strains. It was observed that the appressoria of the ΔCgscd1 mutant
successfully penetrated the onion epidermal cells. However, the formation rate of infectious hyphae in the ΔCgscd1 mutant was extremely low compared with that in the WT strain, and could be recovered in the Cgscd1com strain (Figure 6D). This indicates that deletion of the CgSCD1 gene affected infectious hyphae formation in the ΔCgscd1 mutant.

To test whether a defect in the production of melanin affects cell wall integration in the appressoria of ΔCgscd1, a Nile red staining assay was performed. Nile red is generally used to localize and quantitate lipids, particularly lipid droplets within cells. As shown in Figure 6E, the appressoria in the ΔCgscd1 mutant were stained, but the appressoria of the WT and Cgscd1com strains were not. This suggests that in comparison to WT appressoria, the ΔCgscd1 appressoria were changed in such a way that it was easier for Nile red to penetrate the cell walls and stain the lipids inside. Treatment with lysing enzyme solution was subsequently used to test cell wall sensitivity. Following 2 h of treatment, the conidia, germ tube, and appressoria of the ΔCgscd1 mutant could hardly be observed by light microscopy. The outlines of these structures were considerably fuzzy, and the lipid droplets were stained red and were brighter than those of the WT and Cgscd1com strains (Figure 6E). After 8 h of continuous culture, the cell walls of these structures were completely digested and destroyed by lysing enzymes, leaving only spherical protoplasts. However, most of the melanized appressoria of the WT and Cgscd1com strains retained their shape (Figure 6E). Taken together, the results show that albino appressoria had disintegrated and were more sensitive to lysing enzymes, suggesting that melanin has a protective effect in appressoria.

2.7. Localization of the CgSCD1 Protein

To further investigate the function of the CgSCD1 gene, the subcellular localization of the CgSCD1 protein and the transcription pattern of the CgSCD1 gene in C. gloeosporioides were studied. The levels of cyan fluorescent protein (CFP) fluorescence were assayed at different developmental stages (0, 4, 12, and 72 h) for different structures, including conidia, conidia with initial appressorium formation, conidia with melanized appressorium formation, and vegetative hyphae. The signal was hardly observed in ungerminated conidia, although it increased along with the germination of conidia and eventually reached a maximum level at the melanized appressorium formation stage. The CFP fluorescence levels were observed to correspond with the levels of CgSCD1 gene transcription as determined by qPCR. The signals of CFP fluorescence coincide with the melanization in appressoria at 12 h. Since the signals appeared to be evenly distributed throughout the appressoria, germ tube, and conidia, the CgSCD1 protein likely localized to the cytoplasm in the early germination stage. In vegetative hyphae, we found that the signal could be observed on hyphae cell walls (Figure 7A).

![Figure 7](image_url)

**Figure 7.** Subcellular localization of CgSCD1 at different stages. (A) Cyan fluorescent protein (CFP) fluorescence was detected during the stages of germination, appressorium formation, and vegetative hyphae. Scale bar = 10 µm. (B) qPCR analysis of the CgSCD1 transcript levels. Bars indicate standard deviations of three biological repeats.
3. Discussion

The correlation between melanin and pathogenicity in different fungal species differs due to pathogen diversity. Melanogenic genes are dispensable in virulence in necrotrophic pathogen B. cinerea [26]. In hemibiotrophic pathogens such as M. grisea and C. orbiculare, melanized appressoria play an important role in mechanical penetration, and melanin-deficient mutants cannot infect intact plants. However, little is known about the specific mechanism and role of SCD1 information in virulence. In this study, we firstly cloned and identified the CgSCD1 gene in C. gloeosporioides.

The reference genome database of the Cg14 strain possesses two scytalone dehydratases genes, including the MN566451 (designed in this experiment) and the EQB45228. Phylogenetic tree of fungal scytalone dehydratase orthologs showed that the MN566451 clustered with SCD from Colletotrichum spp. However, the EQB45228 showed lower similarity in the phylogentic tree. The EQB45228 may not have functional redundancy with the MN566451 in the melanin synthesis pathway since the MN566451 deletion mutant △Cgscd1 showed melanin synthesis deficiency in this study. It has been reported that there are two predicted scytalone dehydratases genes in B. cinerea and one is presumed to be a non-functional gene [26]. The EQB45228 may work somewhere else or just be a non-functional gene, which should be investigated by the deletion mutant of the EQB45228 next.

As an intermediate metabolite of the melanin biosynthesis, scytalone was accumulated and secreted into the medium due to the deletion of CgSCD1 in the △Cgscd1 mutant. It is not yet possible to confirm the proportion of scytalone secreted into the medium, but the scytalone remaining inside the fungi does not seem to affect most growth and development of themselves, except for a slight decrease in biomass. Similar phenotypes have been described in previous studies of Bipolaris oryzae [29] and B. cinerea [26]. Interestingly, although there was no difference in the number of appressoria formation among the WT, △Cgscd1, and Cgscd1com strains, the proportion of secondary and tertiary appressorium of △Cgscd1 mutant was increased, presumably due to a feedback effect caused by the unsuccessful penetration behavior of albino appressoria.

Melanin facilitates glycerol accumulation, increasing osmotic pressure in appressoria, which is essential to the penetration process in rice blast pathogen M. grisea. Melanin-deficient mutants of M. grisea are unable to infect intact plants. However, studies in Phakopsora pachyrhizi and C. graminicola showed that there was no change in turgor pressure in albino appressoria [20,21]. In this study, we tested the appressoria turgor pressure of the △Cgscd1 mutant and found that the defect in the production of melanin has no effect on appressoria turgor pressure in C. gloeosporioides. It is still not clear how appressoria maintain osmotic pressure and generate high turgor pressure without melanin.

In the assays of penetrating ability, the △Cgscd1 mutant appressoria successfully penetrated onion epidermal cells, but the number of infectious hyphae were extremely low compared to that in the WT strain. This indicates that appressoria penetration is partially impaired in the △Cgscd1 mutant. The proportion of secondary and tertiary appressoria of the △Cgscd1 mutant was increased, presumably due to a feedback effect caused by the unsuccessful penetration behavior of albino appressoria. We detected that turgor pressure was unchanged. In the Nile red staining assay, the △Cgscd1 mutant was more easily stained and more susceptible to lysing enzyme. Thus, the cell wall integrity may be changed, which may affect appressoria penetration in the △Cgscd1 mutant.

Judging from the intensity of the fluorescent signals at continuous time intervals during the early germination stage of conidia, we found that the signals in melanized appressoria were much stronger than the immature ones, indicating that CgSCD1 performs its function mainly in melanized appressoria. Interestingly, the fluorescence signals of PKS1 in C. graminicola were visible in immature appressoria but not in the mature appressoria with full melanization [20]. We could infer the reasons based on the melanin synthesis pathway: the PKS1 is involved in the first step of the melanin synthesis pathway, and performs its function at the earlier stage (immature appressoria), whereas CgSCD1, located downstream of the melanin synthesis pathway, performs its function mainly at the later stage (melanized appressoria). Of course, due to the diversity in fungal species, it may be necessary to characterize a PKS1 mutant of C. gloeosporioides in order to verify our hypothesis.
4. Materials and Methods

4.1. Fungal Strains and Culture Conditions

The strains used in this study were derived from the WT strain [22] (Table 1). With the exception of strains with specific requirements, all strains were cultured on potato dextrose agar (PDA) medium at 25 °C. The 2-day-old cultured mycelia grown on a cellophane-coated PDA Petri dishes were collected with a cell scraper for DNA and RNA extraction. The liquid medium used in the test was potato dextrose broth (PDB), which was shaken at 180 rpm at 25 °C. The conidia for appressorial formation assay and host inoculation were obtained by scraping 7-day cultures from PDA Petri dishes flooded with sterilized water.

Table 1. List of fungal and bacterial strains and plasmids used in this study.

| Title          | Description                                      |
|----------------|--------------------------------------------------|
| Strains        |                                                  |
| WT             | Wild-type strain EX2016-02                       |
| ΔCgscd1        | A knockout mutant of CgSCD1 gene, NrsR           |
| Cgscd1com      | Ectopic complementation mutant from ΔCgscd1 NrsR,HygR |
| Ag1l           | The strain used for Agrobacterium-mediated transformation |
| Plasmids       |                                                  |
| pNR2           | Contains nourseothricin resistance sequence       |
| pflu6          | Contains cyan fluorescent protein sequence        |
| pAg1-H3        | Agrobacterium-mediated transformation             |
| pAg1-H3-scd1-com| Contains CgSCD1 for complementation along with the CFP sequence |

4.2. Nucleotide Sequencing and Sequence Analysis

Genomic DNA (gDNA) and total RNA were extracted following the method described by Dubey [30]. The complementary DNA (cDNA) was obtained using Reverse Transcription Kit (Takara, Beijing, China). The CgSCD1 gene sequences of gDNA and cDNA were amplified by PCR amplification with the primers F and R (Table 2), respectively. Nucleotide sequencing was performed by Sangon (Sangon Biotech, Shanghai, China). The sequences were submitted to the GenBank database (accession number: MN539625, MN566451). Reference SCD1 protein sequences from other fungi were downloaded from the GenBank database. These SCD1 protein sequences, including OAG15977 in Alternaria alternata, CBF90107 in Aspergillus nidulans, XP_024547967 in Botrytis cinerea, AB004741 in Magnaporthe grisea, KHE84649 in Neurospora crassa, and EGY21953 in Verticillium dahliae, were aligned with MN566451 in C. gloeosporioides using DNAman 9.0 (Lynnon Biosoft, San Ramon, CA, USA). A phylogenetic tree based on this alignment was constructed using the neighbor-joining method in MEGA-X with 1000 bootstrap replicates. Conserved domains were predicted using NCBI Conserved Domain Database (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi).

Table 2. List of oligonucleotides used in this study.

| Oligonucleotides | Sequence 5’→3’                                   |
|------------------|--------------------------------------------------|
| 1F               | GTCCAGATTTTCACTTCAATTCCAG                     |
| 2R               | GCCCGAATCGGAATGCGGCTCTAATGTCGTACACATCTAACATCT |
| 3F               | TGATTACTGAAGATACAGATGCTAATGTGATGTGTCAGACTCAT  |
| 4R               | TTCTCTCAATCTTTACATCTTTACATCTTTAGAGTGAGTTCTCTT |
| F                | ATGGCGTCTCCTGTCTTGCAATACAAATACAAATACAAATACAA |
| R                | TTAGTGNCCGACGACAGACTCTGCTGCTGCTGCTGCTGCTGCTG |
| conF             | GATCGTACAGAGGAGGCTCTCAGGAGGAGGAGGAGGAGGAGGAGG |
| conR             | GCTCAAGACAGTGAGAAGGAGGAGGAGGAGGAGGAGGAGGAGG |
| N-F              | CTAGAGCCCGATTCCGCATTCCGCATTCCGCATTCCGCATTCCGC |
Table 2. Cont.

| Oligonucleotides | Sequence 5′→3′ |
|------------------|----------------|
| N-R             | AGGCTTGATATCTGTAGTTAATCA |
| H-F             | ATGAAAAGCCTGAATTCAGC |
| H-R             | TTCCCTTGCCCTTGAGACGAGTG |
| pAg-comF        | CGACGCGCAGTTGAATTCAGTGTCAGTCAGTTGACAGC |
| pAg-comR        | GAACATGTCGAGACGAGC |
| CFP-F           | CTCAAGTCTGCGGTGGCTCACATGGCCCTGTACAGTCAGA |
| CFP-R           | CCCGGCCCTTGGATACCGAGCTGCAGACGAGT |
| Qpcr-F          | GAAGGTCTCCGATAAGAGGTC |
| Qpcr-R          | CCGTACATCGATCTTCGTC |
| Actin-F         | AGCGGAAAGCCTTGACG |
| Actin-R         | TGGCTGATACCACATCTGACCCA |

4.3. Fungal Transformation

PEG-mediated protoplast transformation was used for the targeted deletion of CgSCD1, as described by Chung [31]. Fungal protoplasts were also prepared according to a method in that paper but with some modifications. Briefly, 2 mL conidial suspension (10^6 conidia per mL) was added into potato dextrose broth (PDB), which was shaken at 180 rpm for 24 h at 28 °C. The hyphae were harvested by filtering through four layers of cheesecloth, and then washed three times by sterilized water. The collected hyphae were digested in 30 mL permeation buffer (1.2 M KCl) containing 0.3 g of lysing enzyme from *Trichoderma harzianum* (L1412, Sigma-Aldrich, USA) at 28 °C with gentle shaking for 4 h. The digested solution was filtered through lens cleaning papers and centrifuged at 3000 × g for 10 min to harvest protoplasts. The protoplasts were suspended with 100 µL STC buffer (1.2 M sorbitol, 50 mM Tris-HCl, pH = 8, 50 mM CaCl2), mixed with 10 µg DNA fragments, and then placed on ice for 15 min. Then, 800 µL PEG (40% PEG 4000 in STC buffer) solution was added with gentle shaking at room temperature for 30 min, and then mixed with 50 mL of liquid regeneration medium (RM). RM was poured into Petri dishes using 10 mL per dish, and placed at 28 °C for 14 h of recovery. The 10 mL of PDA medium containing 100 µg/mL nourseothricin was poured into dishes for the screening of transformed mutants. After 3 days of incubation, the strains that grew on the surface of the PDA medium were transferred to new selective nourseothricin PDA Petri dishes.

4.4. Disruption of the CgSCD1 Gene and Verification

To generate the knockout construct of the CgSCD1 gene, three fragments were amplified, including upstream (1165 bp) of the CgSCD1 gene, downstream (1164 bp) of the CgSCD1 gene, and the nourseothricin resistance (NrsR) cassette (2.1 kb) as the selection marker. The primers for the CgSCD1 gene fragment amplification were designed according to the reference genome data of *C. gloeosporioides* strain Cg14. The primer information is listed in Table 1. The nourseothricin resistance (NrsR) cassette (2.1 kb) was amplified using the N-F and N-R primers from plasmid pNR2 [32]. The PCR mixture included 30 ng genomic DNA, 2U DNA polymerase, and PCR master mix (10149ES03, Yeasen Biotech, Shanghai, China). PCR was carried out with the following cycling program: 3 min at 98 °C, followed by 35 cycles of 98 °C for 10 s, 60 °C for 20 s, and 72 °C for 30 s, and a final extension at 72 °C for 5 min. The amplified fragments were isolated from agarose gel. The three fragments from the first-round PCR were fused by overlapping extensions in the second round of PCR. The products (4.3 kb) from the second round of PCR were used for transformation.

The deletion of the CgSCD1 gene was confirmed on two levels. At the genome level, the CgSCD1 gene and NrsR cassette were amplified to verify the deletion of CgSCD1 and the insertion of NrsR cassette, respectively. Specific primers were designed outside of the cassette region (conF and conR). The primer sets of conF and N-R were used to check for correct insertion at the 5′ region, and the primer sets of N-F and conR were used to check for correct insertion at the 3′ region. The PCR fragments, approximately 3.2 kb from the 5′ and 3′ regions, respectively, were sequenced for verification. At the
RNA level, the expression of CgSCD1 in ΔCgscd1 was tested and compared to WT and ectopic Cgscd1<com> by RT-PCR. The actin-like gene (Cglo_15775 from genomic dates of Cg14 strain) was used as a reference gene.

4.5. Complementation and Cyan Fluorescent Protein (CFP) Fusion Constructs

To confirm the phenotype of ΔCgscd1 conferred by deletion of the CgSCD1 gene, the complementation plasmid pAg1-H3-scd1-com was constructed based on plasmid pAg1-H3 [33]. Two fragments were inserted into the SacI restriction site of pAg1-H3 to generate the pAg1-H3-scd1-com construct. One was a fragment of the native promoter, along with the full-length gDNA fragment of the CgSCD1 gene, which was amplified using the primers pAg-comF and pAg-comR. The other was a fragment of the coding sequence of Anemonia majano cyan fluorescent protein (CFP) and Cochliobolus heterostrophus glyceraldehyde-3-phosphate dehydrogenase terminator sequence, which was amplified from plasmid pflu6 using the primers CFP-F and CFP-R. The CFP fusion was designed for the subcellular study of CgSCD1. The pAg1-H3-scd1-com construct was transformed into A. tumefaciens strain Agl1. The A. tumefaciens-mediated transformation (ATMT) procedure was based on a previously described protocol but with some modifications [34]. The complementation mutant Cgscd1<com> was selected on hygromycin medium and verified by PCR using primers H-F and H-R for the HygR cassette, and primers F and R for the CgSCD1 gene. The PCR cycles and temperature were as follows: 98 °C for 3 min; 35 cycles (98 °C, 10 s; 60 °C, 20 s; 72 °C, 30 s); and a final extension at 72 °C for 5 min.

4.6. Growth and Appressorial Formation Assay

To study the growth rate, the colony diameters of the WT, ΔCgscd1, and Cgscd1<com> strains (cultured as described above) were measured once a day for 5 days. To study the sporulation, the conidium was collected and calculated from 7 days of cultured strain on PDA Petri dishes. The germination rate and appressorium formation of the WT, ΔCgscd1, and Cgscd1<com> strains were performed on glass slides at 25 °C for 12 h of incubation. The conidial suspension (10^6 conidia per mL) of each strain was prepared and placed onto glass slides (20 µL droplets per slide, three replicates). The conidial germination and appressorium development were monitored using a microscope. The conidia were considered germinating when the germination tube was longer than the conidia. Using the WT strain as a control, appressorium formation was considered positive when appressoria were completely melanized.

4.7. Pathogenicity Assay

The fruits of two species, including tomato VF36 and key lime, were used to test the virulence of the WT, ΔCgscd1, and Cgscd1<com> strains. The fruits were stabbed using a sterile needle and wounds were inoculated with 20 µL of conidial suspension (10^6 conidia per mL) or 20 µL sterile water as a control. Ten healthy fruits were inoculated in each treatment. The fruits were kept at 25 °C with 90% relative humidity. The lesions of tomato and key lime were observed and measured at 7 and 21 days post-inoculation (dpi). Approximately 4-week-old detached tomato (VF36) leaves were used for droplet inoculation and incubated with 40 µL conidial suspension (8 × 5 µL droplets per leaf) at 25 °C with 90% relative humidity. DAB staining was used to observe the production of H2O2 in leaves at 36 hours post-inoculation (hpi) [35]. The inoculation method used for the onion epidermis test is detailed in [36]. Quantification of lactophenol cotton blue-stained fungal infection structures (germ tube, appressoria, infectious hyphae) after the infection was performed on glass slides at 24 hpi. A total of 100 conidia were counted for each strain.

4.8. Detection of In Vitro Scytalone Release from ΔCgscd1

To test whether the scytalone was released from the ΔCgscd1 mutant, the ΔCgscd1 and ΔBcpks12 mutants were separately inoculated in a symmetrical orientation about 22 mm from the edge of the dishes and cultured on PDA in Petri dishes (90 mm diameter) in darkness at 25 °C for 14 days. The color of the sclerotium was observed at the location where the hyphae were in contact.
4.9. Turgor Pressures Assay of Appressoria and Nile Red Stain

The conidia were inoculated on glass slides. When the conidium germinated and formed melanized appressoria, the medium was replaced with a dilution series of PEG6000 solutions, corresponding to an external turgor pressure ranging from 1 to 5 MPa [37]. The percentage of collapsed appressoria was quantified after 5 min of incubation in polyethylene glycol solution. Nile red stain solution and PBS buffer were configured as previously described, but with minor modifications [20]. Melanized appressoria on glass slides were prepared, the water was removed, and lysing enzyme solutions were added. Nile red staining was performed after 2 h of digestion. The slides were examined under a fluorescence microscope, with an excitation at 543 nm and emission at 619 nm. At least 100 conidia were counted per strain.

4.10. Gene Expression Analysis by qPCR

qPCR was performed using the Bio-Rad CFX96 Real-Time PCR Detection System (Bio-Rad, USA). PCR amplification was performed with 2 µL of cDNA template in a 25 µL reaction mixture containing 12.5 µL UltraSYBR Mixture (CW0957, Cwbio, Beijing, China) and 2 µL primers (Qpcr-F and Qpcr-R). PCR was carried out with the following cycling program: 95 °C for 10 min and 45 cycles of 95 °C for 30 s, 55 °C for 30 s, and 72 °C for 20 s. The expected qPCR product size was 250 bp. Three biological experiments were conducted. One representative set of results is presented as mean values of $2^{-\Delta\Delta CT} \pm \text{SE}$ for each treatment.

4.11. Subcellular Localization Analysis

The Cgscd1com conidial suspensions were obtained by scraping 7-day old cultures, grown in PDA, which were flooded with sterile water. These suspensions were placed onto glass-slides and incubated at 25 °C (three replicates of 20 µL droplets per slide). Cyan fluorescence was detected at 0, 4, and 12 h using a Zeiss laser scanning microscope (ZEISS, Germany). Images were taken under 40× objective and processed with ZEN 3.0 software. Cyan fluorescence in the hyphae of Cgscd1com was observed on the edge of the colony after 72 h of culture on PDA Petri dishes.

4.12. Statistical Analysis

The results are expressed as mean ± standard deviation (SD) and statistical analyses were carried out with SPSS software (SPSS, Chicago, USA). The statistical significance of the differences observed was analyzed by the Student’s t-test. Differences were considered significant at $p < 0.05$ (*).

Author Contributions: Conceptualization, L.X. and P.Z.; methodology, T.W. and D.R.; formal analysis, T.W., D.R., and X.C.; writing—original draft preparation, T.W.; writing—review and editing, H.N. and P.Z.; visualization, H.G.; project administration, L.X.; funding acquisition, L.X. All authors read and agreed to the published version of the manuscript.

Funding: This research was funded by the National Natural Science Foundation of China (No. 31571902 and 31972121) and the Shanghai Municipal Science and Technology Commission (No. 18391901400).

Acknowledgments: The authors sincerely thank Gang Liu and Xingzhong Liu (Institute of Microbiology, Chinese Academy of Sciences, Beijing, China) for providing strain Agl1 and plasmid pAg1-H3. They would also like to thank Weihua Tang (Institute of Plant Physiology and Ecology, Chinese Academy of Sciences, Shanghai, China) for kindly providing plasmid pflu6.

Conflicts of Interest: The authors declare no conflicts of interest.

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