Assignment of a dubious gene cluster to melanin biosynthesis in the tomato fungal pathogen Cladosporium fulvum

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Abstract

Pigments and phytoxins are crucial for the survival and spread of plant pathogenic fungi. The genome of the tomato biotrophic fungal pathogen Cladosporium fulvum contains a predicted gene cluster (CIPKS1, CIPRF1, CIRDT1 and CITSF1) that is syntenic with the characterized elsinochrome toxin gene cluster in the citrus pathogen Elsinoë fawcettii. However, a previous phylogenetic analysis suggested that CfPks1 might instead be involved in pigment production. Here, we report the characterization of the CIPKS1 gene cluster to resolve this ambiguity. Activation of the regulator CITSF1 specifically induced the expression of CIPKS1 and CIRDT1, but not of CIPRF1. These co-regulated genes that define the CIPKS1 gene cluster are orthologous to genes involved in 1,3-dihydroxynaphthalene (DHN) melanin biosynthesis in other fungi. Heterologous expression of CIPKS1 in Aspergillus oryzae yielded 1,3,6,8-tetrahydroxynaphthalene, a typical precursor of DHN melanin. Δcfpks1 deletion mutants showed similar altered pigmentation to wild type treated with DHN melanin inhibitors. These mutants remained virulent on tomato, showing this gene cluster is not involved in pathogenicity. Altogether, our results showed that the CIPKS1 gene cluster is involved in the production of DHN melanin and suggests that elsinochrome production in E. fawcettii likely involves another gene cluster.

Introduction

Secondary metabolites (SMs) are compounds produced by microbes, plants and insects that are often repurposed as medicines and pesticides. Equally important are SMs with harmful effects, such as mycotoxins and pathogenicity factors that poison animals or promote crop diseases. The vast majority of fungal SMs with a clear biological role in situ are pathogenicity or
virulence factors, also known as effector SMs, which are produced by plant pathogens during infection of their respective hosts [1,2]. Non-specific toxins synthesised by semi-biotrophic and necrotrophic fungi are compounds that necrotise host tissues indiscriminately, whilst host-selective toxins (HSTs) only cause necrosis on plants expressing corresponding susceptibility genes, thereby determining host range [2].

1,8-dihydroxynaphthalene (DHN) melanin is a virulence SM for several plant and human fungal pathogens. DHN melanin is required for the penetration of rice leaves by *Magnaporthe oryzae*, a process mediated by appressoria, dome-shaped cells specialized in piercing the plant cuticle and cell wall [3]. Failure to melanise the fungal cell wall results in immature appressoria that cannot generate the turgor pressure required to penetrate host tissues [4,5]. Tricyclazole, pyroquilon and other commercial compounds that inhibit DHN melanin biosynthesis are highly effective at preventing rice blast [6–8]. The same role in plant penetration was reported in several other plant pathogens, including *Colletotrichum kahawae* and *Diplocarpon rosae*, pathogens of coffee berries and roses, respectively [9,10]. It has been suggested that DHN melanin is also a photodynamic virulence factor used by *Pseudocercospora fijiensis*, the causal agent of black Sigatoka disease, to generate toxic reactive oxygen species during infection of banana [11]. In addition to its role in virulence, DHN melanin provides tolerance to many kinds of abiotic stresses, including radiation and extreme temperatures [9,10]. Accordingly, DHN melanin production is often linked to the development of survival structures. For example, DHN melanin accumulates in the cell wall of conidia and sclerotia of the plant pathogen *Botrytis cinerea*, but it does not play a role in the virulence of this pathogen [12]. In the plant endophytic fungus *Pestalotiopsis fici*, DHN melanin was recently shown to be essential for the development of multicellular conidia [13].

Fungal DHN is produced through a polyketide pathway, which starts with a non-reducing polyketide synthase (nrPKS) [14,15]. The first stable intermediate, 1,3,6,8-tetrahydroxynaphthalene (4THN), can be produced through three distinct biosynthetic routes. In Sordariomycetes such as *Colletotrichum lagenarium* [16,17], the nrPKS carries a bi-functional release domain that produces the hexaketide acetyl THN (ATHN) through Claisen ring closure, and then deacylates ATHN to release the pentaketide 4THN [18]. The nrPKS in the Eurotiomycete fungus *Exophiala dermatitidis* also releases ATHN [19], but the deacetylation step is instead performed by the discrete hydrolase, YG1 [20]. In other Eurotiomycetes fungi such as *Aspergillus* and *Penicillium* species, the nrPKS is a heptaketide synthase that releases YWA1 [21–23], which is deacylated by the hydrolase AYG1 to produce 4THN [21,24,25]. In certain fungal species like *B. cinerea*, two nrPKSs, likely one synthase with a bi-functional release domain (BcPks12) and one hexaketide or heptaketide synthase (BcPks13), are involved in DHN melanin biosynthesis [12]. The subsequent enzymatic steps to convert 4THN to DHN are common to all fungal species; 4THN is first reduced to scytalone by a 4HNR reductase, then dehydrated to 1,3,8-trihydroxynaphthalene (3THN) by the dehydratase SCD1 [15]. 3THN is reduced to vermelone by the reductase 3HNR, then dehydrated by SCD1 to yield DHN [15]. These reductions, especially 3HNR, are the target of tricyclazole [26]. Finally, DHN is polymerized into melanin by multicopper oxidases [27–30]. These different pathways have been invoked to explain the difference in pigmentation between brown-black fungi, including *C. lagenarium*, *M. oryzae* and *C. heterosporus*, and blueish-green fungi like *A. fumigatus* that might polymerize YWA1 in addition to DHN [30]. The genes encoding enzymes involved in DHN melanin biosynthesis and polymerization are organized in a gene cluster in *A. fumigatus* [30], and *Penicillium marneffei* [31], but they are partially clustered in *Alternaria alternata* and *Cochliobolus heterostrophus* [32,33] and tend to be dispersed in other fungal genomes [15].

It must be noted that certain fungal species produce another kind of melanin that is synthesized from L-3,4-dihydroxyphenylalanine (DOPA) through the action of tyrosinases and

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laccases [34]. Although the DHN melanin genes can be present in fungal genomes, the DOPA melanin pathway is the major route employed by certain fungal species, as exemplified by the pine needle pathogen Dothistroma septosporum [35].

Cladosporium fulvum is a non-obligate, biotrophic fungus that causes tomato leaf mold disease. *C. fulvum* shows limited filamentous growth on *in vitro* media in the dark, forming small sporulating colonies. They exhibit a green-brown colour, which was linked to the production of the pigment cladofulvin [36]. *C. fulvum* colonies harbour a grey colour when cladofulvin is not produced, which is likely due to the production of another pigment [36]. *C. fulvum* is known to reproduce asexually only and production of cladofulvin is primarily observed in conidia [37]. Despite a high potential chemical diversity with 23 predicted-functional SM core genes [38,39], the pigment cladofulvin produced by the *claG* gene cluster remains the only detectable SM [36,40]. It was suggested and later shown that the repression of cladofulvin biosynthetic genes is required for biotrophic growth of *C. fulvum* [37,38].

*CfPKS1* is another nrPKS core gene that shows a similar expression profile during infection of tomato leaves, i.e. downregulation [38]. *CfPKS1* belongs to a predicted gene cluster containing genes that encode a prefoldin chaperone (*CfPRF1*), a reductase (*CfRDT1*) and a transcription factor (*CfTSF1*) [38]. Previous comparative genomic analyses indicated that the *CfPKS1* gene cluster is homologous to the *Elsinoë fawcettii* gene cluster responsible for elsinochrome production, a light-activated toxin involved in the virulence of this pathogen on citrus hosts [41,42]. However, the phylogeny of *CfPKS1* suggested that it is also closely related to nrPKSs involved in DHN melanin biosynthesis [38]. In another study, *CfPKS1* was strongly up-regulated in the *C. fulvum* deletion mutant Δcfwor1 during growth on agar [43]. The hyper-black appearance of Δcfwor1 colonies and the absence of detectable SMs suggested that *CfPKS1* might be involved in the production of polymerized DHN melanin in *C. fulvum*. Such ambiguity between elsinochrome and DHN production remains unresolved.

Here, we report the functional characterization of the *CfPKS1* gene cluster by targeted gene deletion (*CfPKS1*), over-expression of the predicted local regulator (*CfTSF1*), and heterologous expression in *Aspergillus oryzae*. We provide chemical evidence of the pigment produced by this pathway and assessed the role of this compound in pathogenicity and biotrophic growth of *C. fulvum*.

**Results**

**Definition of the *CfPKS1* gene cluster in Cladosporium fulvum**

The *CfPKS1* gene cluster (Fig 1A) was initially predicted solely through its homology and synteny with the characterized elsinochrome gene cluster in *E. fawcettii*, as only minimal gene expression within this gene cluster had been observed during the growth of wild-type *C. fulvum* under diverse conditions [38]. The gene cluster includes *CfTSF1*, a gene predicted to encode a pathway-specific transcription factor [38,41]. To up-regulate and clearly define the *CfPKS1* gene cluster, wild-type *C. fulvum* was transformed with a plasmid containing *CfTSF1* fused to the promoter region of the nitrogen-regulated *C. fulvum* Avr9 gene [44]. The resulting OE.CfTSF1 transformant (S1 Fig) does not show any *in vitro* difference compared to wild type, but this transformant is not pathogenic on tomato (S2 Fig). Although random insertion of the expression cassette in a pathogenicity gene cannot be excluded, this loss of pathogenicity is likely due to the up-regulation of *CfTSF1* because the *Avr9* promoter induces high-expression when *C. fulvum* enters the plant and colonizes leaf tissues [38,44]. Both transformant and parental strain were grown in PDB and then induced in B5 medium without nitrogen (B5-N) for 48 hours to induce gene expression. Transcriptional profiling by RT-qPCR showed that the relative expression of *CfPKS1*, *CfTSF1* and *CfRDT1* was 1.7, 14.6 and 46.5-fold higher,
respectively, in the OE.CfTSF1 transformant than in wild type (each t-test P-value < 0.0001), whilst the predicted border genes are not co-regulated (Fig 1B). In contrast, CfPRF1 was not co-regulated and is therefore unlikely part of the CfPKS1 biosynthetic pathway. The up-regulation of the CfPKS1 gene cluster is specific to the over-expression of CfTSF1 because the gene
cluster is not significantly activated when the regulator of cladofulvin production (CfClaE; 40) is over-expressed (Fig 1B).

In a previous study, the CfPKS1 core gene was found to be highly up-regulated in Δcfwor1 deletion mutants [43]. The published RNA-seq data of this mutant confirms that all genes from the predicted gene cluster are co-regulated, but CfPRF1 and the predicted border genes are not (Fig 1C). Consistent with previous phylogenetic analyses of 4THN synthases [35,38,45], CfPKS1, CfRDT1 and CfTSF1 are all orthologous to genes involved in DHN melanin biosynthesis (Figs 2A–2C). In other fungi, this pathway involves three other genes, AYG1, 4HNRe and SCD1 [12,34]. Orthologues of these genes were identified on different scaffolds in the genome of C. fulvum (Table 1; Figs 2C–2E) and all are significantly up-regulated in the Δcfwor1 deletion mutants (Fig 1C). In contrast, paralogues of these genes are not differentially expressed or are significantly down-regulated in the Δcfwor1 deletion mutants (Fig 1C).

CfPks1 is a polyketide synthase that releases 4THN

CfPKS1 is orthologous to other characterized 4THN synthases from fungi of different orders (Fig 2A). In C. lagenarium, the 4THN synthase ClPks1 carries a bi-functional TE domain that releases 4THN [18]. CfPks1 was expressed heterologously in A. oryzae M-2-3 in order to determine whether or not it catalyses the same reactions as ClPks1. Ethyl acetate extracts of transformants contained three major products, 1–3, bearing UV signatures diagnostic of aromatic polyketides (Fig 3). Product 1 (RT = 4.9 min; UV max = 210, 261, 307 nm; m/z (ES−) 205 [M-H]−) was identified as flavolin by comparing its UV and mass spectra to published data (S3 and S4 Figs) and was confirmed by High-Resolution Mass Spectrometry (HRMS; exact mass 207.0283; S5A Fig). Flavolin is a spontaneously oxidised degradation product of 4THN. Product 2 (RT = 5.4 min; UV max = 244, 327 nm; m/z (ES−) 233 [M-H]−) harbours the same chemical formula as benzopyran according to HRMS (exact mass 235.0600; S5B Fig), which is a compound known as a shunt metabolite of the 4THN hexaketide pathway [18]. Product 3 (RT = 6 min; UV max = 244, 325 nm; m/z (ES−) 191 [M-H]−) was identified as 4THN by HRMS (exact mass 193.0498; S5C Fig). These results clearly show that CfPks1 produces the same intermediate as ClPks1 and thus exhibits the same enzymatic activity.

LC-MS analyses of organic extracts obtained from OE.CfTSF1 transformant after 48h growth on induction medium did not detect any precursor of DHN melanin. This observation could be due to a delay between the transcriptional induction and production of the compounds in significant amount as there was no pigmentation difference with wild type. Alternatively, it could suggest that DHN is efficiently polymerized, which is not easily extractable from cell walls with regular chemical methods.

CfPKS1 is needed for proper pigmentation of C. fulvum, but it is not required for pathogenicity on tomato

To confirm the heterologous expression results and obtain further insights in the biological role of the CfPKS1 gene cluster, the CfPKS1 gene was replaced by a deletion cassette containing the hygromycin resistance marker gene via homologous recombination (S1 Fig). Two confirmed independent deletion mutants and an ectopic transformant were selected for further analysis. Both Δcfpks1 deletion mutants were yellow-orange compared to the grey-green ectopic transformant and wild type (Fig 4A). The wild-type strain shows similar coloration to the Δcfpks1 deletion mutants when it is grown in the presence of pyroquilon, an inhibitor of the DHN melanin pathway (Fig 4A; [7]). In contrast, it was not coloured differently when grown in the presence of hydroquinone, an inhibitor of DOPA melanin (Fig 4A; [7]). These results contrast with the situation in D. septosporum, a close relative species of C. fulvum that
produces DOPA melanin despite the presence of *DsPKS1*, the orthologue of *CfPKS1* and other 4THN synthase genes [35]. Together, these results show that the polymerization of DHN melanin contributes to the pigmentation of *C. fulvum*. The Δcfpks1 mutants do not manifest any other obvious developmental or physiological defects.

Because DHN melanin is a pathogenicity factor in several plant pathogens, we addressed the possibility that this role was also true in *C. fulvum*. Tomato plants were inoculated with wild-type *C. fulvum*, an ectopic transformant and two independent Δcfpks1 deletion mutants.

Fig 2. Phylogenetic analysis of proteins involved in DHN melanin biosynthesis. Maximum likelihood phylogenetic trees were built to resolve the evolutionary relationship of (A) *CfPks1* to characterized non-reducing polyketide synthases; (B) *CfTsf1* to characterized regulators of DHN melanin biosynthesis; (C) *CfRdt1* and *CfRdt2* to reductases involved in the reduction of 1,3,8-trihydroxy- and 1,3,6,8-tetrahydroxy-naphthalene, respectively; (D) *CfAyg1* to characterized hydrolases involved in polyketide deacetylation; and (E) *CfScd1* to characterized scytalone dehydratases. *Elsinoe fawcettii* proteins assigned to the elsinochrome gene cluster are included [41]. Non-characterized homologues are also included for species in which the polyketide synthase involved in DHN synthesis is characterized. The name of characterized proteins and accession numbers (GenBank, SwissProt or Joint Genome Institute (JGI) protein ids) are indicated next to the species acronym. *Clafu*: *Cladosporium fulvum*; *Dotse*: *Dothistroma septosporum*; *Claph*: *Cladosporium phlei*; *Elsfa*: *Elsinoe fawcettii*; *Coche*: *Cochliobolus heterostrophus*; *Cocmi*: *Cochliobolus miyabeanus*; *Podan*: *Podospora anserina*; *Sorma*: *Sordaria macrospora*; *Magor*: *Magnaporthe oryzae*; *Colla*: *Colletotrichum lagenarium*; *Pesfi*: *Pestalotiopsis fici*; *Nodsp*: *Nodulisporium sp.*; *Glalo*: *Glarea locyozensis*; *Botci*: *Botrytis cinerea*; *Exode*: *Exophiala dermatitidis*; *Aspfl*: *Aspergillus flavus*; *Aspoc*: *Aspergillus ochraceus*; *Alta*: *Alternaria alternata*; *Ophfl*: *Ophiostoma fuscum*; *Penmar*: *Penicillium marneffei*; *Verda*: *Verticillium dahliae*.

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Whilst the growth of all strains was similar at 4 and 8 dpi, the \( \Delta cfpks1 \) deletion mutants had significantly outgrown the control strains by 12 dpi (Fig 4B). By 16 dpi, this difference in growth became strikingly clear, as the white-orange deletion mutants had colonized a greater leaf surface area than the control strains (Fig 4C).

**Discussion**

**The \( Cfpks1 \) gene cluster is involved in DHN melanin biosynthesis**

A previous phylogenetic analysis suggested that \( Cfpks1 \) is an orthologue of \( Epfs1 \), which encodes the nrPKS required for elsinochrome production [35,38,42]. The predicted elsinochrome gene cluster is also present in \( C. fulvum \), except for the putative transporter \( ECT1 \)

![CfPks1](https://doi.org/10.1371/journal.pone.0209600.g003)
However, the same phylogenetic analysis showed that CfPKS1 and EfPK1 are orthologous to 4THN synthases involved in the biosynthesis of DHN melanin. Similar observations were reported in other phylogenetic analyses [35,45]. Given that PKSs from a monophyletic clade tend to produce the same products, the different chemical structures of DHN melanin and elsinochrome contradict the phylogeny of their respective nrPKSs. The orthologous 4THN synthase in *C. lagenarium* has been characterized in detail; ClPks1 synthesizes the hexaketide 2-acetyl-1,3,6,8-tetrahydroxynaphthalene (ATHN) that is cyclized and deacylated by its bi-functional TE domain to give 4THN [18]. In this report, we showed that *C. fulvum* CfPks1 is also a synthase that releases 4THN because it produced both the hexaketide benzopyran and

**Fig 4. Characterization of *Cladosporium fulvum* Δcfpks1 deletion mutants.** (A) *In vitro* growth of wild-type *C. fulvum*, ectopic transformant and Δcfpks1 deletion mutants on Potato Dextrose Agar (PDA) medium; and wild-type *C. fulvum* on PDA supplemented with 30 mg.L\(^{-1}\) DHN biosynthesis inhibitor (pyroquilon) or DOPA melanin biosynthesis inhibitor (hydroquinone) [7]. (B) Quantification of fungal growth during tomato infection by wild type, an ectopic transformant and Δcfpks1 deletion mutants. (C) Tomato leaves infected by wild type, an ectopic transformant and Δcfpks1 deletion mutants at 16 days post-inoculation (dpi).

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the pentaketide 4THN (Fig 3). The precursor ATHN was not detected in our experiments, which might be due to the presence of an expressed homologue of the hydrolase Ayg1 in \textit{A. oryzae} that would efficiently deacetylate ATHN to yield 4THN [18, 20].

In characterized nrPKSs involved in DHN melanin biosynthesis, aromatisation of the nascent ACP-bound polyketide to give a monocyclic intermediate is catalysed by the product template (PT) domain [14,16,18,46]. Closure of the second ring and release of the polyketide requires the Claisen/Dieckmann cyclase class of TE domains (TE/CLC) [14,18,46]. The absence of a functional TE domain results in the release of pyrone shunt products \textit{via} spontaneous O-C cyclization [18]. The co-detection of 4THN and benzopyran in \textit{A. oryzae}:\textit{CfPKS1} suggests that the TE domain of \textit{CfPKS1} was not fully functional, which might be due to the absence of tailoring enzymes.

Ayg1 and its orthologues were shown to be involved in the deacetylation of a hexaketide or heptaketide precursor in the DHN melanin pathway described in blue/green fungi [24]. However, it was recently shown in the plant pathogen \textit{V. dahiae} that \textit{AYG1} might also be important for DHN biosynthesis in brown/black fungi [47]. Our finding that \textit{CfAYG1} is present in \textit{C. fulvum} genome and co-regulated with \textit{CfPKS1} suggests that this gene also plays a role in the DHN biosynthetic pathway in \textit{C. fulvum}, which requires further investigation. Although another parologue is present in \textit{C. fulvum}, it is not co-regulated with \textit{CfPKS1} and thus is unlikely to be involved in 4THN biosynthesis (Figs 1 and 2). It is noteworthy that \textit{Yg1} in \textit{E. dermatitidis} is actually a parologue of characterized Ayg1 in other fungal species (Fig 2). The true orthologue of Ayg1 in \textit{E. dermatitidis} must also be further investigated to ascertain its involvement in 4THN production.

Our phylogenetic analysis showed that co-regulated genes at the \textit{CfPKS1} locus in \textit{C. fulvum} and genes from the predicted elsinochrome gene cluster (\textit{PKS1}, \textit{RDT1} and \textit{TSF1}) in \textit{E. fawcettii} are orthologous to genes involved in DHN biosynthesis (Fig 2). A gene cluster for elsinochrome biosynthesis was recently characterized in \textit{Stagonospora nodorum} [45]. This gene cluster was shown to be related to gene clusters involved in the biosynthesis of cercosporin, a compound of the perylenequinone family that is structurally related to elsinochrome [48–50]. Together with our characterization of \textit{CfPKS1} as a 4THN synthase, these observations suggest that the elsinochrome gene cluster in \textit{E. fawcettii} has not been accurately assigned. Instead, a cercosporin-like gene cluster is certainly involved in the production of elsinochrome as reported in \textit{S. nodorum}. The fungus \textit{Cladosporium phlei} produces phleichrome, a perylenequinone that is structurally related to elsinochrome and cercosporin [48]. The \textit{C. phlei} nrPKS gene \textit{CpPks1}, orthologous to \textit{CfPKS1} and \textit{EfPKS1}, was assigned to phleichrome production [51]. Similarly, we suggest that \textit{CfPKS1} is involved in DHN melanin production and another cercosporin-like nrPKS is responsible for phleichrome production. Such false assignments suggest that crosstalk and interdependencies between gene clusters might be more important than previously thought [52].

**DHN melanin is not a virulence factor in \textit{C. fulvum}**

The biosynthesis of fungal DHN melanin has been extensively studied because of its diverse roles in fungal biology [15]. Melanin contributes to virulence in animal and plant pathogens, with the latter linked to the formation of host-invading appressoria [15]. In species that do not produce appressoria, melanin confers resilience to chemical and abiotic stresses [12]. \textit{C. fulvum} does not differentiate appressoria, which likely explains why DHN melanin is not involved in the pathogenicity of this fungus. The pigmentation of \textit{C. fulvum} relies on two compounds, the greyish DHN melanin and yellow-orange-purple (depending on pH) cladofulvin [36,40]. Cladofulvin is also not produced during plant infection, but instead protects the
fungus from environmental stresses such as UV light and cold temperatures [37]. Although untested, DHN melanin likely plays a similar role in the survival of *C. fulvum ex planta*.

Although further investigation is required to fully exclude that the expression cassette has inserted within a pathogenicity gene, the observed loss of pathogenicity of the *OE.CfTSF1* transformant suggests that adequate regulation and downregulation of DHN melanin production during leaf colonization is required for full pathogenicity in *C. fulvum*. In *A. fumigatus* and *A. nidulans*, melanin biosynthesis is initiated in endosomes that carry the enzymes that produce DHN, which is then polymerized within the cell wall by multicopper oxidase and laccase enzymes [53]. Melanin is polymerised in the cell wall, forming several layers of globular particles that grow thicker over time, strengthening the cell wall [54]. It is accepted that hyphal tip elongation requires enzymes to weaken the cell wall in order to incorporate new components, which are then cross-linked to rigidify the cell wall [55]. Thus, the abnormal accumulation of melanin in fungal cell walls is likely to modify its physical properties and increases its rigidity. We observed that the runner hyphae of *C. fulvum OE.TSF1* transformants branched infrequently and the few successful penetration events resulted in colonizing hyphae that rapidly stopped growing (S2 Fig). This phenotype can be explained by the elevated production of DHN melanin and its abnormal accumulation in cell walls, which might reduce sensitivity to chemotactic gradients and then arrest growth of colonizing hyphae by preventing essential fungal tip remodelling [55]. An *in planta* analysis of fungal cell walls would address this hypothesis. Alternatively, DHN melanin accumulating in the cell wall could be recognized by plant cells, leading to the activation of plant defences and resistance. Similar *in planta* activation might be detrimental to the virulence of other plant pathogens irrespective of whether or not DHN melanin is a pathogenicity factor.

**Conclusion**

Using complementary approaches (gene expression, phylogeny, heterologous expression), our study confidently assigned the *CfPKS1* gene cluster to DHN melanin in *C. fulvum*. It suggests that orthologous gene clusters in other species have been wrongly assigned to toxin production, including elsinochrome in *E. fawcettii* and phleichrome in *C. phlei*. Further investigations in these fungal species are needed to address this ambiguity and are likely to provide important insights on pathway crosstalk that might lead to incorrect gene cluster assignment.

**Experimental procedures**

Most of the methods were performed as described in Griffiths et al. (2016) and Griffiths et al. (2018) [37,40].

**Fungal strains employed in this study**

*C. fulvum* 0WU [39] was the parental strain used to perform transformation and gene deletion experiments. *C. fulvum* was grown on potato dextrose agar (PDA) plates at 20˚C in the dark. For inhibitor experiments, PDA medium was supplemented with 30 mg.L⁻¹ pyroquilon or hydroquinone (Sigma-Aldrich, Zwijndrecht, The Netherlands). *A. oryzae* M-2-3 strain was used to perform heterologous expression [56].

**Phylogeny**

The protein sequence of *CfPks1* and of selected characterized nrPKSs from groups II, III and IV [57] were aligned using Muscle [58] and poorly aligned regions of the alignment were removed using Gblocks, with half allowed gap positions for Ayg1, Rdt and Scd1 alignments, and allowing
all gapped positions for the Tsf1 alignment [59]. Maximum-likelihood phylogeny was calculated using PhyML 3.1 [60] with the LG+G+I substitution model as determined by Modelgenerator v851 [61] and SH approximate likelihood ratio test to evaluate branch support.

Non-characterized homologues were retrieved from the Joint Genome Institute MycoCosm portal (genome.jgi.doe.gov; [62]) using BlastP (with default parameters) [63]. All Homologues of 4Hnr (AAG29497.2) and Sdh1 (BAA34046.1) from M. oryzae and of Ayg1 (AAF03354.1) from A. fumigatus were sought in the predicted proteome of C. fulvum. Homologues of CfRdt1, CfTsf1, CfRdt2, CfScd1 and CfAyg1 were sought in selected genomes. For each protein, retrieved homologues and characterized proteins were aligned using Muscle; sequences with large deletions or insertions were manually removed. Neighbour-Joining phylogenetic trees were built with the JTT substitution model using MEGA 7 [64] in order to ascertain orthology. Identified orthologues were then aligned again and analysed following the same process as for CfPks1 described above, but using the LG+G substitution model for Ayg1 and Scd1 trees.

**Generation of OE.CfTSF1 and OE.CfTSF1::GFP transformants**

The putative local regulator from the CfPKS1 gene cluster, CfTSF1, was amplified by PCR using Phusion Flash High-Fidelity PCR Master Mix (Life Technologies, Carlsbad, CA) from C. fulvum genomic DNA using the primer pair PacI_CfTSF1_Forward and NotI_CfTSF1_Reverse (S1 Table). Plasmid pFBTS3 contains the promoter of the nitrogen-regulated Avr9 gene [36,44]. The CfTSF1 amplicon and pFBTS3 were cut using PacI and NotI restriction enzymes (Fermentas Fast Digest, Waltham, MA), cleaned with Zymogen DNA Clean & Concentrator (Baseclear, Leiden, The Netherlands), and ligated using T4 DNA polymerase (Promega, Madison, WI) to yield pFBTS3-CfTSF1. Escherichia coli DH5α cells were transformed with the ligation mix using a standard heat-shock protocol and transformants were selected in lysogeny broth (LB)-kanamycin agar (50 μg.ml⁻¹). Plasmids were extracted from transformants and screened by restriction digest analysis using PacI and NotI in a double digestion. A plasmid bearing the correct restriction pattern was sent to Macrogen (Amsterdam, The Netherlands) for sequencing of the insert. Agrobacterium tumefaciens AGL1 was transformed with pFBTS3-CfTSF1 by electroporation, and plated on LB-kanamycin agar (50 μg.ml⁻¹). One positive transformant was picked, verified and named AT-pFBTS3-CfTSF1. This plasmid was introduced to C. fulvum using A. tumefaciens-mediated transformation as previously described [65]. Transformants were selected on PDA medium supplemented with hygromycin (100 μg.ml⁻¹). Several transformants and wild-type C. fulvum were grown for 5 days in potato-dextrose broth (PDB; Oxoid, Altrincham, UK) and then transferred to Gamborg B5 medium without nitrogen (B5-N) in order to induce the Avr9 promoter [37,44]. Total RNA was extracted and cDNA synthesis was performed as previously described [65]. The induction of the CfPKS1 biosynthetic gene cluster was confirmed by RT-qrtPCR using primers listed in S1 Table. One transformant showing the expected strong induction of genes at the CfPKS1 locus was selected and named OE.CfTSF1.

Using the same methods, A. tumefaciens AGL1 was transformed with plasmid pRM254, which contains GFP and geneticin-resistance genes [66] to yield AT-pRM254 strain. The plasmid was introduced into the OE.CfTSF1 transformant as described above. Transformants were selected on PDA medium supplemented with geneticin (100 μg.ml⁻¹). Transformants were picked and screened for GFP fluorescence. One transformant was selected and named OE.CfTSF1::GFP.

**Generation of Δcfpks1 deletion mutants**

The plasmid for targeted gene replacement of CfPKS1 was generated following the same procedure as described in Griffiths et al. (2016) [40]. The upstream (US) and downstream (DS)
regions flanking of \(CfPKS1\) were amplified using primers 1 + 2 and 3 + 4, respectively (S1 Table), and cloned into \(pDONR-P4-P1R\) and \(pDONR-P2-P3\) vectors. The final gene replacement plasmid was assembled in a LR reaction (Invitrogen) that combined the \(pDONR-P4-P1R::US\_CfPKS1,\ pDONR-P2-P3::DS\_CfPKS1,\ p221\_GFP\_HYG\ (pDONR containing a cassette with GFP and HYG resistance marker genes) and the destination vector \(pDEST\_R4-R3\) [40,66]. One correct sequenced plasmid was chosen and named \(pDest43\_Δcfpks1\). This plasmid was introduced into \(C.\ fulvum\) 0WU using the \(A.\ tumefaciens\) transformation method as described in Okmen et al. (2013) [65]. Transformants were selected on PDA plates containing hygromycin (100 \(μg.mL^{-1}\)). Genomic DNA of each strain was isolated using a Zymo Research Genomic DNA Clean & Concentrator™ (Baseclear), according to the manufacturer’s recommendations. PCR and quantitative real-time PCR were performed to screen for double crossovers and measure the number of inserted deletion cassettes, respectively (S1 Fig and S1 Table).

**Plant inoculation and determination of fungal growth**

Inoculation of tomato with \(C.\ fulvum\) wild-type, deletion mutant and transformant strains was carried out according to a previously described method [67]. To determine fungal growth, the fourth composite leaf of infected tomato plants was harvested at 4, 8, and 12 days post-inoculation (dpi) and flash frozen in liquid nitrogen. Samples were ground to a fine powder in liquid nitrogen, and total RNA was extracted from 100 \(mg\) of material using a Zymogen Direct-zol RNA MiniPrep kit (Baseclear) according to the manufacturer’s recommended protocol. cDNA synthesis was performed using 100–2,000 \(ng\) of total RNA and M-MLV reverse transcriptase (Promega), following the manufacturer’s protocol. To assess \(C.\ fulvum\) growth during infection, the \(actin\) gene of this fungus was targeted by qrtPCR using the \(Cf-actin\_RT-qrtPCR\_F/Cf-actin\_RT-qrtPCR\_R\) primer pair method [67]. For sample calibration, the \(Solanum\ lycopersicum\ actin\) gene was targeted using the \(Sl-actin\_qrtPCR\_F/Sl-actin\_qrtPCR\_R\) primer pair method [67]. The same cDNA samples were used to measure the expression of genes at the \(CfPKS1\) locus by qrtPCR using previously reported methods and primers [38]. Additional oligonucleotides (S1 Table) were designed and their efficiency determined as described in [38]. Results were analyzed according to the \(2^{ΔCt}\) method [68] and are the average of three biological replicates.

**Microscopic examination of GFP-expressing strains**

Imaging of infected tomato leaves was performed using a spinning disc confocal microscope (Nikon Ti microscope body (Shinagawa, Tokyo, Japan), Yokogawa CSUX1 scanner (Musashino, Tokyo, Japan), Photometrics Evolve camera (Tucson, AZ), Metamorph software (Molecular Devices, Sunnyvale, CA), 491 nm laser line; 60x oil 1.40NA objective). Z-stacks were acquired with an internal spacing of 0.5 \(μm\). All images were processed using Fiji software [69].

**Construction of vectors for heterologous expression and generation of A. oryzae M-2-3 transformants**

The cloning of \(CfPKS1\) in heterologous expression vectors was performed as described in Griffiths et al. (2016) [40]. Briefly, \(CfPKS1\) was amplified from \(C.\ fulvum\ OE.CfTSF1\) transformant cDNA (grown on B5-N medium) by PCR using primers 5 + 6 (S1 Table) and cloned into \(NotI\)-linearized \(pEYA2\) using recombination in \(S.\ cerevisiae\) BMA 64 to generate plasmid \(pEYA2-CfPKS1\) [40,53]. \(CfPKS1\) was transferred into the expression vector \(pTAex3GS\) using LR clonase (Invitrogen), and the resulting \(pTAex3GS-CfPKS1\) plasmid was introduced in \(A.\ oryzae\) M-2-3.
oryzae M-2-3 using PEG-mediated transformation as described in Griffiths et al. (2016) [40]. The starch-inducible taka-amylase promoter (PamyB) controls the expression of CfPKS1. The final vector contains the arginine biosynthesis gene (argB) for selection of fungal transformants.

Secondary metabolite extraction and analysis by LC-MS

Selected A. oryzae transformants containing CfPKS1 were grown on dextrose-peptone-yeast extract (DPY) agar plates at 30˚C until the whole plates were covered. The cultures were freeze-dried and then homogenised with a pestle and mortar. The homogenate was resuspended in water, acidified to pH4 with HCl, and then twice extracted with ethyl acetate. The organic phase was recovered and dried under a nitrogen flow. Samples were resuspended in acetonitrile (CH₃CN), centrifuged at 20,000 x g for 5 min in a microcentrifuge tube and then transferred to a 1 mL clear glass shell vial (WAT025054c).

HPLC analysis with a Waters Symmetry reverse phase 5 μm, C18, 100 Å column (WAT046980) was carried out on a Waters 600S system. The sample was eluted with a variable gradient of solvents (A) H₂O and (B) CH₃CN (both containing 0.1% trifluoroacetic acid) at a flow rate of 1 mL.min⁻¹. The following gradient was used: 0 min, A (95%); 10 min, A (10%); 12 min, A (10%), 15 min, A (0%), 16 min, A (95%), 20 min, A (95%). UV spectra were obtained using a 996-photodiode array (PDA) detector and analysed with the Waters Empower software.

LC-MS data were obtained using a Waters LC-MS system composed of a Waters 2767 autosampler, Waters 2545 pump system, a Phenomenex Kinetex column (2.6 μm, C18, 100 Å, 4.6 × 100 mm) equipped with a Phenomenex Security Guard precolumn (Luna C5 300 Å) eluted at 1 mL.min⁻¹. Detection was by Waters 2998 Diode Array detector between 200 and 400 nm and Waters SQD-2 mass detector operating simultaneously in ES+ and ES- modes between 100 m/z and 650 m/z. Solvents were: A, HPLC grade H₂O + 0.05% formic acid; B, HPLC grade MeOH containing 0.045% formic acid; and C, HPLC grade CH₃CN containing 0.045% formic acid. Gradients were as follows: Kinetex/ CH₃CN: 0 min, 10% C; 10 min, 90% C; 12 min, 90% C; 13 min, 10% C; 15 min, 10% C. Samples were generally diluted to 1 mg.mL⁻¹ and 10 μL injected (10 μg). Data capture and analysis, including peak integration, was performed using MassLynx 4.1 software (Waters).

Semi-preparative LC-MS, compound purification and structure determination

Purification of compounds was achieved using a Waters mass-directed autopurification system comprising a Waters 2767 autosampler, Waters 2545 pump system, a Phenomenex Kinetex Axia column (5 μm, C18, 100 Å, 21.2 × 250 mm) equipped with a Phenomenex Security Guard precolumn (Luna C5 300 Å) eluted at 20 mL.min⁻¹ at ambient temperature. Solvent A, HPLC grade H₂O + 0.05% formic acid; Solvent B, HPLC grade CH₃CN + 0.045% formic acid. The post-column flow was split (100:1) and the minority flow was made up with HPLC grade MeOH + 0.045% formic acid to 1 mL.min⁻¹ for simultaneous analysis by diode array (Waters 2998) and ESI mass spectrometry in positive and negative modes (Waters SQD- 2). Detected peaks were collected into glass test tubes. Combined tubes were evaporated under a flow of dry N₂ gas and weighed. HRMS data were measured using Waters Q-Tof Premier operating in ES⁺ mode.

Supporting information

S1 Fig. Molecular characterization of Δcfpks1 deletion mutants and OE.CfTSF1 transformant.
(PNG)
S2 Fig. Pathogenicity assay of the *Cladosporium fulvum* OE.CfTSF1 transformant and microscopic observation of the OE.CfTSF1::GFP over-expression transformant on tomato leaves.
(PNG)

S3 Fig. Diode array chromatograms (left) and Total Ion Chromatograms (TICs; right) of ethyl acetate extracts from *Aspergillus oryzae* transformants expressing CfPks1.
(PNG)

S4 Fig. UV and MS spectra of major products produced by *Aspergillus oryzae* transformants expressing CfPks1.
(PNG)

S5 Fig. High Resolution Mass Spectrometry (HRMS) data measured for products produced by *Aspergillus oryzae* transformants expressing CfPks1.
(PNG)

S1 Table. Oligonucleotides used in this study.
(DOCX)

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References

1. Collemare J, Lebrun M-H. Fungal secondary metabolites: ancient toxins and novel effectors in plant–microbe interactions. In: Martin F, Kamoun S, editors. Effectors in Plant–Microbe Interactions. Oxford: Wiley-Blackwell; 2011. pp. 377–400.

2. Stergiopoulos I, Collemare J, Mehrabi R, de Wit PJ. Phytotoxic secondary metabolites and peptides produced by plant pathogenic *Dothideomycete* fungi. *FEMS Microbiol. Rev.* 2013; 37:67–93. https://doi.org/10.1111/1574-6976.2012.00349.x PMID: 22931109
3. Howard RJ, Valent B. Breaking and entering: host penetration by the fungal rice blast pathogen Magnaporthe grisea. Annu. Rev. Microbiol. 1996; 50:491–512. https://doi.org/10.1146/annurev.micro.50.1.491 PMID: 8905089

4. Chumley F, Valent B. Genetic analysis of melanin-deficient, nonpathogenic mutants of Magnaporthe grisea. Mol. Plant-Microbe Interact. 1990; 3:135–143.

5. Howard RJ, Ferrari MA, Roach DH, Money NP. Penetration of hard substrates by a fungus employing enormous turgor pressures. Proc. Natl. Acad. Sci. USA. 1991; 88:11281–11284. PMID: 1837147

6. Chen H, Han X, Qin N, Wei L, Yang Y, Rao L, et al. Synthesis and biological evaluation of novel inhibitors against 1,3,8-trihydroxynaphthalene reductase from Magnaporthe grisea. Bioorg. Med. Chem. 2016; 24:1225–30. https://doi.org/10.1016/j.bmc.2016.01.053 PMID: 26860927

7. Lee JK, Jung HM, Kim SY. 1,8-dihydroxynaphthalene (DHN)-melanin biosynthesis inhibitors increase erythritol production in Torula corallina, and DHN-melanin inhibits erythrose reductase. Appl. Environ. Microbiol. 2003; 69:3427–34. https://doi.org/10.1128/AEM.69.6.3427-3434.2003 PMID: 12788746

8. Woloshuk CP, Sisler HD, Vigil EL. Action of the antipenetrant, tricyclazole, on appressoria of Pyricularia oryzae. Physiol. Plant Pathol. 1983; 22:245–259.

9. Chen Z, Nunes MA, Silva MC, Rodrigues C.J. Jr. Appressorium turgor pressure of Colletotrichum kahawae might have a role in coffee cuticle penetration. Mycologia. 2004; 96:1199–1208. PMID: 21148942

10. Gachomo EW, Seufferheld MJ, Kotcheno SO. Melanization of appressoria is critical for the pathogenicity of Diplodia rosea. Mol. Biol. Rep. 2010; 37:3583–3591. PMID: 20204524

11. Beltrán-García MJ, Prado FM, Oliveira MS, Ortiz-Mendoza D, Scalfo AC, Pessoa A Jr, et al. Single-tissue molecular oxygen generation by light-activated DHN-melanin of the fungal pathogen Mycosphaerella fijiensis in black Sigatoka disease of bananas. PLoS One. 2014; 9:e91616. https://doi.org/10.1371/journal.pone.0091616 PMID: 24646830

12. Schumacher J. DHN melanin biosynthesis in the plant pathogenic fungus Botrytis cinerea is based on two developmentally regulated key enzyme (PKS)-encoding genes. Mol. Microbiol. 2016; 99:729–48. https://doi.org/10.1111/mmi.12626 PMID: 26514268

13. Zhang P, Wang X, Fan A, Zheng Y, Liu X, Wang S, et al. A cryptic pigment biosynthetic pathway uncovered by heterologous expression is essential for conidial development in Pestalotiopsis fici. Mol Microbiol. 2017; 105:469–483. https://doi.org/10.1111/mmi.13711 PMID: 28517364

14. Cox RJ. Polyketides, proteins and genes in fungi: Programmed nano-machines begin to reveal their secrets. Org. Biomol. Chem. 2007; 5:2010–2026. https://doi.org/10.1039/b704420h PMID: 17581644

15. Langfelder K, Streibel M, Jahn B, Haase G, Brakhage AA. Biosynthesis of fungal melanins and their importance for human pathogenic fungi. Fungal Genet. Biol. 2003; 38:143–58. PMID: 12620252

16. Fujii I, Mori Y, Watanabe A, Kubo Y, Tsugi G, Ebizuka Y. Heterologous expression and product identification of Colletotrichum lagenarium polyketide synthase encoded by the PKS1 gene involved in melanin biosynthesis. Biosci. Biotechnol. Biochem. 1999; 63:1445–52. https://doi.org/10.1271/bbb.63.1445 PMID: 10501004

17. Takano Y, Kubo Y, Shimizu K, Mise K, Okuno T, Furusawa I. Structural analysis of PKS1, a polyketide synthase gene involved in melanin biosynthesis in Colletotrichum lagenarium. Mol. Gen. Genet. 1995; 249:162–7. PMID: 7500937

18. Vagsfad AL, Hill EA, Labonte JW, Townsend CA. Characterization of a fungal thioesterase having Claisen cyclase and deacetylase activities in melanin biosynthesis. Chem. Biol. 2012; 19:1525–34. https://doi.org/10.1016/j.chembiol.2012.10.002 PMID: 23261597

19. Feng B, Wang X, Hauser M, Kaufmann S, Jentsch S, Haase G, et al. Molecular cloning and characterization of WdPKS1, a gene involved in dihydroxynaphthalene melanin biosynthesis and virulence in Wangiella (Exophiala) dermatitidis. Infect. Immun. 2001; 69:1781–94. https://doi.org/10.1128/IAI.69.3.1781-1794.2001 PMID: 11179356

20. Wheeler MH, Abramczyk D, Puckhaber LS, Naruse M, Ebizuka Y, Fujii I, Szansizlo PJ. New biosynthetic step in the melanin pathway of Wangiella (Exophiala) dermatitidis: evidence for 2-acetyl-1,3,6,8-tetrahydroxynaphthalene as a novel precursor. Eukaryot Cell. 2008; 7:1699–711. https://doi.org/10.1128/EC.00179-08 PMID: 18676950

21. Chiang YM, Meyer KM, Praseuth M, Baker SE, Bruno KS, Wang CC. Characterization of a polyketide synthase in Aspergillus niger whose product is a precursor for both dihydroxynaphthalene (DHN) melanin and naphtho-γ-pyrene. Fungal Genet. Biol. 2011; 48:430–7. https://doi.org/10.1016/j.fgb.2010.12.001 PMID: 21176790

22. Tsai HF, Chang YC, Washburn RG, Wheeler MH, Kwon-Chung KJ. The developmentally regulated alb1 gene of Aspergillus fumigatus: its role in modulation of conidial morphology and virulence. J. Bacteriol. 1998; 180:3031–8. PMID: 9620950
23. Watanabe A, Fuji I, Tsai H, Chang YC, Kwon-Chung KJ, Ebizuka Y. *Aspergillus fumigatus alb1* encodes naphthopyrone synthase when expressed in *Aspergillus oryzae*. FEMS Microbiol. Lett. 2000; 192:39–44. https://doi.org/10.1111/j.1574-6968.2000.tb09356.x PMID: 11040426

24. Fuji I, Yasuoka Y, Tsai HF, Chang YC, Kwon-Chung KJ, Ebizuka Y. Hydrolytic polyketide shortening by agy1p, a novel enzyme involved in fungal melanin biosynthesis. J. Biol. Chem. 2004; 279:44613–20. https://doi.org/10.1074/jbc.M406758200 PMID: 15310761

25. Tsai HF, Fuji I, Watanabe A, Wheeler MH, Chang YC, Yasuoka Y, et al. Pentaketide melanin biosynthesis in *Aspergillus fumigatus* requires chain-length shortening of a heptaketide precursor. J. Biol. Chem. 2001; 276:29292–8. https://doi.org/10.1074/jbc.M101998200 PMID: 11350964

26. Thompson JE, Fahnstock S, Farrall L, Liao DJ, Valent B, Jordan DB. The second naphthol reductase of fungal melanin biosynthesis in *Magnaporthe grisea*: tetrahydroxynaphthalene reductase. J Biol Chem. 2000; 275:34867–72. https://doi.org/10.1074/jbc.M006659200 PMID: 10956664

27. Lin SY, Okuda S, Ikeda K, Okuno T, Takano Y. LAC2 encoding a secreted laccase is involved in appressorial melanization and conidial pigmentation in *Colletotrichum orbiculare*. Mol. Plant Microbe Interact. 2012; 25:1552–61. https://doi.org/10.1094/MPMI-05-12-0131-R PMID: 22934563

28. Saitoh Y, Izumitsu K, Morita A, Shimizu K, Tanaka C. ChMCO1 of *Coelhobulus heterostrophus* is a new class of metallo-oxidase, playing an important role in DHN-melanization. Mycotechnology. 2010; 51:327–336.

29. Sugareva V, Härtl A, Brock M, Hübner K, Rohde M, Heinekamp T, et al. Characterisation of the laccase-encoding gene abz2 of the dihydroxynaphthalene-like melanin gene cluster of *Aspergillus fumigatus*. Arch. Microbiol. 2006; 186:345–55. https://doi.org/10.1007/s00203-006-0144-2 PMID: 16988817

30. Tsai HF, Wheeler MH, Chang YC, Kwon-Chung KJ. A developmentally regulated gene cluster involved in conidial pigment biosynthesis in *Aspergillus fumigatus*. J. Bacteriol. 1999; 181:6469–77. PMID: 10519399

31. Woo PC, Tam EW, Chong KT, Cai JJ, Tung ET, Ngan AH, et al. High diversity of polyketide synthase genes and the melanin biosynthesis gene cluster in *Penicillium marneffei*. FEBS J. 2010; 277:3750–8. https://doi.org/10.1111/j.1742-4658.2010.07776.x PMID: 20718860

32. Kimura N., Tsuge T. Gene cluster involved in melanin biosynthesis of the filamentous fungus *Alternaria alternata*. J. Bacteriol. 1993; 175:4427–35. PMID: 8392512

33. Fetzner R, Seither K, Wenderoth M, Herr A, Fischer R. *Alternaria alternata* transcription factor CmrA controls melanization and spore development. Microbiology. 2014; 160:1845–54. https://doi.org/10.1099/mi.0.079046-0 PMID: 24972701

34. Teichert I, Nowrousian M. Evolution of genes for secondary metabolism in fungi. In: Pöggeler S, Wöstehoff J, editors. The Mycota XIV, Evolution of fungi and fungal-like organisms. Springer-Verlag Berlin Heidelberg: 2011. pp. 231–255.

35. Ozturk IK, Chettri P, DupONT PY, Barnes I, McDougall RL, Moore GG, et al. Evolution of polyketide synthase genes and the melanin biosynthesis gene cluster in *Penicillium marneffei*. FEBS J. 2010; 277:3750–8. https://doi.org/10.1111/j.1742-4658.2010.07776.x PMID: 20718860

36. Griffiths S, Saccomanno B, de Wit PJ, Collemare J. Regulation of secondary metabolite production in the fungal tomato pathogen *Cladosporium fulvum*. Fungal Genet. Biol. 2015; 84:52–61. https://doi.org/10.1016/j.fgb.2015.09.009 PMID: 26415644

37. Griffiths S, Mesarich CH, Overdijk EJR, Saccomanno B, de Wit PJGM, Collemare J. Down-regulation of cladofulvin biosynthesis is required for biotrophic growth of *Cladosporium fulvum* on tomato. Mol Plant Pathol. 2018; 19:369–380. https://doi.org/10.1111/mpp.12527 PMID: 27997759

38. Collemare J, Griffiths S, Iida Y, Karimi Jashni M, Battaglia E, Cox RJ, et al. Secondary metabolism and biotrophic lifestyle in the tomato pathogen *Cladosporium fulvum*. PLoS One. 2014; 9:e85877. https://doi.org/10.1371/journal.pone.0085877 PMID: 24465762

39. de Wit PJ, van der Burgt A, Ökmen B, Stergiopoulos I, Abd-Elsalam KA, Aerts AL, et al. The genomes of the fungal plant pathogens *Cladosporium fulvum* and *Dothistroma septosporum* reveal adaptation to different hosts and lifestyles but also signatures of common ancestry. PLoS Genet. 2012; 8:e1003088. https://doi.org/10.1371/journal.pgen.1003088 PMID: 23209441

40. Griffiths S, Mesarich CH, Saccomanno B, Vaisberg A, de Wit PJ, Cox R, et al. Elucidation of cladofulvin biosynthesis reveals a cytochrome P450 monoxygenase required for arthraquinone dimerization. Proc. Natl. Acad. Sci. USA. 2016; 113:6851–6. https://doi.org/10.1073/pnas.1603528113 PMID: 27274078

41. Chung KR, Liao HL. Determination of a transcriptional regulator-like gene involved in biosynthesis of elsinochrome phytotoxin by the citrus scab fungus, *Elsinoë fawcettii*. Microbiology. 2008; 154:3556–66. https://doi.org/10.1099/mic.0.01941-0 PMID: 18957608
42. Liao HL, Chung KR. Genetic dissection defines the roles of elsinochrome phytotoxin for fungal pathogenesis and conidiation of the citrus pathogen Elsinoe fawcettii. Mol. Plant Microbe Interact. 2008; 21:469–79. https://doi.org/10.1094/MPMI-21-4-0469 PMID: 18321192

43. Ökmen B, Collemare J, Griffiths S, van der Burgt A, Cox R, de Wit PJ. Functional analysis of the conserved transcriptional regulator CfWor1 in Cladosporium fulvum reveals diverse roles in the virulence of plant pathogenic fungi. Mol. Microbiol. 2014; 92:10–27. https://doi.org/10.1111/mmi.12535 PMID: 24521437

44. van den Ackerveken GF, Dunn RM, Cozijnsen AJ, Vossen JP, van den Broek HW, de Wit PJ. Nitrogen limitation induces expression of the avirulence gene avr9 in the tomato pathogen Cladosporium fulvum. Mol. Gen. Genet. 1994; 243, 277–285. PMID: 8190081

45. van den Ackerveken GF, Dunn RM, Cozijnsen AJ, Vossen JP, van den Broek HW, de Wit PJ. Nitrogen limitation induces expression of the avirulence gene avr9 in the tomato pathogen Cladosporium fulvum. Mol. Gen. Genet. 1994; 243, 277–285. PMID: 8190081

46. Watanabe A, Ebizuka Y. Unprecedented mechanism of chain length determination in fungal aromatic polyketide synthases. Chem. Biol. 2004; 11:1101–6. https://doi.org/10.1016/j.chembio.2004.05.015 PMID: 15324811

47. Fan R, Klosterman SJ, Wang C, Subbarao KV, Xu X, Shang W, et al. Vayg1 is required for microsclerotium formation and melanin production in Verticillium dahliae. Fungal Genet. Biol. 1997; 98:1–11. https://doi.org/10.1016/j.fgb.2016.11.003 PMID: 27866941

48. Daub ME, Herrero S, Chung KR. Photoactivated perylenequinone toxins in fungal pathogenesis of plants. FEMS Microbiol. Lett. 2005; 252:197–206. https://doi.org/10.1016/j.femsle.2005.08.033 PMID: 16165316

49. Chen H, Lee MH, Daub ME, Chung KR. Molecular analysis of the cercosporin biosynthetic gene cluster in Cercospora nicotianae. Mol. Microbiol. 2007; 64:755–70. https://doi.org/10.1111/j.1365-2958.2007.05689.x PMID: 17462021

50. Newman AG, Townsend CA. Molecular characterization of the cercosporin biosynthetic pathway in the fungal plant pathogen Cercospora nicotianae. J. Am. Chem. Soc. 2016; 138:4219–28. https://doi.org/10.1021/jacs.6b00633 PMID: 26938470

51. So KK, Chung YJ, Kim JM, Kim BT, Park SM, Kim DH. Identification of a polyketide synthase gene in the synthesis of phleichromone in the phytopathogenic fungus Cladosporium phlei. Mol. Cells. 2015; 38:1105–10. https://doi.org/10.14348/molcells.2015.0208 PMID: 26612679

52. Bergmann S, Funk AN, Scherlach K, Schroeck V, Shelest E, Horn U, et al. Activation of a silent fungal polyketide biosynthesis pathway through regulatory cross talk with a cryptic nonribosomal peptide synthetase gene cluster. Appl. Environ. Microbiol. 2010; 76:8143–9. https://doi.org/10.1128/AEM.00683-10 PMID: 20952652

53. Upadhyay S, Xu X, Lowry D, Jackson JC, Roberson RW, Lin X. Subcellular compartmentalization and trafficking of the biosynthetic machinery for fungal melanin. Cell Rep. 2016; 14:2511–8. https://doi.org/10.1016/j.celrep.2016.02.059 PMID: 26972005

54. Eisenman HC, Nosanchuk JD, Webber JB, Emerson RJ, Camesano TA, Casadevall A. Microstructure of cell wall-associated melanin in the human pathogenic fungus Cryptococcus neoformans. Biochemistry. 2005; 44:3683–93. https://doi.org/10.1021/bi047731m PMID: 15751945

55. Riquelme M. Tip growth in filamentous fungi: a road trip to the apex. Annu. Rev. Microbiol. 2013; 67:587–609.

56. Pahirulzaman KAK, Williams K, Lazarus CM. A toolkit for heterologous expression of metabolic pathways in Aspergillus oryzae. Methods Enzymol. 2012; 517:241–260. https://doi.org/10.1016/B978-0-12-404634-4.00012-7 PMID: 23084942

57. Throckmorton K, Wiemann P, Keller NP. Evolution of chemical diversity in a group of non-reduced polyketide gene clusters: using phylogenetics to inform the search for novel fungal natural products. Toxins (Basel). 2015; 7:3572–607.

58. Edgar RC. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Research. 2004; 32, 1792–1797. https://doi.org/10.1093/nar/gkh340 PMID: 15034147

59. Castronova J. Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. Mol. Biol. Evol. 2000; 17: 540–552. https://doi.org/10.1093/oxfordjournals.molbev.a026334 PMID: 10742046

60. Guindon S, Gascuel O. A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst. Biol. 2003; 52:696–704. PMID: 14530136
61. Keane TM, Creevey CJ, Pentony MM, Naughton TJ, Mcinerney JO. Assessment of methods for amino acid matrix selection and their use on empirical data shows that ad hoc assumptions for choice of matrix are not justified. BMC Evol. Biol. 2006; 6:29. https://doi.org/10.1186/1471-2148-6-29 PMID: 16563161

62. Grigoriev IV, Nikitin R, Haridas S, Kuo A, Ohm R, Otillar R, et al. MycoCosm portal: gearing up for 1000 fungal genomes. Nucleic Acids Res. 2014; 42:D699–704. https://doi.org/10.1093/nar/gkt1183 PMID: 24297253

63. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J. Mol. Biol. 1990; 215:403–410. https://doi.org/10.1016/S0022-2836(05)80360-2 PMID: 2231712

64. Kumar S, Stecher G, Tamura K. MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. Mol. Biol. Evol. 2016; 33:1870–4. https://doi.org/10.1093/molbev/msw054 PMID: 27004904

65. Ökmen B, Etalo DW, Joosten MH, Bouwmeester HJ, de Vos RC, Collemare J, et al. Detoxification of α-tomatine by Cladosporium fulvum is required for full virulence on tomato. New Phytol. 2013; 198:1203–14. https://doi.org/10.1111/nph.12208 PMID: 23448507

66. Mehrabi R, Mirzadi Gohari A, da Silva GF, Steinberg G, Kema GH, de Wit PJ. Flexible gateway constructs for functional analyses of genes in plant pathogenic fungi. Fungal Genet. Biol. 2015; 79:186–92. https://doi.org/10.1016/j.fgb.2015.03.016 PMID: 26092806

67. Mesarich CH, Griffiths SA, van der Burgt A, Ökmen B, Beenen HG, Etalo DW, et al. Transcriptome sequencing uncovers the Avr5 avirulence gene of the tomato leaf mold pathogen Cladosporium fulvum. Mol. Plant Microbe Interact. 2014; 27:846–57. https://doi.org/10.1094/MPMI-02-14-0050-R PMID: 24678832

68. Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the 2(−ΔΔC(T)) method. Methods. 2001; 25, 402–408. https://doi.org/10.1006/meth.2001.1262 PMID: 11846609

69. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, et al. Fiji: an open-source platform for biological-image analysis. Nat. Methods. 2012; 9:676–82. https://doi.org/10.1038/nmeth.2019 PMID: 22743772