**Genome-Based Discovery of Polyketide-Derived Secondary Metabolism Pathways in the Barley Pathogen *Ramularia collo-cygni***

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*Ramularia collo-cygni* causes *Ramularia* leaf spot (RLS) disease of barley. The fungus develops asymptptomatically within its host until late in the growing season, when necrotic lesions become visible on upper leaves. Fungal secondary metabolites (SM) have been proposed as important factors in RLS lesion formation but the biosynthetic pathways involved remain largely unknown. Mining the *R. collo-cygni* genome revealed the presence of 10 polyketide synthases (PKS), 10 nonribosomal peptide synthetases (NRPS), and 3 hybrid PKS-NRPS (HPS) identified within clusters of genes with predicted functions associated with secondary metabolism. SM core genes along with their predicted transcriptional regulators exhibited transcriptional coexpression during infection of barley plants. Moreover, their expression peaked during early stages of host colonization and preceded or overlapped with the appearance of disease symptoms, suggesting that SM may manipulate the host to promote colonization or protect *R. collo-cygni* from competing organisms. Accordingly, *R. collo-cygni* inhibited the growth of several fungi in vitro, indicating that it synthesized and excreted antifungal agents. Taken together, these findings demonstrate that the *R. collo-cygni* genome contains the genetic architecture to synthesize a wide range of SM and suggests that coexpression of PKS and HPS is associated with competitive colonization of the host and early symptom development.

*Ramularia* leaf spot (RLS) is a late-season disease of barley (*Hordeum vulgare*) occurring in temperate regions worldwide. Typical RLS symptoms are often reddish-brown in color, delineated by the leaf vein, surrounded by a chlorotic halo, and visible on both sides of infected leaves (Huss 2002). Symptoms can also be observed on stems, heads, and awns in severely infected crops. As a result, RLS is considered a major disease of barley causing typical yield penalties of approximately 20% but these losses can be as high as 70% (Havis et al. 2015). RLS can also reduce grain size by up to 4%, which devalues the crop (Cromey et al. 2002; McGrann and Havis 2017).

RLS is caused by the filamentous fungus *Ramularia collo-cygni*, a Dothideomycete species within the order Capnodiales. *R. collo-cygni* was recently confirmed to be closely related to *Zymoseptoria tritici*, the fungus responsible for Septoria leaf blotch of wheat (Crous et al. 2001; McGrann et al. 2016). *R. collo-cygni* colonization initially occurs asymptometrically before turning to necrotrophic growth (Kaczmarek et al. 2017). The transition from asymptomatic to necrotrophic development appears linked with changes in environmental conditions, leading to breakdown of the plant antioxidant system as RLS symptoms generally develop concomitantly with leaf senescence (Schützendübel et al. 2008). For example, plants grown under high light conditions display higher disease levels than plants grown under normal light levels (Makepeace et al. 2008; McGrann and Brown 2018; Peraldi et al. 2014). Moreover, barley plants with altered redox homeostasis or delayed senescence also appear to show differential expression of RLS symptoms (McGrann et al. 2014, 2015a,b). The long latent period combined with the importance of seed transmission in *R. collo-cygni* dissemination (Havis et al. 2013) and the effects of abiotic stress on disease symptom formation has led to *R. collo-cygni* being described as an endophyte which can cause disease if it enters its necrotrophic stage (McGrann and Brown 2018; McGrann and Havis 2017). The exact causes of the transition from endophytic to necrotrophic lifestyle are currently unknown but the action of phytotoxic secondary metabolites (SM) has been proposed as a plausible mechanism (Heiser et al. 2003; Walters et al. 2008).

Many Dothideomycete fungi synthesize toxic SM during disease development (Stergiopoulos et al. 2013). SM such as the host-specific toxin victorin, produced by the Victoria blight pathogen of oat *Bipolaris victoriae* (syn. *Cochliobolus victoriae*), are required for pathogenicity (Lorang et al. 2004; Navarre and Wolpert 1999), whereas others such as the nonhost-specific phytotoxin dothistromin produced by *Dothistroma septosporum*, responsible for Dothistroma needle blight of pine trees, act as virulence factors mediating disease severity (Kabir et al. 2015). *R. collo-cygni* is known to produce light-activated phytotoxic SM called rubellins that have been isolated from *R. collo-cygni*-infected barley leaves (Miethbauer et al. 2003) and in vitro cultures (Heiser et al. 2004; Miethbauer et al. 2003).
Six rubellin derivatives, named alphabetically from A to E plus 14-dehydro rubellin D, have been found in *R. collo-cygni* cultures (Miethbauer et al. 2008). Rubellins are nonhost-specific toxins that induce fatty acid peroxidation in vitro upon light activation (Heiser et al. 2004). Because light influences RLS symptom expression (Makepeace et al. 2008; Peraldi et al. 2014), the link between RLS and the action of rubellins in planta was presumed.

Most SM produced by plant-pathogenic fungi are derived from four core biosynthesis pathways that are controlled by polyketide synthases (PKS), nonribosomal peptide synthetases (NRPS), terpene cyclases (TC; synonym: terpene synthases), and dimethylallyl tryptophan synthases (DMATS) (Muriain-Gonzalez et al. 2015). Rubellin is an anthraquinone-derived toxin structurally related to characterized fungal SM such as cladofulvin, emodin, and aflatoxin, produced by *Cladosporium fulvum*, *Aspergillus nidulans*, and *A. flavus*, respectively. Based on the structural similarity between rubellin and other anthraquinone SM, rubellin biosynthesis has been proposed to occur through a polyketide pathway (Miethbauer et al. 2008). Considering that SM often mediate the interactions between a pathogen and its host, understanding the potential of *R. collo-cygni* for producing such compounds may provide insights into the nature of these interactions. Here, we investigated secondary metabolism in *R. collo-cygni* using the recently sequenced genome of this fungus (McGrann et al. 2016). We identified gene clusters involved in the synthesis of polyketide-derived SM and demonstrated their transcriptional coregulation during *R. collo-cygni* colonization of the barley host and early stages of RLS symptom development.

**RESULTS**

**Identification of secondary metabolism-related core genes in *R. collo-cygni***

Using Basic Local Alignment Search Tool (BLAST) combined with domain analysis, the *R. collo-cygni* genome was found to contain 21 genes encoding for proteins containing domains associated with PKS and 25 genes encoding proteins exhibiting a domain associated with NRPS. However, only 10 putative PKS possessed the three domains ketosynthase (KS), acyl-transferase (AT), and acyl carrier protein (ACP) required for prototypical PKS functionality. Of the ten putatively functional PKS, three exhibited the domain organization of non-reducing (NR)-PKS and seven had the organization of highly and partially reducing PKS (Fig. 1). A further three genes were classified as hybrid PKS-NRPS (HPS) genes because they encoded proteins containing a typical highly reducing PKS module fused to a single NRPS module consisting of condensation, adenylation, and peptidyl carrier protein domains (Fig. 1). The best BLAST hit for each *R. collo-cygni* PKS and HPS is given in Table 1. Furthermore, 10 NRPS appeared to have the domain organization found in functional type A and type B NRPS proteins. Three NRPS proteins were monomodular,

![Fig. 1. Domain organization of the proteins encoded by *Ramularia collo-cygni* polyketide synthase and hybrid polyketide synthase/nonribosomal peptide synthetase (PKS-NRPS) genes. PKS and hybrid PKS-NRPS domains are labeled from N-terminus to C-terminus: starter unit acyl carrier transacylase (SAT), ketosynthase (KS), acyl transferase (AT), product template (PT), dehydratase (DH), methyl transferase (MT), enoyl reductase (ER), acyl carrier protein (ACP), condensation (C), adenylation (A), peptidyl carrier protein (PCP), thiolesterase (TE), and reductase (R). Only PKS-NRPS genes with domain structures found in functional secondary metabolite (SM) core proteins are shown.](image-url)
two were bimodular, and five were multimodular (Supplementary Fig. S1). Corresponding GenBank accession numbers for the *R. collo-cygni* SM core genes are presented in Supplementary Table S1. Compared with a previous analysis that identified only four TC and no DMATS in the *R. collo-cygni* genome (McGrann et al. 2016), it appears that *R. collo-cygni* possesses at least 27 SM core genes. A similar number of SM core genes was obtained using the alternative detection methods Antibiotics and Secondary Metabolite Analysis Shell (AntiSMASH) and Secondary Metabolite Unique Regions Finder (SMURF) which identified 26 and 18 SM core genes, respectively (Supplementary Table S2). The number of core genes identified in *R. collo-cygni* is similar to that found in other closely related species of the class Dothideomycete such as *Z. tritici*, *C. fulvum*, and *Pseudocercospora fijiensis* (de Wit et al. 2012; McGrann et al. 2016; Ohm et al. 2012; Noar and Daub 2016). Thus, the *R. collo-cygni* genome encodes for an appropriate number of SM core genes but these do not appear to reflect completely the lifestyle of the fungus because no correlation was apparent between fungi with different lifestyles in the order Capnodiales (Supplementary Table S3).

**Prediction of *R. collo-cygni* PKS and HPS secondary metabolism gene clusters.**

The genomic locus around each putatively functional PKS and HPS gene was investigated by in silico genome walking. Eight of the identified PKS and all three HPS core genes appeared to be located in a cluster containing several genes with a predicted function associated with secondary metabolism (Fig. 2). Pks8 and Pks2 were not located within potential SM gene clusters but these PKS sequences were located at an end of their respective contig or scaffold; thus, it is possible that SM-related genes are present near these core genes but these regions were not completely assembled in the draft genome assembly. The largest cluster, located on contig 58 near Pks6, was over 45.2 kb in length and contained 16 putative SM-related genes and two predicted genes which currently have no transcript evidence based on RNAseq data (McGrann et al. 2016). The smallest cluster spanned 17 kb on contig 282 and contained five SM-related genes, including the PKS core gene (Pks10) and a TC gene (Fig. 2, gene Tc1) previously identified in the *R. collo-cygni* genome (McGrann et al. 2016). Of the eight putative SM clusters located around PKS, four appeared to have more than one predicted core gene present within the cluster. The cluster on contig 118 contained two genes with predicted PKS domains; however, only Pks3 exhibited the typical domain organization of functional PKS. Two other clusters, located on contig 280 and contig 58 near Pks9 and Pks6, both contained an additional SM gene exhibiting a domain architecture found in nonfunctional NRPS. These genes were designated Nps12 (contig 280) and Nps13 (contig 58). Of the 11 clusters surrounding PKS or HPS core genes, six possessed at least one gene with a predicted function associated with cellular transport from either the major facilitator superfamily (MFS) or ATP-binding cassette family, which are known to be involved in secondary metabolism processes, including protecting fungi against their own toxic SM (Gardiner et al. 2005). Two putative SM clusters surrounding a PKS core gene (Pks1 and Pks6, identified on scaffold m24 and contig 58, respectively) and two located near a HPS gene (Hps1 and Hps2, identified on contig 1 and contig 14, respectively) contained predicted transcription factors with GAL4 domains, also known as Zn(II)$_2$Cys$_6$ DNA-binding domains, that have been shown to regulate secondary metabolism in fungal pathogens (Brakhage 2013).

Three of the SM-related gene clusters found in *R. collo-cygni* exhibited high similarity to clusters involved in the biosynthesis of SM previously characterized in another fungal species (Fig. 3; Supplementary Table S4). Four of the six genes identified in the cluster located on contig 30 containing Pks5 are similar to genes present in the betaenone biosynthesis cluster identified in the oilseed rape pathogen *Phoma betae* (syn. *Pleospora betae*) (Ugai et al. 2015) (Fig. 3A). Pks1 was located in a cluster exhibiting similarity to an SM-cluster involved in monodictyphenone production in *A. nidulans* (Chiang et al. 2010) (Fig. 3B). Similarly, the locus on contig 14 surrounding Hps2 comprises genes showing similarity to all seven genes present in the chaetoglobosin A biosynthetic cluster identified in *Penicillium expansum* as well as two genes not identified in the *P. expansum* cluster (Schimmann and Hertweck 2007) (Fig. 3C). The clusters identified on contigs 282, 118, 58, 1, and 38, surrounding Pks10, Pks3, Pks6, Hps1, and Hps3, respectively, all showed similarity to SM clusters present in other fungal species but the end SM products of these clusters are currently unknown (Supplementary Fig. S2). Because all of the *R. collo-cygni* clusters that showed similarity to SM gene clusters in other fungi typically exhibited gene inversions and rearrangements within the *R. collo-cygni* locus, it remains to be determined whether or not *R. collo-cygni* can synthesize these SM products.

The relationship between *R. collo-cygni* core genes and core genes in other fungal species was further examined by a
Fig. 2. Organization of the genetic loci around *Ramularia collo-cygni* secondary metabolism core genes. Arrowheads indicate predicted direction of transcription for each open reading frame. The end of a cluster is marked by a double oblique bar and the end of a contig or scaffold is marked by a single vertical bar. Brackets indicate the size of the cluster. Each cluster is drawn to scale. Secondary metabolite core genes with domain structures that would indicate the resultant protein is nonfunctional are marked with an asterisk.
peaking at 7 dpi and slowly decreasing to minimal levels at 21 dpi, whereas Hps2 was highly expressed at both 5 and 7 dpi before declining to a low level of expression from 10 dpi onward (Fig. 5). The expression profile of several NRPS indicated that these SM core genes were also most abundant at 5 and 7 dpi (Supplementary Fig. S5). SM core gene transcription was highest when foliar R. collo-cygni biomass was at its lowest. Concomitant with the expression of RLS symptoms, there is a decline in the host’s antioxidant system (Schützendübel et al. 2008) which is associated with elevated levels of reactive oxygen species hydrogen peroxide in the plant (McGrann and Brown 2018). Therefore, R. collo-cygni catalase expression was measured over the time course of disease development as an experimental control gene that would be predicted to increase transcript abundance as disease developed, similar to the interaction between the related fungus Z. tritici and wheat (Keon et al. 2005). As R. collo-cygni biomass increased over the experiment time course along with RLS symptom expression, transcript levels of R. collo-cygni catalase, a hydrogen peroxide scavenger, also increased (Fig. 5). These results suggest that elevated R. collo-cygni SM core gene expression was distinct from fungal biomass accumulation and preceded by or overlapped with early disease symptom expression.

Because the expression profile of most of the R. collo-cygni SM core genes investigated here showed similar patterns of transcription, the expression of previously characterized fungal transcriptional regulators as well as SM pathway-specific transcription factors (TF) was investigated to determine possible coregulation of these genes. Putative R. collo-cygni homologs of the zinc-finger TF PacC, CreA, and AreA, involved in responses to pH, carbon, and nitrogen conditions, respectively; VeA and LaeA, two components of the light-activated velvet complex which regulates diverse biological processes, including asexual reproduction and secondary metabolism; and pathway-specific TF AflR and AflJ, both known to regulate genes involved in the aflatoxin and dothistromin biosynthetic cluster in A. flavus and D. septosporum, were identified in the R. collo-cygni genome using BLASTp (Supplementary Table S5). Transcript profiles of PacC and AreA were similar to those observed for most of the SM core genes, with highest transcript levels at 5 and 7 dpi before a general decline in gene expression from 10 dpi onward (Fig. 6A). The pathway-specific TF AflR and AflJ were also most highly expressed during the early stages of R. collo-cygni colonization (Fig. 6C). Both VeA and LaeA exhibited slightly different expression profiles, with the highest transcription of these genes between 5 and 10 dpi (Fig. 6B). These results indicate that several transcription regulators with known roles in SM regulation in plant-pathogenic fungi appear coregulated with most of the R. collo-cygni SM core genes examined. The increased expression of SM core genes during the early stages of R. collo-cygni colonization suggests that, if these SM are indeed produced by R. collo-cygni, the SM products of the genes investigated here could play a role in early plant–pathogen interactions such as manipulating the host to aid fungal colonization or to protect the fungus from competing organisms.

**In vitro antifungal activity of R. collo-cygni.**

To assess antifungal activity of SM produced by R. collo-cygni, in vitro dual-culture competition assays between R. collo-cygni and common fungal pathogens of barley were used. The presence of R. collo-cygni significantly inhibited the growth of all of the tested competing fungi examined (Fig. 7). Growth of Pyrenophora teres, the causative agent of net blotch, was inhibited by 66% (P = 0.041) and 80% (P = 0.002) on alkyl ester agar (AEA) and potato dextrose agar (PDA) media.

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**Fig. 3.** Synteny between selected *Ramularia collo-cygni* secondary metabolism clusters and gene clusters in other fungi. Syntenic relationship between predicted *R. collo-cygni* secondary metabolism clusters and gene clusters from A, *Phoma betae*; B, *Aspergillus nidulans*; and C, *Penicillium expansum* are shown. Arrowheads indicate open reading frame predicted transcription direction. Putative orthologous genes identified by reciprocal BLAST are marked by dotted lines. The end of a cluster is marked by a double oblique bar and the end of a contig or scaffold is marked by a single vertical bar.
respectively. *Fusarium poae* growth was inhibited on both media by more than 50% in the presence of *R. collo-cygni* (AEA, *P* < 0.001; PDA, *P* < 0.001) and that of *Fusarium graminearum* by more than 60% (AEA, *P* < 0.001; PDA, *P* < 0.001). *R. collo-cygni* also had an antagonistic effect on *Rhynchosporium commune*, responsible for barley leaf scald, growth (*P* < 0.001) on PDA. A red coloration of the medium surrounding *R. collo-cygni* cultures that was particularly visible on PDA when grown in competition with *P. teres* and *Rhynchosporium commune* (Fig. 7B) suggests that *R. collo-cygni* releases compounds into the growing medium that inhibits the growth of other fungi. Previous reports have demonstrated that the one characterized SM family synthesized by *R. collo-cygni*, the rubellins, can produce a red coloration in culture media (Heiser et al. 2003). Together, these data illustrate that SM could provide *R. collo-cygni* with a competitive advantage over other microorganisms during host colonization.

**DISCUSSION**

Genomic analysis of the *R. collo-cygni* genome reveals the potential to produce several different SM.

Genomes of filamentous species of Ascomycetes and, in particular, of Dothideomycetes possess a large number of SM-related genes which result in the ability to produce a wide range of chemical compounds (Bushley and Turgeon 2010; Kroken et al. 2003). The *R. collo-cygni* genome contains at least 23 potentially functional SM core genes belonging to the PKS, NRPS, and HPS families. This finding contrasts with results of McGrann et al. (2016), which reported 19 PKS, 14 NRPS, and 4 TC but did not report any dimethylallyl tryptophan synthases or HPS. This variation highlights the importance of analyzing domain structure rather than relying on BLAST similarity alone to identify SM core genes, which can lead to misidentification of putatively nonfunctional genes as well as excluding HPS. Core genes tend to be organized within the *R. collo-cygni* genome in SM-related gene clusters. Clusters exhibiting similarity with previously reported fungal SM clusters involved in the biosynthesis of characterized SM were identified in the *R. collo-cygni* genome (Fig. 3). The cluster located at the *Pks5* loci contained putative homologs to all the genes involved in betaenone biosynthesis in *Phoma betae* (Ugai et al. 2015). Betaenones are phytotoxins associated with leaf spot disease of sugar beet and induce leaf chlorosis (Ichihara et al. 1983). The cluster located near the core gene *Hps2* contained putative homologs of the seven genes involved in the biosynthesis of chaetoglobosin A in *Penicillium expansum* and two additional genes, a putative O-methyltransferase and an MFS transporter.

![Fig. 4. Development of Ramularia leaf spot (RLS) in barley Century seedlings. A, Representative images of RLS symptoms on barley seedlings over a 21-day time course following inoculation with *Ramularia collo-cygni* DK05 Rcc001 ss2; dpi = days postinoculation B, Decline in green leaf area (GLA) retention (black line) and increase in RLS symptom development (gray bars) measured over the 21-day inoculation time course as the area under the disease progress curve (AUDPC). AUDPC is presented as a proportion of the maximum AUDPC possible over the entire inoculation course. Time points that showed significantly different development of RLS symptoms (top row) or differences in GLA retention (bottom row) from each other (*P* < 0.05) are indicated by different letters. C, *R. collo-cygni* DNA levels (in picograms) in barley prophyl leaves measured by quantitative polymerase chain reaction. Fungal DNA levels (in picograms) were calculated using a standard curve produced from a dilution series of *R. collo-cygni* DNA. Time points that showed significantly different (*P* < 0.05) amounts of fungal DNA are indicated by different letters.](Vol. 31, No. 9, 2018 / 967)
Chaetoglobosins are known to have cytotoxic properties (Ohtsubo et al. 1978) and to exhibit antifungal activity (Zhang et al. 2013). Therefore, although, to date, only rubellins were isolated from \textit{R. collo-cygni}, it appears that \textit{R. collo-cygni} has the genetic capacity to produce a wide range of SM, including several phytotoxins and antifungal agents. \textit{R. collo-cygni} gene clusters with similarity to uncharacterized SM biosynthetic clusters, where the function of the SM has not yet been determined in other species, were also discovered, further highlighting the potential diversity of the \textit{R. collo-cygni} SM arsenal.

The pathway responsible for rubellin biosynthesis has yet to be determined but the chemical structure of this SM may

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\textbf{Fig. 5.} Expression of selected \textit{Ramularia collo-cygni} polyketide synthases (PKS) and hybrid polyketide synthase/nonribosomal peptide synthetase (HPS) genes during \textit{Ramularia} leaf spot development in barley seedlings. \textbf{A}, Pks1; \textbf{B}, Pks2; \textbf{C}, Pks3; \textbf{D}, Pks4; \textbf{E}, Pks5; \textbf{F}, Pks10; \textbf{G}, Hps1; \textbf{H}, Hps2; \textbf{I}, Hps3; and \textbf{J}, Catalase. Data are presented as mean normalized expression values ± standard error over a 21-day time course postinoculation. The expression of the \textit{R. collo-cygni} catalase gene is shown as a control.
provide clues toward better understanding its production. Chemically, rubellins resemble several fungal polyketide-derived SM such as endocrocin, emodin, and cladofulvin that all exhibit a dioxoanthracene backbone characteristic of anthraquinones and are synthesized by the NR-PKS EncA, MdpG, and Clag, in A. fumigatus, A. nidulans, and C. fulvum, respectively (Chiang et al. 2010; Griffiths et al. 2016; Lim et al. 2012). Three NR-PKS (Pks3, Pks1, and Pks2) were identified in the R. collo-cygni genome. Of these R. collo-cygni NR-PKS, Pks3 can probably be excluded from the biosynthesis of rubellin or other anthraquinone-derived metabolites because the domain organization of Pks3, which includes a methyltransferase domain and lacks a product template domain, differs significantly from the SAT-KS-AT-PT-ACP organization of EncA, MdpG, and Clag. Furthermore, the phylogenetic analysis suggests that the protein encoded by Pks3 is not related to proteins involved in anthraquinone biosynthesis. The domain organization of Pks2 is similar to that of other NR-PKS, including AjPksA and DsPksA, involved in the production of the anthraquinone-derived A. flavus aflatoxin and D. septosporum dothistromin, as well as the PKS encoded by CTB1, Pks1, and Cppks1 involved in the biosynthesis of the perylenequinone toxins cercosporin, elsinochrome, and phleichrome, isolated from Cercospora nicotianae, Elsinoë fawcettii, and Cladosporium phlei, respectively (Choquer et al. 2005; Liao and Chung 2008; So et al. 2015). Of these NR-PKS, the phylogenetic

![Graphs showing expression of putative Ramularia collo-cygni homologs of characterized fungal secondary metabolism regulators during Ramularia leaf spot development in barley seedlings.](image-url)
analysis suggests that \textit{Pks2} is more closely related to the elsinochrome synthase \textit{Pks1} and the phleichrome synthase \textit{CpPks1} than \textit{AfPksA} and \textit{DsPksA}, which are recovered in a different clade. These results suggest that \textit{Pks2} may be involved in either the production of an anthraquinone-derived SM or the biosynthesis of SM potentially belonging to the perylenequinone family.

\textit{Pks1} encodes a protein exhibiting the same domain organization as the proteins encoded by \textit{MdpG}, \textit{EncA}, and \textit{ClaG}. \textit{R. collo-cygni} PKS \textit{Pks1} was also closely related to \textit{MdpG}, \textit{EncA}, and \textit{ClaG}, as well as \textit{Acas}, which is responsible for the biosynthesis of atrochrysone carboxylic acid, a precursor of endocrocin, emodin, chrysophanol, and cladofulvin (Awakawa et al. 2009; Griffiths et al. 2016). A putative role for \textit{Pks1} in emodin or chrysophanol biosynthesis was further supported by the colocalization of \textit{Pks1} with genes exhibiting sequence similarity to those identified in the monodictyphenone biosynthetic cluster involved in emodin and chrysophanol production (Fig. 3B) (Chiang et al. 2010). Considering that the rubellins reported from \textit{R. collo-cygni}, especially rubellin A

![Growth inhibition assays between \textit{Ramularia collo-cygni} and common foliar fungal pathogens of barley.](image)
and B, and uredinorubellins (Miethbauer et al. 2008) exhibit a chemical structure resembling that of a dimer of chrysophanol, Pks1 appears to be a strong PKS candidate involved in the rubellin biosynthetic pathway. Functional characterization of this gene and its genetic locus may provide further insights into the pathways that regulate rubellin biosynthesis.

**R. collo-cygni secondary metabolism core gene transcripts are most abundant during the early stages of R. collo-cygni colonization.**

Transcript levels of core genes associated with potential secondary metabolism pathways in *R. collo-cygni* were most abundant during the asymptomatic and the earliest lesion formation phases following *R. collo-cygni* colonization (Fig. 5). Transcription of SM-related genes in *Colletotrichum orbiculare*, *C. higginsianum*, and *C. gloeosporioides* were most abundant during the early stages of infection (Alkan et al. 2015; Gan et al. 2013; O’Connell et al. 2012). In the *R. collo-cygni* sister species *Z. tritici*, SM gene transcripts were most abundant during the transition phase from biotrophic to necrotrophic growth (Rudd et al. 2015). SM produced during this transition phase could be involved in protecting the fungus from plant defense mechanisms or inducing host cell death to facilitate fungal development during necrotrophic colonization. However, some secondary metabolism-associated transcripts were found to be most abundant during later stages of disease development in *Z. tritici*, including sporulation (Cairns and Meyer 2017; Palma-Guerrero et al. 2016). Similarly, in the closely related *D. septosporum*, four core genes, including three PKS, were upregulated during the later stages of infection (Bradshaw et al. 2016). The variation in SM transcription profiles between different species of Dothideomycetes highlights the complex nature of SM functions in these fungi and may provide insights into how SM affects the unique pathology of the associated diseases.

Regulation of fungal SM-related genes is the result of a finely tuned balance between requirements for fungal development and secondary metabolism (Calvo et al. 2002). As a result, SM-related genes are often coregulated, facilitating rapid control of SM production. *R. collo-cygni* SM core gene expression appeared to be coregulated with several well-characterized fungal transcriptional regulators, including the pH-dependent transcription regulator *PacC* as well as components of the light-regulated velvet complex. However, the similar expression profile of *R. collo-cygni* *PacC* (Fig. 6A) with SM gene transcription in *R. collo-cygni* does not conclusively indicate that this gene is involved in regulating SM gene expression in *R. collo-cygni*. The observation that *PacC*-specific 5'-GCCARG-3' binding sites (Tilburn et al. 1995) were present within 1,000 bp of the start codon of all the tested *R. collo-cygni* PKS genes (data not shown), except *Pks3*, which has a slightly different expression profile, provides further evidence that may indicate that *PacC* could be involved in PKS core gene regulation in *R. collo-cygni*. Further studies on the possible role of *PacC* and other well-characterized fungal SM gene regulators are required to ascertain the true role of such genes in SM gene transcription in *R. collo-cygni*.

The velvet complex differentially regulates fungal SM biosynthesis in a light-dependent manner (Wiemann et al. 2010). This complex, which is formed by the three proteins VelB, VeA, and LaeA, is involved in regulating SM-related gene expression (Bayram et al. 2008). Most of the *R. collo-cygni* core genes examined showed higher transcript levels at the earliest stages of fungal colonization concomitant with higher levels of VeA and LaeA transcription (Fig. 6B). Similar results were found in other fungal species such as *A. nidulans*, *F. fujikuroi*, *F. oxysporum*, and *B. maydis*, where the velvet complex regulates SM biosynthesis (López-Berges et al. 2013; Park et al. 2012; Wiemann et al. 2010; Wu et al. 2012). These data indicate that the velvet complex may be involved in SM gene expression regulation in *R. collo-cygni*; however, the elevated transcription of VeA and LaeA at 10 dpi, when transcription of most *R. collo-cygni* SM core genes was declining, suggests that either control of secondary metabolism in *R. collo-cygni* may be under the control of other transcriptional regulators or the velvet complex may have another function in *R. collo-cygni*. Obtaining *R. collo-cygni* mutants impaired in *LaeA* or VeA expression may provide useful insights into the role played by the velvet complex in this fungus.

The two NR-PKS *Pks2* and *Pks1*, with the highest sequence similarity to *D. septosporum* *PksA*, the core gene involved in dothistromin biosynthesis, and *Pks5*, which was identified in a betaenone-like cluster (Fig. 3), were most abundant at 5 and 7 dpi, suggesting that, if the SM synthesized by these PKS are phytotoxins, then these SM could be produced early during *R. collo-cygni* colonization and stored. A red compound was observed inside vesicles within *R. collo-cygni* hyphae, indicating that metabolites such as rubellin could be stored within the fungus until needed (McGrann et al. 2016). However, the presence of SM in vesicles does not necessarily correspond to a storage phase. In *A. parasticus*, several steps in the biosynthesis of the highly toxic SM aflatoxin are carried out in specialized vesicles termed aflatoxisomes (Chanda et al. 2009). Therefore, the previous observations of red-colored compounds stored in vesicles within *R. collo-cygni* hyphae may correspond to a specific phase in *R. collo-cygni* SM biosynthesis rather than a storage phase. Alternatively, the necrotic lesions observed as early as 7 dpi (Fig. 4) during RLS development indicate the action of a cell-death inducer. It is possible that it is these early-colonization phase cell-death inducers that are being observed in hyphal vesicles although, because cell death can be induced through a number of pathways, it is unclear whether the observed cell death is linked with the action of a phytotoxin, SM, or some other product.

Transcript levels of the three HPS identified were also the highest during the early stages of colonization. If the chaetoglobosin A-like cluster located at the *Hps2* locus is functional, the expression profile of *Hps2* suggests that this HPS may synthesize an antifungal agent early during the infection of the host plant. Considering the slow-growing nature of *R. collo-cygni* (Kaczmarek et al. 2017), which initially develops as an endophyte, chaetoglobosin production could act to eliminate fungi competing for the same ecological niche as *R. collo-cygni*. In vitro antifungal activity of *R. collo-cygni* was observed against common fungal pathogens of barley (Fig. 7). The antimicrobial activity of rubellin toxins produced by *R. collo-cygni* is reported by Miethbauer et al. (2008) as being light dependent. The assay used here was undertaken in constant darkness, suggesting that the antimicrobial effect of *R. collo-cygni* on in vitro fungal growth may not be linked with rubellin activity. Therefore, other SM such as chaetoglobosin or other non-SM compounds may be responsible for the observed antifungal activity and function to confer an ecological advantage to *R. collo-cygni*. Obtaining *R. collo-cygni* mutants in which SM core genes have been deleted may provide useful insights into the role played by SM in the biology of this fungus.

In silico analysis of the *R. collo-cygni* genome reveals that this fungus, like many other members of class Dothideomycetes, has the genetic potential to produce several classes of SM, including hybrid polyketide/nonribosomal peptides such as chaetoglobosin or polyketides such as betaenones or the previously characterized anthraquinone rubellin (Heiser et al. 2004; Miethbauer et al. 2003). Although the identification of
the complete \emph{R. collo-cygni} secondary metabolism core genes. The genome of \emph{R. collo-cygni} isolate DK05 Rcc001, collected from the susceptible spring barley cultivar Braemar in Denmark in 2005, was recently sequenced and estimated to be 30.3 Mb with 90x coverage and 11,617 predicted gene models (McGrann et al. 2016). RNAseq data were generated from total RNA extracted from mycelium of DK05 grown in vitro on PDA plates for 10 to 12 days and confirmed the expression of 8,514 of the predicted 11,617 genes under these conditions. A database containing the corresponding 11,617 \emph{R. collo-cygni} protein models was created in GeneiousR9 software (Biomatters Ltd., Auckland, New Zealand). Protein sequences of PKS and NRPS involved in the synthesis of model fungal SM were downloaded from the National Center for Biotechnology Information (NCBI) GenBank database (Supplementary Table S6). These sequences were used to interrogate the \emph{R. collo-cygni} genome using protein BLASTp (BLASTp) (Altschul et al. 1990) to identify putative homologs in \emph{R. collo-cygni}. Significant \emph{R. collo-cygni} matches were retained if the BLASTp search with at least one of the fungal query proteins had an \emph{e}-value < $1.e^{-10}$. The protein sequence of the corresponding \emph{R. collo-cygni} gene models was then used to interrogate InterProScan and the PKS-NRPS analysis website to identify typical PKS or NRPS domains. Results from these analyses were combined and proteins that did not contain at least one of the PKS or NRPS domains were eliminated. Genes encoding the selected proteins were positioned on contigs or scaffolds within the \emph{R. collo-cygni} genome assembly using tBLASTn. For comparison purposes, the SM core gene complement of the \emph{R. collo-cygni} genome was also analyzed using the programs AntiSMASH and SMURF.

**Identification of putative secondary metabolism clusters.**

To identify putative SM clusters, in silico genome walking was carried out both up- and downstream along \emph{R. collo-cygni} contigs or scaffolds where each core gene was located. StarOrf (Massachusetts Institute of Technology, Cambridge, MA, U.S.A.) was used to determine open reading frames (ORF) along each contig or scaffold, with a minimal ORF size of 120 bp. The protein sequences of each ORF predicted in the genetic space surrounding each putative core gene were used to interrogate the NCBI nonredundant protein database with BLASTp with an \emph{e}-value cut-off < $1.e^{-10}$. Each protein sequence of the best-match fungal BLAST hit was downloaded from GenBank and used to interrogate the \emph{R. collo-cygni} protein models by BLASTp and the annotation of each gene in every cluster was confirmed. Borders of a gene cluster were delineated by a 5-kb region of noncoding DNA or when three consecutive genes with no predicted function associated with SM production were identified. Comparative analysis of predicted \emph{R. collo-cygni} SM clusters with SM clusters in other fungal species was performed with the AntiSMASH website (Blin et al. 2017) and similarity between genes within clusters was confirmed for each gene by reciprocal BLASTp.

**Phylogenetic analysis of \emph{R. collo-cygni} PKS and HPS proteins.**

Protein sequences of PKS and HPS involved in the biosynthesis of characterized fungal SM used in previous phylogenetic studies (Collemare et al. 2014; Fisch 2013) as well as sequences from fungi related to \emph{R. collo-cygni} showing sequence similarity, based on BLASTp, to known SM core genes were downloaded from the GenBank database (Supplementary Table S7). Full-length sequences of PKS and HPS were aligned using MAFFT, and poorly aligned sequences were manually removed in Aliview (Larsson 2014). Phylogenetic trees were built using the RAxML, version 8.2.4 ( Stamatakis et al. 2005), with 100 rapid bootstrap replicates and the protein model PROT-GAMMAAUTO, where model testing is part of the algorithm.

**R. collo-cygni inoculation of barley seedlings.**

The RLS-susceptible spring barley cultivar Century was grown until growth stage 12/13 (Zadoks et al. 1974) in a controlled environment cabinet (Snijders Scientific, Tilburg, The Netherlands) at 18°C, with a photoperiod of 16 h of light and 8 h of darkness supplemented with light at 250 µmol m$^{-2}$ s$^{-1}$ and 80% relative humidity. \emph{R. collo-cygni} isolate Rcc001 DK05 ss2 was maintained and inoculum prepared as previously described (Makepepe et al. 2008; Peraldi et al. 2014). Inoculum was applied at a rate of 0.5 ml/seedling, as previously reported (Makepepe et al. 2008). Inoculated plants were placed in a plastic tray (50 by 40 by 5 cm), covered with a clear plastic lid to ensure maximum humidity, and incubated in the dark for 48 h at 18°C. Following incubation, the light regime was restored and, 72 h later, the clear plastic lids were removed. Development of RLS was assessed at 5, 7, 10, 12, 15, 18, and 21 dpi. Visual assessment was carried out on the prophyll leaf and disease severity was estimated as the percentage of the leaf area covered by RLS lesions. The area under the disease progress curve (Shaner and Finney 1977) was calculated to report the development of the disease over time. GLA retention was also visually assessed at each time point because RLS development has been associated with leaf senescence (Schützendübel et al. 2008). Data show the mean ± standard error calculated on six independent inoculation experiments, each of them containing two biological replications of 20 leaves each.

**Microscopic assessment of \emph{R. collo-cygni} colonization.**

Colonization by \emph{R. collo-cygni} during RLS development was followed using light microscopy (Leica DMRBE; Leica Microsystems, Wetlzar, Germany). Leaves were harvested at 5, 7, 10, 12, 15, 18, and 21 dpi and cleared in 70% ethanol for 7 days prior to staining with 0.05% trypan blue in lacto-glycerol and briefly destaining with 100% ethanol. For each leaf, a minimum of 120 stomata were observed over its full length. Fungal development stages and the associated host response was scored in three categories based on those described previously (McGrann and Brown 2018): (i) proportion of stomata penetrated by fungal hyphae, (ii) proportion of stomata surrounded by necrosis, and (iii) the proportions of stomata with fungal conidiophore-like structures emerging. Data show the mean ± standard error calculated on six independent experiments, each of them containing two biological replications of two leaves each.

**R. collo-cygni DNA quantification during RLS development.**

Two leaves were harvested from Century plants at each time point between 5 and 21 dpi and snap frozen in liquid nitrogen, and DNA was extracted using the DNeasy Plant Mini Kit (Qiagen Ltd., Hilden, Germany). \emph{R. collo-cygni} DNA
quantification based on an internal transcribed spacer region was carried out using qPCR as described by Taylor et al. (2010). Each reaction volume of 25 µl consisted of 1× iQ Supermix (Bio-Rad, Hercules, CA), forward primer RamF6 (5'-CGTACCTCCACCTCACAG-3') and reverse primer RamR6 (5'-CCCTCAGATAAGTTCGC-3') at 400 nmol liter⁻¹, 5 µl of DNA template (5 ng µl⁻¹), R. collo-cygni probe Ram6 (5'-GGACTCCCGTCGAGGTTGCGCATC-3') at 150 nmol liter⁻¹, and molecular-grade water to the final volume. Reactions were performed in a Mx3000P thermocycler (Agilent Technologies, Santa Clara, CA) using the following program: initial denaturation of 10 min at 95°C; followed by 50 cycles of 95°C for 20 s, 55°C for 20 s, and 72°C for 20 s; followed by a final extension of 95°C for 1 min. Potential primer-dimer formation and confirmation of single gene-specific product amplification was monitored at the end of each reaction by melt curve analysis. Data show the mean quantity (in picograms) of R. collo-cygni DNA calculated from a standard curve produced by a 10-fold dilution series of R. collo-cygni DNA. R. collo-cygni DNA levels were assessed in five independent inoculation experiments.

Expression profiling of R. collo-cygni SM-related genes.
Transcript levels of putative R. collo-cygni SM-related genes were assessed as RLS developed 5 to 21 dpi with Rec001 DK05 ss2. For each biological replicate, two leaves were harvested and pooled prior to snap freezing in liquid nitrogen. Total RNA was extracted using the RNeasy Mini Kit (Qiagen Ltd.) following the manufacturer’s instructions. Contaminating genomic DNA was removed from RNA samples using TURBO DNase I (Ambion, Thermo Fisher Scientific, Austin, TX). cDNA was synthesized from 1 µg of total RNA using the SuperScript III First-Strand Synthesis System (Invitrogen, Carlsbad, CA). cDNA was diluted 10-fold in sterile distilled water and used in qRT-PCR using the SybrGreen Jump Start Taq system (Sigma, Dorset, U.K.) and a Mx3000P thermocycler (Agilent Technologies). The stability of four R. collo-cygni reference genes—elongation factor 1-α, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), α-tubulin, and actin—was assessed under the conditions of this experiment using GeNorm (Vandesompele et al. 2002). The two most stable reference genes (GAPDH and α-tubulin) were used for cDNA normalization. Gene-specific primers (Supplementary Table S8) for R. collo-cygni genes of interest were designed using Primer3. Primers with amplification efficiency ranging from 80 to 110% were used for transcript quantification based on the E⁻ΔΔCt method (Livak and Schmittgen 2001). Data show the mean normalized expression values ± standard error of three independent experiments, each of them containing two biological replicates.

In vitro fungal growth inhibition assays.
Growth inhibition of commonly occurring barley pathogens F. poae, F. graminearum, Pyrenophora teres, and Rhynchosporium commune by Ramularia collo-cygni isolate Rec001 DK05 ss2 was assessed in dual-plate culture assays. The assay was carried out in 9-cm Petri dishes (Sarstedt, Numbrecht, Germany), on two different media, AEA and PDA (Sigma) containing streptomycin at 5 µg ml⁻¹. Growth inhibition of Rhynchosporium commune was only tested on PDA due to the slow growing nature of both Rhynchosporium commune and R. collo-cygni on AEA. To compensate for the differential in vitro growth rate between R. collo-cygni and F. poae, F. graminearum, and P. teres, an 8-mm plug of R. collo-cygni mycelium was extracted from a 14-day-old culture plate, placed 2 cm from the border of the Petri dish, and allowed to grow for 14 days in isolation. Mycelial plugs from competing fungal species were taken from 14-day-old cultures and placed 3 cm away from the R. collo-cygni culture. Considering the slow-growing nature of both R. collo-cygni and Rhynchosporium commune, the mycelial plugs of the two fungi were added to the culture plate at the same time and placed 3 cm apart. All fungal cultures were grown in a Sanyo MIR-254 incubator (Sanyo Electric Ltd., Osaka, Japan) in constant darkness at 15°C. Fungal radial growth was measured after 10 days for P. teres and F. poae, 7 days for F. graminearum, and 21 days for Rhynchosporium commune. The assay was repeated at least three times for each fungus, with experimental replicates consisting of at least three individual plates per assay.

Data analysis.
All statistical tests were performed in GenStat, v.16 (Payne et al. 2009). Data were analyzed by analysis of variance using generalized linear models (GLM) unless stated otherwise. Each GLM assessed how the various factors that contributed to the design of the different experiments affected variation in the scored phenotype. Disease score data from the R. collo-cygni time-course inoculation experiment were Logit+ transformed (McGrann et al. 2014) prior to analysis. GLA retention during RLS development was assessed with a linear mixed model of repeated measures using the uniform correlation/split plot in time covariance matrix (McGrann et al. 2015a). The factors day, experiment, and tray were fixed in the model whereas the leaf–day interaction term was a random factor. The microscopy experiment was analyzed using a GLM where each score category was analyzed separately as a proportion of the total number of stomata scored on each leaf using a binomial distribution as the link function. Growth inhibition effects of R. collo-cygni on each pathogen were assessed separately using a GLM assessing variation attributable to experimental replicates and the direction of fungal growth from the point of inoculation either toward the R. collo-cygni culture or toward the edge of the culture plate. Data are presented as growth inhibition calculated as the ratio (percentage) of radial fungal growth toward R. collo-cygni culture from point of inoculation/radial fungal growth toward the edge of the culture plate from point of inoculation. Growth on either PDA or AEA was assessed separately. After modeling, treatments were compared within each experiment using t test probabilities and applying Fisher’s least significant difference to unplanned comparisons.

LITERATURE CITED


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AUTHOR-RECOMMENDED INTERNET RESOURCES


InterProScan: https://www.ebi.ac.uk

Primers3: http://primer3.ut.ee

MAFFT: https://www.ebi.ac.uk