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Pathogen Genetic Control of Transcriptome Variation in the *Arabidopsis thaliana* – *Botrytis cinerea* Pathosystem

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ABSTRACT In plant–pathogen relations, disease symptoms arise from the interaction of the host and pathogen genomes. Host–pathogen functional gene interactions are well described, whereas little is known about how the pathogen genetic variation modulates both organisms’ transcriptomes. To model and generate hypotheses on a generalist pathogen control of gene expression regulation, we used the *Arabidopsis thaliana*–*Botrytis cinerea* pathosystem and the genetic diversity of a collection of 96 *B. cinerea* isolates. We performed expression-based genome-wide association (eGWA) for each of 23,947 measurable transcripts in *Arabidopsis* (host), and 9267 measurable transcripts in *B. cinerea* (pathogen). Unlike other eGWA studies, we detected a relative absence of locally acting expression quantitative trait loci (*cis*-eQTL), partly caused by structural variants and allelic heterogeneity hindering their identification. This study identified several distantly acting *trans*-eQTL linked to eQTL hotspots dispersed across *Botrytis* genome that altered only *Botrytis* transcripts, only *Arabidopsis* transcripts, or transcripts from both species. Gene membership in the *trans*-eQTL hotspots suggests links between gene expression regulation and both known and novel virulence mechanisms in this pathosystem. Genes annotated to these hotspots provide potential targets for blocking manipulation of the host response by this ubiquitous generalist necrotrophic pathogen.

KEYWORDS Host–pathogen interaction; pathosystem; *Arabidopsis*; *Botrytis cinerea*; RNA-sequencing; genome-wide association; dual transcriptome

Introduction

Infectious disease results from an interaction between host and pathogen driven by the genetics of both organisms. The mechanisms of plant–pathogen interactions are often divided into qualitative, in which a few genetic variants of large effect shape binary disease outcomes, or quantitative, in which a spectrum of outcomes arise from the interaction of polygenic variation in the host and pathogen. The past decades have witnessed the unraveling of the molecular basis of large-

effect loci on both the host and the pathogen sides that control qualitative interactions (Giraldo and Valent 2013; Marone *et al.* 2013; Meng and Zhang 2013; Cui *et al.* 2015; Lo Presti *et al.* 2015). In the qualitative model, alternative alleles at these genes create sweeping differences in the transcriptome and phenotypic responses to infection in the host and pathogen via differential recognition events surrounding their proteins. However, plant–pathogen interactions cover a full range of genetic architectures, from few genes of large effect to many genes of small effect (Poland *et al.* 2009; Kou and Wang 2010; Lannou 2012). In contrast to qualitative systems, quantitative plant–pathogen interactions exhibit a lack of virulence/resistance genes that explain large proportions of the variance in the disease outcome in the population (Poland *et al.* 2009; Kou and Wang 2010; St. Clair 2010; Roux *et al.* 2014). These interactions are highly polygenic, with genetic variation influencing diverse molecular

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mechanisms, extending beyond host–pathogen perception and large-effect arms-race loci (Glazebrook 2005; Nomura *et al.* 2005; Goss and Bergelson 2006; Rowe and Kliebenstein 2008; Barrett *et al.* 2009; Corwin *et al.* 2016a; Bartoli and Roux 2017; Wu *et al.* 2017; Atwell *et al.* 2018 *preprint*; Fordyce *et al.* 2018; Soltis *et al.* 2019). It is, however, unclear how these polygenic molecular interactions alter higher-order phenotypes such as virulence or the transcriptome of both species. There is conflicting evidence on the balance of such pathosystem. Some studies and traits indicate that genetic variation in the pathogen dominates the interaction (Bartha *et al.* 2017; Wang *et al.* 2018a). Other studies find a balanced contribution of plant and pathogen genetics (Corwin *et al.* 2016a; Soltis *et al.* 2019). Thus, there is a need to develop genomic approaches to understand how polygenic variations affect the genomic response of both organisms.

Polygenic variation in the pathogen should influence numerous genes that shift the pathogen's transcriptome and cause differential expression of various virulence mechanisms. This virulence mechanism variation will affect the host and alter the host's resistance-associated transcriptome. Thus, by measuring the transcriptome in both the pathogen and the host, it should be possible to map how genetic variation in the pathogen is conveyed through the pathogen's transcriptome, and concurrently, how the host's transcriptome responds. Recent work has shown that it is possible to measure the pathogen's transcriptome in planta in the *Arabidopsis thaliana*–*Pseudomonas syringae* pathosystem, leading to new hypotheses about virulence (Nobori *et al.* 2018). In the *A. thaliana*–*Botrytis cinerea* pathosystem, the genetic interactions are dominated by complex small-effect loci that display a high degree of interaction between the host and pathogen (Denby *et al.* 2004; Finkers *et al.* 2007; Finkers *et al.* 2008; Rowe and Kliebenstein 2008; Anuradha *et al.* 2011; Fu *et al.* 2017; Fordyce *et al.* 2018). In this pathosystem, a cotranscriptome study with simultaneous analysis of the host and pathogen's transcripts was recently done through single-sample RNA-sequencing (Zhang *et al.* 2017; Zhang *et al.* 2019). This cotranscriptome approach allowed the description of key virulence networks in the pathogen and resistance responses within the host (Zhang *et al.* 2017; Zhang *et al.* 2019). Further, this study revealed a single network of transcripts from pathogen and host species. However, these studies did not assess the genetic architecture behind these cotranscriptome interactions.

Genome-wide association mapping (GWA) that identifies expression quantitative trait loci (eQTL; SNPs correlated with variations in transcript expression) can reveal the genetic architecture behind these cotranscriptome interactions. Previous eQTL studies revealed that the SNPs that cause differential transcript accumulation can be parsed into *cis* or *trans* effects. Locally acting (*cis*) eQTL indicate regulatory variation within or near the expressed gene itself. *trans*-eQTL reveal SNPs that are acting at a distance to affect regulatory

processes influencing the expression of the transcript. A *trans*-eQTL that affects many transcripts is classified as a hotspot. Such *trans*-acting hotspot SNPs may influence regulatory processes that, in turn, influence numerous transcripts.

eQTL analysis has been utilized to study host–pathogen interactions, albeit with a focus either on host or pathogen. Frequently, these studies focus on the host's response, such as mapping how host loci control host gene expression over time, using either traditional QTL mapping or GWA analysis (Chen *et al.* 2010; Hsu and Smith 2012; Zou *et al.* 2012; Allen *et al.* 2016; Christie *et al.* 2017). Additional studies have begun to invert this scheme by looking at how genetic variation in the pathogen influences the host transcriptome to identify pathogen loci modulating host expression levels, and thus to identify candidate loci for interspecific signals (Saeij *et al.* 2007; Wu *et al.* 2015; Guo *et al.* 2017). These studies attest to the potential to identify pathogen loci that influence host gene expression. However, previous studies have thus far addressed pathogen populations with limited genetic variation, and thus identify the few polymorphic loci between strains with strongest effects on transcriptomic variation (Wu *et al.* 2015; Guo *et al.* 2017).

The genomes of both the host and the pathogen harbor extensive genetic diversity that has been successfully used for genetic mapping to identify loci controlling virulence in combination with transcriptomics and genomics (Denby *et al.* 2004; Rowe and Kliebenstein 2008; Dalmais *et al.* 2011; Schumacher *et al.* 2012; Zhang *et al.* 2017; Atwell *et al.* 2018 *preprint*; Soltis *et al.* 2019). Expanding the scope of these studies, we performed a cotranscriptome analysis using a single wild-type *A. thaliana* Columbia-0 (Col-0) host accession, and *B. cinerea* pathogen transcriptomes were measured using a diverse *B. cinerea* population (Zhang *et al.* 2017; Zhang *et al.* 2019). We conducted a GWA analysis of both host and pathogen transcriptomes to identify loci in the *B. cinerea* genome that may affect the transcriptomes of either or both organisms (Zhang *et al.* 2017; Zhang *et al.* 2019). The loci tagged by these SNPs have an explicit directionality of effect, as genetic causality must arise within the pathogen and then extend to the host. Our analysis found mostly small-effect polymorphisms dispersed throughout the *B. cinerea* genome, with several *trans*-eQTL hotspots. These hotspot loci are associated with specific host or pathogen transcript coexpression modules and variation in lesion size. There was no identifiable overlap in the hotspots that influenced the host's or the pathogen's transcriptome, suggesting a surprisingly independent basis of transcriptional regulation of host and pathogen by the *B. cinerea* genome. Most of the hotspot-loci-tagged genes have no previous association to plant–pathogen virulence interactions. This generates a set of *B. cinerea* loci that have regulatory potential in controlling the *A. thaliana* and *B. cinerea* interaction via modulation of gene expression to influence the lesions outcome.

Materials and Methods

Previously published transcriptomics

For this study, we utilized the exact previously published transcriptome data involving 1164 RNA-sequencing libraries available under the NCBI accession number SRP149815 (Zhang *et al.* 2017; Zhang *et al.* 2019). A brief recitation of the materials and methods used to generate the transcriptome data are provided below. This large number of libraries measure the interaction of three *A. thaliana* genotypes (Col-0, *coi1-1*, and *npr1-1*) with a collection of 96 *B. cinerea* genotypes that were isolated as single spores from natural infections of fruit and vegetable tissues collected in California and internationally (Zhang *et al.* 2017; Atwell *et al.* 2018 preprint; Zhang *et al.* 2019; Caseys *et al.* 2020 preprint). We focused the analysis for this eGWA study on the *A. thaliana* wild-type accession Col-0. The transcriptome data were generated using fourfold replication of the full-randomized complete block experimental design across two independent experiments for all interactions (*i.e.*, two fully independent randomized biological replicates in each of two experiments). Leaves from Col-0 were harvested 5 weeks after sowing, and individually inoculated in a detached leaf assay with spores of each of 96 *B. cinerea* isolates (Zhang *et al.* 2017; Zhang *et al.* 2019). Whole leaves were sampled at 16 hr post-inoculation, before visible lesion formation, and flash-frozen for RNA isolation while lesion area was measured at 72 hr postinoculation. RNA-sequencing libraries were prepared as described in Kumar *et al.* 2012; Zhang *et al.* 2017; and Zhang *et al.* 2019. Reads were aligned to both the *A. thaliana* TAIR10.25 and the *B. cinerea* B05.10 ASM83294v1 complementary DNA reference genomes, and gene counts were pooled, summed across gene models, and normalized (Langmead *et al.* 2009; Li *et al.* 2009; Van Kan *et al.* 2017).

GWA of gene expression profiles

As phenotype for the GWA, we used the *z*-scaled model-adjusted least square means of normalized gene counts obtained from a negative binomial generalized linear model for both the *A. thaliana* and *B. cinerea* transcriptomes (Zhang *et al.* 2017; Zhang *et al.* 2019). GWA was implemented using a univariate linear mixed model in a genome-wide efficient mixed model association (GEMMA; Zhou and Stephens 2012). GWA was performed using PLINK binary ped format files with a standardized relatedness matrix calculated in GEMMA. The relatedness matrix accounts for population structure among *B. cinerea* isolates. Individual SNP significances were extracted as *P*-values from the score test. We used haploid binary SNP calls with <10% missing values and the default 1% minor allele frequency (MAF) was increased to 20%. The use of MAF > 0.20 should help to limit the false positive error given the population size (Tabangin *et al.* 2009). The mapping was based on 96 isolates with a total of 237,878 SNPs mapped to the *B. cinerea* B05.10 genome (Atwell *et al.* 2018 preprint). We ran GEMMA once per

phenotype, across 9267 *B. cinerea* gene expression profiles and 23,947 *A. thaliana* gene expression profiles.

Significance threshold by GWA of permuted transcripts

Given the large number of traits used for GWA (33,214 traits), full permutation tests assessing the significance of all associations is unfeasible. To query for potential patterns of association that may exist randomly and fix global thresholds of likely nonrandom association of SNPs with transcript variation, we performed a comparative GWA analysis of randomized assignment of each transcriptional profile across the 96-isolate collection. This analysis included five independent permutations of 9267 randomized *B. cinerea* transcripts and 23,947 randomized *A. thaliana* transcripts. Using these permutations, we compiled the top 5% of random *P*-values from each gene across all the permutations and utilized the median of these values as conservative genome-wide permutation thresholds, $1.96e^{-5}$ for the *B. cinerea* transcriptome and $2.9e^{-5}$ for the *A. thaliana* transcriptome. In *B. cinerea*, this threshold identified an average of 4756 SNPs associated with the expression of 461 transcripts (range 3843–5584 SNPs and 410–499 transcripts) across the five random permutations. In *A. thaliana*, this threshold identified an average of 16,446 SNPs associated with the expression of 1201 transcripts (range 13,040–22,359 SNPs and 1129–1260 transcripts) across the five random permutations. Permutation approaches are often more effective than *a priori* *P*-value thresholds to determine significance thresholds of GWA studies with large number of phenotypes (Evans and Cardon 2006).

eQTL hotspot identification, significance, and annotation

For the hotspot analysis we utilized the top single SNP per transcript using either the *B. cinerea* or *A. thaliana* transcripts alone to query for hotspots. This provides 9267 SNP associations for the 9267 *B. cinerea* transcripts and 23,947 SNP associations for the 23,947 *A. thaliana* transcripts. An eQTL hotspot was defined as a SNP having the most significant effect (top SNP) on the abundance of multiple transcripts. For an initial approach to deciding upon significant hotspots in each data set, we randomized the position of these positions across the full 237,878 SNPs and repeated this 1000 times. This analysis gave a maximum hotspot across all permutations as three transcripts for *B. cinerea* and six transcripts for 23,947 *A. thaliana*. This permutation approach, however, does not account for potential effects of coexpression on hotspot detection. Thus, we also estimated thresholds by using the five random permutations GWA analysis to query and measure the number of transcripts randomly associated with each SNP. From each permutation, we identified the position of the top SNP for each *B. cinerea* or *A. thaliana* transcript and found the top hotspot in each permutation. These maximum eQTL hotspot sizes across all the permutations were 11 *B. cinerea* and 80 *A. thaliana* transcripts. Thus, we conservatively defined significant hotspots

as SNPs associated with >20 transcripts in *B. cinerea* and 150 transcripts in *A. thaliana*.

To document the putative functionality of eQTL hotspots, we annotated SNPs to genes using the identity of the nearest gene within a 2-kb window. This window was chosen because the average linkage disequilibrium decay in the *B. cinerea* genome is <1 kb (Atwell *et al.* 2018 preprint). This loss of allelic association over short distances allows tagging SNPs to particular genes with some confidence. While the majority of annotated genes had single SNP hotspots, three genes had two independent SNP hotspots. Two genes on chromosome 12 denoting *A. thaliana* gene expression hotspots appear closely linked, but are separated by ~80 kb on the *B. cinerea* genome.

Annotation of gene ontology and modules

We annotated functions to *B. cinerea* genes using the BotPortal resource (<http://dx.doi.org/10.15454/IHYJCX>) and looked for patterns indicating signal peptides for secretion using the SignalP-5.0 Server (<http://www.cbs.dtu.dk/services/SignalP-5.0/>). We looked for functional overrepresentation among the genes targeted by each *A. thaliana* eQTL hotspot using the PANTHER overrepresentation test implemented by plant gene ontology (GO) term enrichment from TAIR (Lamesch *et al.* 2011; Cadic *et al.* 2013). Over-enrichment of genes found in the previous *B. cinerea* and *A. thaliana* transcriptome modules and the eQTL hotspots were tested by the hypergeometric test (Subramanian *et al.* 2005; Zhang *et al.* 2017; Zhang *et al.* 2019).

Biosynthetic-pathway-focused cis-eQTL analysis

All of the SNPs within the three clustered biosynthetic pathways were described previously (Zhang *et al.* 2019). Within each pathway, the SNPs were used to cluster the isolates by hierarchical clustering using the R package pvclust. Distances were estimated using the unweighted pair group method with arithmetic mean distance, with correlation distance and 1000 bootstrap replications (Suzuki and Shimodaira 2015). The 95% bootstrap probability values are shown in green, while the approximately unbiased *P*-values are reported in red. Clustering is drawn according to those edges with strong support under both estimations.

Data availability statement

Strains are available upon request. Supplemental Material, File S1 contains information on functional annotation of the *B. cinerea* genes targeted by the *B. cinerea* trans-eQTL hotspots. File S2 contains information on functional annotation of the *A. thaliana* genes targeted by the *B. cinerea* trans-eQTL hotspots. File S3 contains GO summary analysis of the *A. thaliana* genes targeted within each *B. cinerea* trans-eQTL hotspot. File S4 contains summary information on the top SNP hits from GWA of each *B. cinerea* expression trait (transcript). SNP data and R scripts are available in Dryad (<https://doi.org/10.25338/B83P56>). Transcriptome data involving 1164 RNA-sequencing libraries are available under

the NCBI accession number SRP149815 (Zhang *et al.* 2017; Zhang *et al.* 2019). Gene annotations for *B. cinerea* genes are available on the BotPortal online platform (<http://dx.doi.org/10.15454/IHYJCX>). Supplemental material available at figshare: <https://doi.org/10.25386/genetics.8069480>.

Results

eQTL indicate polygenic transcriptome modulation

To understand how natural genetic variation in the pathogen influences both the host and pathogen transcriptomes, we performed expression GWA across all genes expressed in each species within the *A. thaliana*–*B. cinerea* pathosystem. This analysis incorporated the *z*-scaled expression profiles of 9267 *B. cinerea* genes and 23,947 Col-0 *A. thaliana* genes, mapped as individual traits across 96 diverse *B. cinerea* isolates. For each trait, we used a GEMMA univariate linear mixed model that estimates the significance of effects of each SNP on the focal trait as a *P*-value after accounting for potential effects of population structure within the *B. cinerea* isolates (Zhou and Stephens 2012). For GEMMA, we used a previously described *B. cinerea* genome-wide SNP data set of 237,878 SNPs with a conservative minimum MAF of 0.20 (Atwell *et al.* 2018 preprint; Soltis *et al.* 2019).

Using permutation estimated thresholds (see *Materials and Methods*), we identified 461,723 *B. cinerea* SNPs associated with transcriptional variation in 1616 *B. cinerea* genes and 978,693 SNPs associated with 5213 *A. thaliana* genes (Figure 1). In comparison to the random permutations, this suggests a false discovery rate for SNPs of ~1% using the $1.96e^{-5}$ threshold for *B. cinerea* (4756 permutation/461,723 empirical) and 1.6% using the $2.9e^{-5}$ threshold for *A. thaliana* (16,446 permutation/978,693 empirical).

The range of significant SNPs per transcript is wide with 0–16,814 SNPs (median = 0, mean = 49.8) for *B. cinerea* and 0–24,622 SNPs (median = 0, mean = 40.8) for *A. thaliana*. The observation of transcript variation associated with a large number of significant SNPs has been seen in other eQTL analyses (Figure S1) (West *et al.* 2007; Wang *et al.* 2018b). Given the wide range of significant SNPs per transcript and how it could shape the analysis, we restricted the analysis of the top SNP per transcript (SNP with the lowest *P*-value) to survey genomic patterns. We further tested how conservative our threshold was by comparing the permuted top to the empirical top *P*-value across all SNPs per transcript. This showed that the *P*-value of empirical top SNP per trait for 69% of the *B. cinerea* transcripts, and for 58% of the *A. thaliana* transcripts, is lower than permuted transcripts top SNP across all five permutations. By comparing the pattern of trans-hotspots across the genome using all the top SNP per transcripts and only top significant SNP per transcripts showed a similar image of the trans-hotspots, with SNP below the threshold reinforcing the trans-hotspots (Figure 2, A and B).

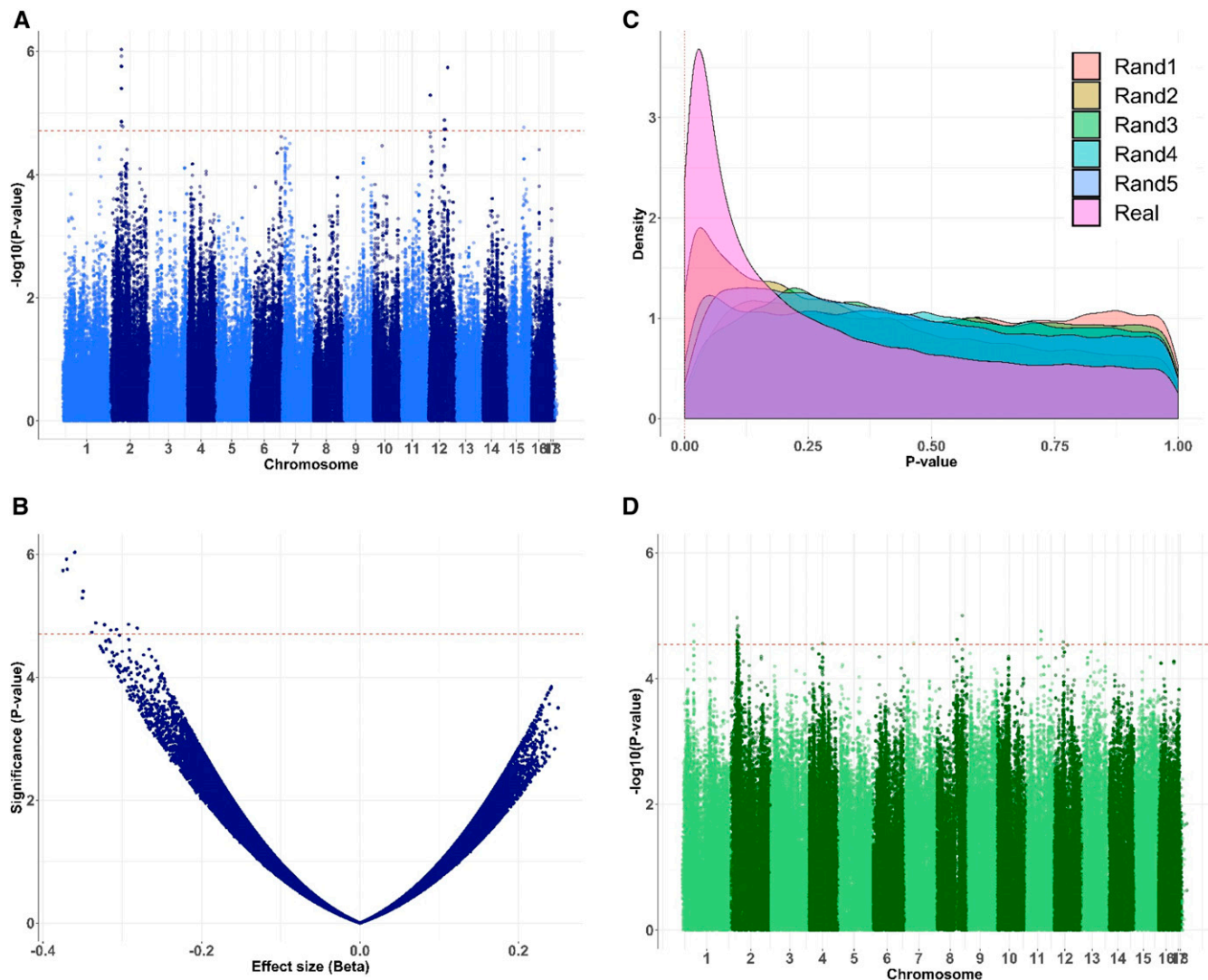


Figure 1 Examples of expression-based GWA mapping in *B. cinerea* genome. The two transcripts plotted correspond to the mean of significant SNPs with 50 and 41 significant SNPs, respectively. (A–C) GWA results for the mapping of the expression of *Bcin01g09490*, a methyltransferase. Red dashed lines indicate significance at $P = 1.96e-05$. (A) Manhattan plot of log-scaled P -values of SNPs associations across the 18 chromosomes of *B. cinerea*. (B) Relation between significance and effect size as estimated by GEMMA. (C) P -value density in the real expression data vs. the five randomizations. (D) A cross-species eQTL Manhattan plot for the host's expression of *AT1G28370* (an ethylene response factor) across the 18 chromosomes of *B. cinerea*. The red dashed line indicates significance at $P = 2.9e-5$.

Therefore, we focused on all the top significant SNP per transcript for the remaining analysis. This restrictive analysis on SNPs that are most likely to be associated with transcript variation accounts for the fact that nearly every transcript has a highly significant heritability ascribed to the *B. cinerea* genome (Zhang *et al.* 2017; Zhang *et al.* 2019). To account for the potentially low resolution in genomic signal encompassed by the single top SNP, we also assessed any general pattern using the top 10 SNPs.

Absence of observed transcriptome cis-effect

A hallmark of within-species eQTL mapping studies using either GWA or structured mapping populations in a wide range of species is the occurrence of *cis*-diagonals (Brem

et al. 2002; Schadt *et al.* 2003; Monks *et al.* 2004; Keurentjes *et al.* 2007; West *et al.* 2007; Zou *et al.* 2012). This occurs because polymorphisms proximal to the gene, *cis*-eQTL or *cis*-SNPs, frequently have large effects on the transcripts accumulation measured. To test if our analysis identifies a similar *cis*-diagonal in the *B. cinerea* transcriptome, we plotted the position of the transcript's genomic position against the top SNP for all the *B. cinerea* transcripts. As *cis*-diagonals are generally enriched in SNPs with the largest effects on transcript abundance, the top GWA SNP should identify this diagonal, assuming there are no intervening technical or biological issues. However, there was no evidence supporting a *cis*-diagonal (Figure 2). This pattern held whether we examined the top SNP per transcript (Figure 2A)

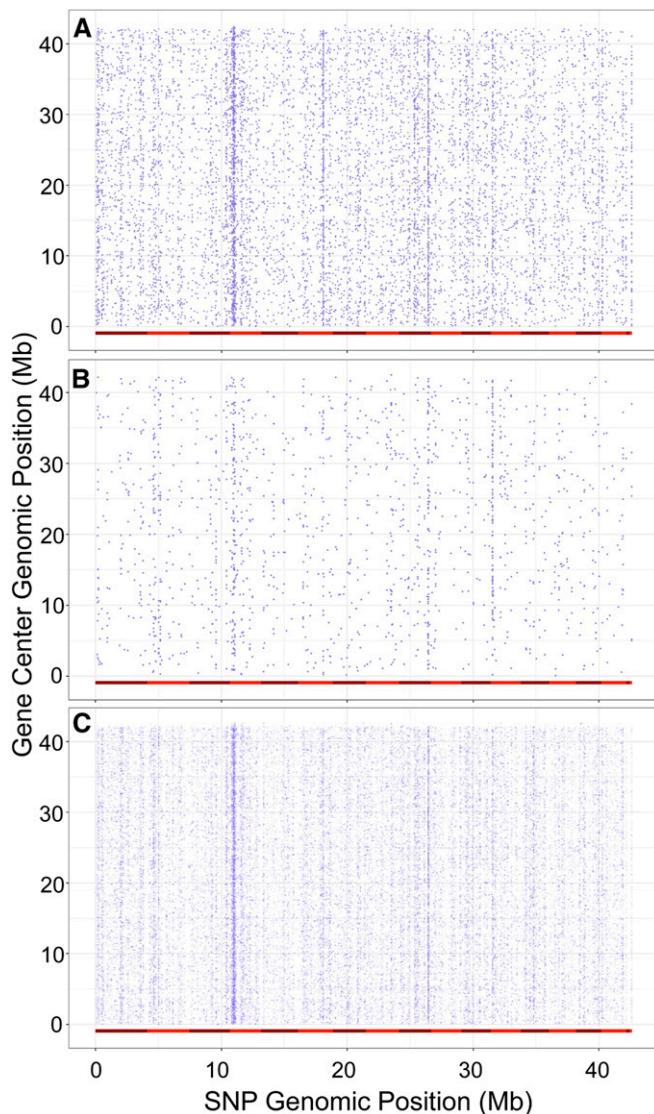


Figure 2 Comparing *B. cinerea* gene center to position of top associated SNP. For all 9267 transcripts mapped to the *B. cinerea* genome, we retained only the SNPs with highest probability (lowest P -value) of significant effect on expression for each transcript. (A) The single top SNP per all transcripts. (B) The single top SNP for only those transcripts that have at least one SNP crossing the $1.96e^{-5}$ significance threshold. (C) The top 10 SNPs per all transcripts. The 18 chromosomes of *B. cinerea* are delimited by differentially shaded red bars along the x -axis, while positions indicate individual SNPs. The y -axis depicts the same chromosome alignment, but positions are the center of each mapped transcript. Vertical striping of SNP positions indicates genomic locations of putative *trans*-eQTL hotspots.

or the top 10 SNPs per transcript (Figure 2C). In contrast, there is evidence for *trans*-eQTL hotspots: distant loci that modulate expression variation across many of the pathogen genes (Figure 2).

Because any significance threshold may complicate the identification of a *cis*-signal, we converted all the P -values assigned to SNPs per transcript into their percentile rank. This conversion was performed by sorting all P -values in ascending order, assigning ranks values to each P -value and

converting these values to percentage of the total number of SNPs. We then identified all the SNPs within each *B. cinerea* transcript or within 1 kb of the start or end of the transcript, and extracted the percentile ranks for these SNPs. The number of local SNPs ranged from 1 to 352 SNPs, with an average of 16 local SNPs per transcripts. The distribution of ranks across all the local SNPs was flat and similar to a random sampling of P -values, further indicating no general *cis*-enrichment within this data set (Figure S2). We next limited this percentile rank analysis to the local top SNP for each transcript because each *cis*-region may have only one or a few true associations masked by noncausal SNPs. As a random comparison, we simulated a data set of percentile rank by random draw of 16 SNPs out of all SNPs within defined *cis*-defined boundaries and repeated this 9267 times (the number of *B. cinerea* transcripts). For each simulated transcript, we extracted the top SNP and repeated the whole simulation five times. This showed that that *cis*-SNPs are not enriched for higher significance than the genome, and might even be less frequent than expected by random chance (Figure S2). As such, we do not detect evidence for overrepresentation of *cis*-effect loci.

Allelic heterogeneity and structural variation potentially masking cis-effects

Haplotype heterogeneity or allele frequency may complicate the accurate identification of *cis*-polymorphisms and hide a *cis*-diagonal (Chan *et al.* 2010; Rivas *et al.* 2011; Visscher *et al.* 2017). To assess these possibilities, we used three clustered biosynthetic pathways: the botcinic acid biosynthetic pathway (13 genes, 55.8 kb, chromosome 1), botrydial biosynthetic pathway (7 genes, 26 kb, chromosome 12), and a putative cyclic peptide pathway (10 genes, 46.5 kb, chromosome 1) (Deighton *et al.* 2001; Colmenares *et al.* 2002; Porquier *et al.* 2016; Zhang *et al.* 2019). These pathways have known presence/absence polymorphisms segregating in the studied *B. cinerea* population and were expected to hold *cis*-patterns, yet none were detected by our GWA analysis (Siewers *et al.* 2005; Pinedo *et al.* 2008). Additionally, genes of the botcinic acid and botrydial biosynthetic pathways are variable across the *Botrytis* genus (Valero-Jiménez *et al.* 2019). Critically, the transcription profiles of the genes within these pathways are highly correlated across isolates, suggesting pathway-specific control of expression variation (Zhang *et al.* 2019). This known presence/absence variation suggested that the absence of *cis*-SNPs in these loci may represent false negatives. To test if these pathways have undetected *cis*-SNPs, we extracted all the SNPs within each biosynthetic cluster and investigated haplotype diversity across the 96 *B. cinerea* isolates.

We first investigated the botcinic acid cluster that contains multiple distinct haplotypes (Figure 3A). To test for haplotype-specific effects on transcript expression within a pathway, we z -scaled each individual transcript and averaged across all the genes in the pathway to get a pathway expression value (Kliebenstein *et al.* 2006). The pathway expression

identified a group of 12 isolates (cluster 4 in Figure 3B) with distinctly lower expression level than the other groups. Investigating the short-read sequences and SNPs showed that these 12 isolates share a 78.6-kb deletion that removes the entire botcinic acid biosynthetic cluster, from *Bcboa1* to *Bcboa17* (Figure 3). After accounting for this major deletion, we found no remaining significant effect of group membership in the remaining groups on the expression profile ($F(1,74) = 0.36$; $P = 0.55$). However, within each of these groups, independent isolates have low pathway expression, suggesting loss-of-expression polymorphisms (e.g., isolates Noble Rot and Apple517) (Figure 3B). While each of these isolates contain independent smaller deletions, the origins of the loss of expression of the genes of botcinic acid pathway remain unknown. Functional analysis of the botcinic acid biosynthetic cluster has thus far identified one transcription factor (*Bcboa13*) that controls the expression of the cluster (Porquier *et al.* 2019). However, none of the SNPs within or near *Bcboa13* were significantly associated with variation in expression of the botcinic acid biosynthesis genes. These patterns may suggest undetected *cis*-effect polymorphisms in addition to the large common deletion and independent additional events.

We then investigated the botrydial and putative cyclic peptide biosynthetic pathways for additional evidence of undetected *cis*-acting genetic variation. These two pathways exhibit a lack of *cis*-effect SNP patterns similar to the botcinic acid pathway. Hierarchical clustering within each of these pathways by SNP variation divided the isolate population into two groups that were not associated with mean pathway expression (Figure S3 and Figure S4, both bipartite divisions were supported by >95% of bootstraps). While there was no obvious structural variation in the botrydial pathway, the cyclic peptide pathway contained small deletions within the intergenic regions that correlate with low expression. Furthermore, two isolates with partial deletions within the genes early in the pathway exhibited very low pathway expression (1.05.16 and 1.05.22) (Figure S4).

Overall, we identified *cis*-acting deletions in two of three of the clustered biosynthetic pathways. This highlights the potential of structural variants that often fall below the minor allele cutoffs and compromise the detection of *cis*-effects by GWA within *B. cinerea*. Testing whether insertion and deletion events account for the majority of localized control of expression variation would require both long-read sequencing to accurately identify these structural variants and computational approaches that can blend SNP and insertion/deletion information.

Detection of *trans*-eQTL hotspots

Using the top SNP per transcript, the GWA analysis identified a strong signature of *trans*-eQTL hotspots (Figure 2): polymorphisms that may influence the regulation of numerous genes in *trans*. These hotspots were detected using both the *B. cinerea* and *A. thaliana* transcriptome with solely the top SNP per transcript (Figure 4 and Figure 5). Using a permutation

approach, we fixed conservative thresholds for eQTL hotspots to 20 transcripts for *B. cinerea* and 150 transcripts for *A. thaliana*. This identified 25 *trans*-eQTL hotspots dispersed across the *B. cinerea* genome that modulate either the host (12 SNPs) or pathogen (13 SNPs) transcriptomes (Figure 5, Figure 6, and Table 1). Hotspot SNPs had an average MAF of 27% (1% SE) with no correlation between the MAF and the number of transcripts associated with a particular SNP within either the *Arabidopsis* or *Botrytis* transcripts. Further, the SNPs at different hotspot were not cosegregating across the isolates.

The use of a cotranscriptome approach should theoretically identify *trans*-eQTL hotspot affecting *B. cinerea* transcripts that create an eQTL hotspot affecting the host's transcriptome. However, the GWA analysis detected no cross-species-connected eQTL hotspots across the two transcriptomes (Figure 5). This result was consistent using either the single top SNP or the top 10 SNPs per transcript (Figure S5). The absence of cross-species overlap in eQTL hotspots suggests that the pathogen's influence on the host's transcriptome is not limited to major interactions between *trans*-eQTL hotspots, but can involve molecularly constrained changes in the pathogen that are magnified in the host's response. One possible scenario is a SNP that alters the expression of a single effector gene or mechanism in the pathogen that does not affect the pathogen but instead affects the host. For example, altering expression of the botcinic acid biosynthetic cluster would alter the accumulation of that phytotoxic metabolite and cause large responses in the host. Similarly, mutations that do not affect transcript abundance, like missense polymorphisms that alter protein function, could equally lead to a lack of overlap in hotspots. Finally, it is possible that the high level of genetic variation in *B. cinerea* may decrease our power to detect the cross-species *trans*-eQTL hotspots. Deeper analysis into the transcriptome and downstream responses could elucidate how restricted responses in the pathogen transcriptome translate to sweeping responses in the host. Future studies using these eQTL hotspots as *a priori* candidates for control of transcript variation in both host and pathogen may need to increase power to detect more modulation overlap across the two transcriptomes.

eQTL hotspot modules

To better understand the *trans*-eQTL hotspots, we examined the genes influenced by each hotspot. We first collected the GO annotations within each species to test if the hotspot transcripts were enriched for some specific functionality. The *B. cinerea* GO annotations showed a preponderance of enzyme, signal peptides for secretion, and transcription factor annotations, but no specific molecular insights arose, largely because the majority of genes had no annotation (Table 1, Table S1, File S1, and File S4). In contrast, GO analysis of the *A. thaliana* transcripts showed that three of the hotspots have an overrepresentation of photosynthesis-related functions within their targeted genes (Table 1, File S2, and

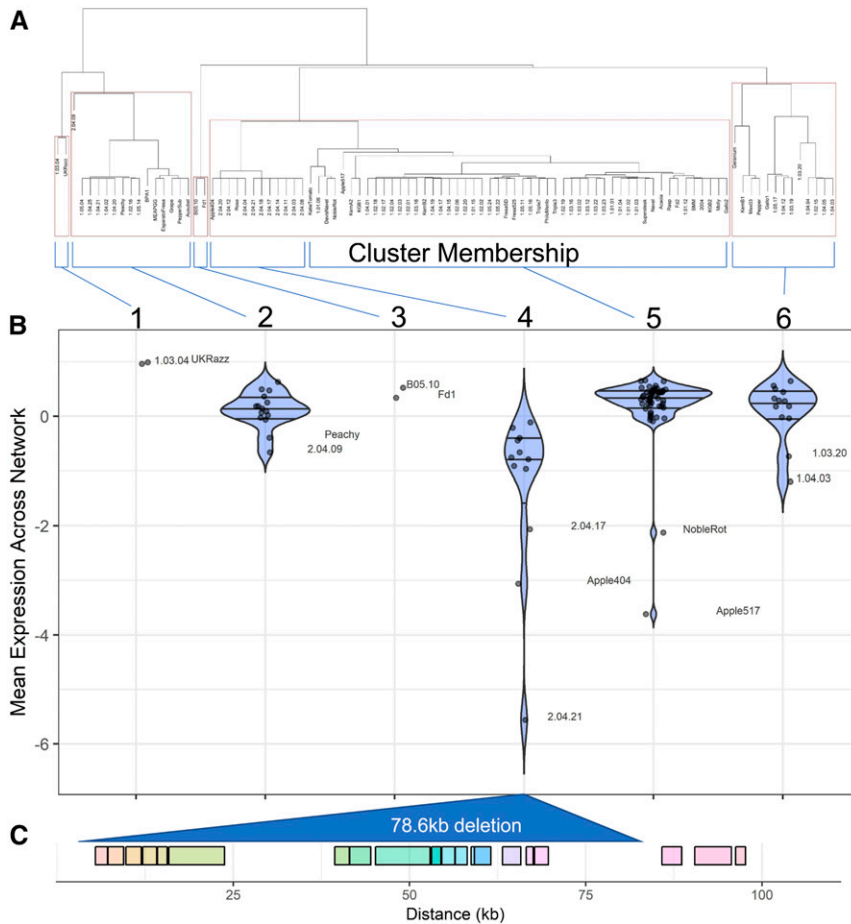


Figure 3 Analysis of the botcinic acid gene cluster on *B. cinerea* chromosome 1. (A) An SNP-based hierarchical clustering of *B. cinerea* isolates. The hierarchical clustering was based on unweighted pair group method with arithmetic mean distance, with 1000 bootstrap replications. Red boxes indicate clustering with >95% confidence. (B) Violin plots of botcinic acid network expression within *B. cinerea* groups. The mean network expression was obtained by converting expression of each gene across the isolates into its corresponding z-score and then averaging across the z-scores. Below is the gene model of the gene cluster, with each rainbow-colored box delimiting single genes (including Bcboa1 to Bcboa13 and 5 additional genes). The large 78.6-kb deletion in the isolate cluster 4 major deletion is indicated as a triangle.

File S3). Downregulation of photosynthesis transcripts is a hallmark of plant immune processes (Bilgin *et al.* 2010; Jiang *et al.* 2017). Two other *A. thaliana* hotspots primarily affect genes associated with abiotic stress responses. Only two of the *A. thaliana* hotspots influence expected plant defense loci, including chitin response and microbe defenses. This suggests that the *B. cinerea* genes underlying these hotspots specifically influence defined networks within the host.

In previous work, we had defined key transcript modules within both the host and pathogen transcriptomes that connected to virulence (Zhang *et al.* 2017; Zhang *et al.* 2019). Thus we tested for overlap between the *trans*-eQTL hotspot-defined modules and previously defined transcript modules using a hypergeometric enrichment test (Zhang *et al.* 2017; Atwell *et al.* 2018 preprint; Zhang *et al.* 2019). Nine of the 11 *B. cinerea* eQTL hotspots were enriched for transcripts present in one or more of four major *B. cinerea* coexpression modules identified when grown on *A. thaliana* (Figure 6). An additional six *B. cinerea* modules did not share any transcript membership with detected eQTL hotspots. Similarly, nine of the *A. thaliana* eQTL hotspots were enriched for transcripts from two of the major *A. thaliana* modules when infected with *B. cinerea* (Figure 6). These two *A. thaliana* modules contain genes that function in jasmonate/salicylic acid

signaling processes and camalexin biosynthesis (network I), or photosynthesis (network IV). Interestingly, these links are not limited to a single hotspot, but have strong connections across several different eQTL hotspots, suggesting that these *A. thaliana* modules are influenced by the pathogens polygenic architecture (Figure 6).

eQTL hotspot candidate genes

To generate working hypotheses on the possible causal basis of the eQTL hotspots, we investigated the candidate genes underlying the associated SNPs. The 12 *B. cinerea* hotspots that influence *A. thaliana* transcripts are located within 11 genes, including four enzymes and two genes associated with isolate compatibility (Table 1). The 13 *B. cinerea* hotspots that influence *B. cinerea* expression profiles were associated with 11 genes, including four enzymes (Table 1). However, only one of these 22 genes was previously linked to virulence functions in *B. cinerea* or other fungi. *Bccwh41* (*Bcin16g01950*), a glycoside hydrolase whose homolog shows increased expression in virulent strains of *Ustilago maydis* on *A. thaliana* (Martínez-Soto *et al.* 2013). To test if any of these 22 eQTL hotspot candidate genes may influence virulence in *B. cinerea*, we used the existing coexpression and virulence data to compare the level of expression of these

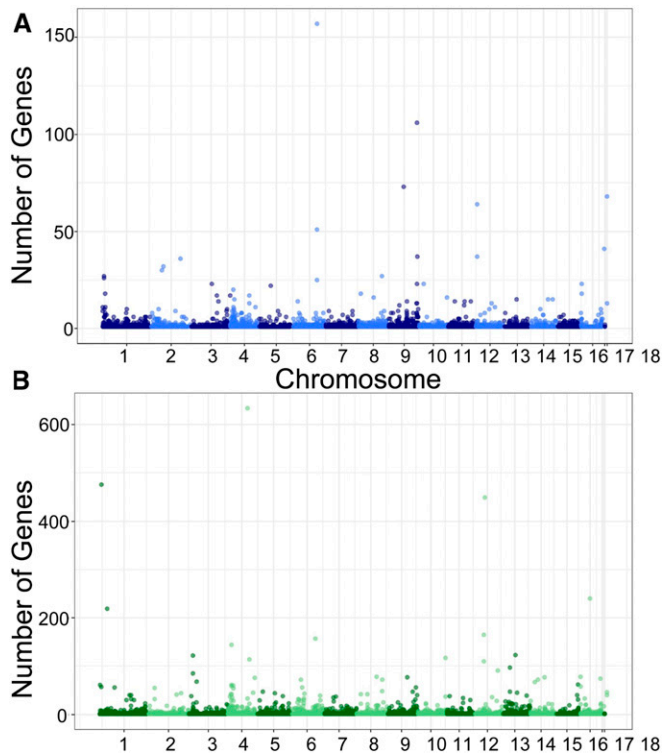


Figure 4 Frequency and positions of *trans*-eQTL hotspots in the *B. cinerea* genome. (A) Number of *Botrytis* transcripts associated with *trans*-eQTL SNPs aligned along the 18 chromosomes of *B. cinerea* (x-axis). (B) Number of *Arabidopsis* transcripts associated with cross-species eQTL SNPs aligned along the 18 chromosomes of *B. cinerea* (x-axis).

22 genes to existing virulence measurements (Zhang *et al.* 2017; Atwell *et al.* 2018 *preprint*; Zhang *et al.* 2019). Transcript accumulation for three *B. cinerea* hotspot genes and two of the *A. thaliana* hotspot genes are strongly positively correlated to lesion size variation, while none are negatively correlated with lesion size (Table 1) (Zhang *et al.* 2019). Further, we utilized a previous GWA analysis of virulence of these same isolates on *A. thaliana* to test for any overlap. This showed that one of the *B. cinerea* hotspot genes (*Bcin16g00010*, *SsuA*/THI5-like) is a top GWA hit, controlling lesion size across host genotypes and association methods (Table 1) (Atwell *et al.* 2018 *preprint*). Together, this suggests that these genes are likely candidates for controlling both the host and pathogen transcriptomes.

Discussion

Dispersed interactions across host and pathogen genomes

Using cotranscriptome GWA, we identified 25 *trans*-eQTL hotspots dispersed across the *B. cinerea* genome that modulate either the host or pathogen transcriptomes. This contrasts with previous cross-species eQTL studies, which identified one or only a few cross-species eQTL hotspots (Wu *et al.* 2015; Guo *et al.* 2017). Further, most of the *B.*

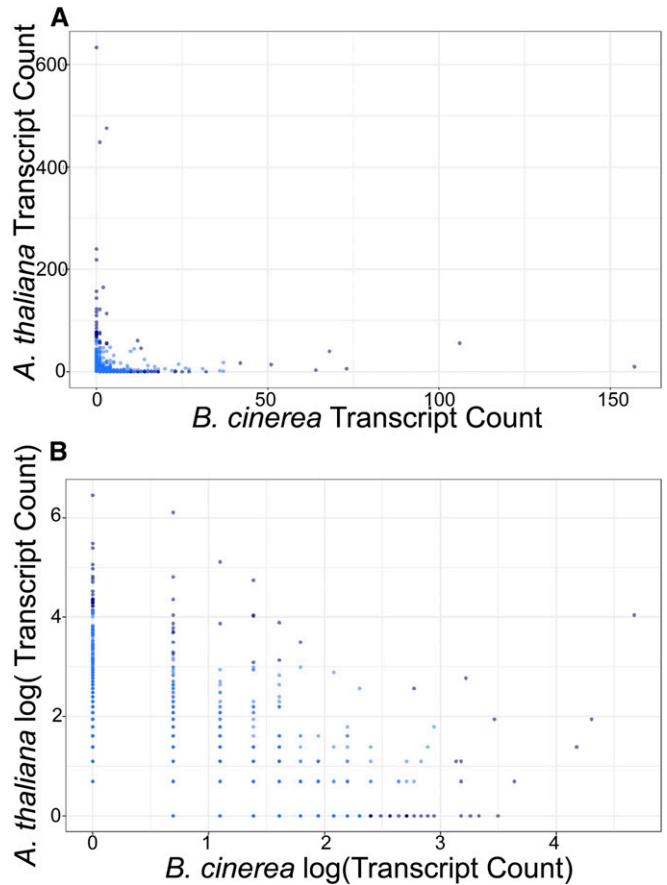


Figure 5 Cross-species hotspot comparison in the *B. cinerea* genome. For each SNP that is a top hit for one or more transcripts, the number of associated transcripts is counted, across both the *B. cinerea* transcriptome and the *A. thaliana* transcriptome.

cinerea genetic variation detected in our study is distant from the affected *B. cinerea* transcripts, *i.e.*, located in *trans*. These *trans*-eQTL hotspots influence expression variation for five major *B. cinerea* modules containing genes dispersed across the genome (Zhang *et al.* 2019). In particular, the eQTL hotspots influenced the expression of specific *B. cinerea* co-expression networks (vesicle/virulence, translation/growth, exocytosis regulation, and peptidase). These candidate polymorphisms are spread throughout the genome and the detected eQTL hotspots are not in regions of the genome with significantly elevated genetic variation. Further, the genetic targets of these eQTL are dispersed across the plant and pathogen genomes (Zhang *et al.* 2019). As such, *B. cinerea* does not fit the model of what might be expected in filamentous fungi showing multiple-speed genome evolution due to varying selective pressures influencing the genome. In such specialist fungi with closer coevolution to their host species than *B. cinerea*, diverse fungal virulence effectors are clustered in regions of the genome containing enhanced rates of mutation and polymorphism, while the rest of the genome shows slower evolutionary rates (Dong *et al.* 2015). If this scenario were true for the *Botrytis* genome, it would predict clustering of the GWA hits to a few locations rather than

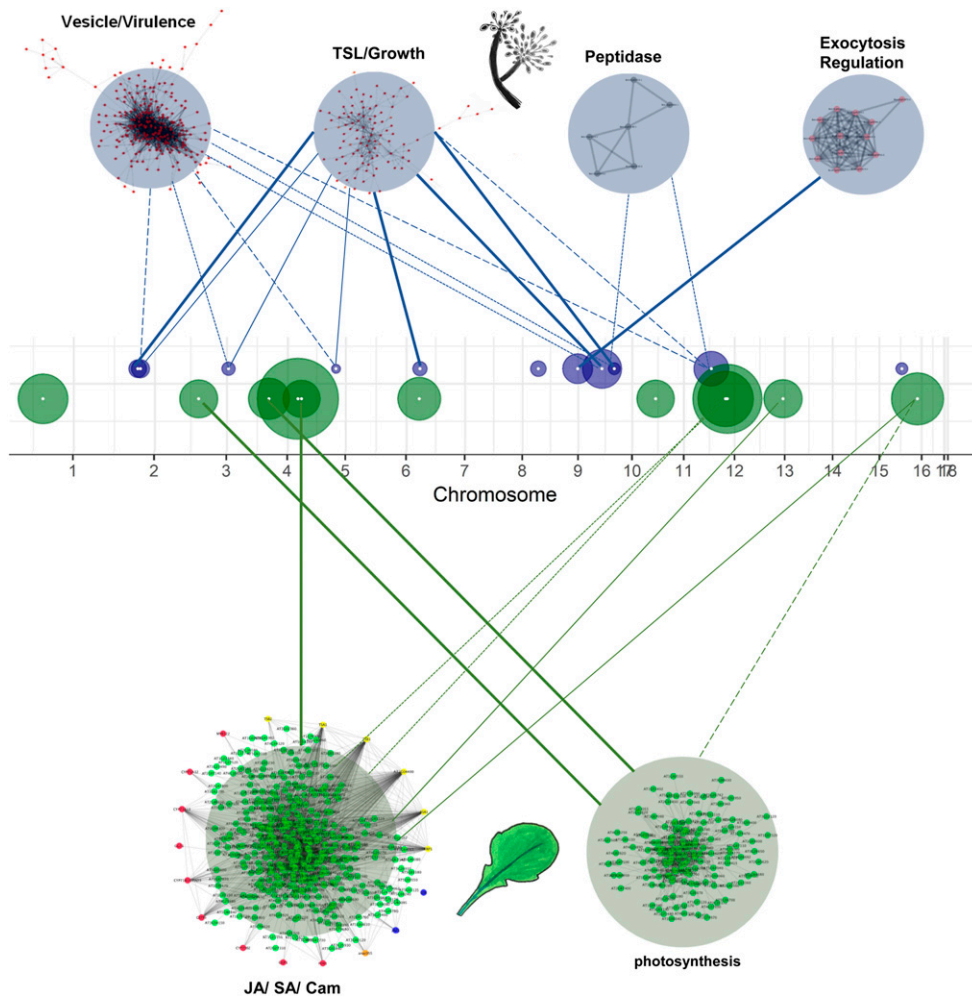


Figure 6 Genes associated with eQTL hotspots are in virulence and defense coexpression networks. Circles along the *B. cinerea* 18 chromosomes are eQTL hotspots, centered at the gene containing the eQTL, and with radius proportional to the number of transcripts associated with this hotspot. The gene center is marked with a white dot. Hotspots for *B. cinerea* transcripts are drawn in blue, hotspots for *A. thaliana* transcripts are drawn in green. The network names are based on biological functions from gene ontology analysis of network members, from Figure 4 of Zhang *et al.* (2019) and Figure 6 of Zhang *et al.* (2017). The *A. thaliana* networks depicted are the most inclusive of the host-dependent networks, from *npr1-1*. Links between hotspots and coexpression networks are drawn according to the number of genes shared between them. Variable line weight represents the percent of hotspot target genes shared with the coexpression network; 1–25% is dashed, 25–50% is dotted, 50–75% is solid, 75–100% is bold.

distribution across the genome, as was found. This is consistent with previous findings of high genome-wide diversity in *B. cinerea*, and virulence and host specificity mapping to large swaths of the pathogen genome, including 16 of 18 chromosomes (Atwell *et al.* 2018 preprint; Caseys *et al.* 2020 preprint). Similar eQTL analyses in the multispeed genome filamentous fungi are required to test whether eQTL in pathogens with a multispeed genome truly cluster within the highly polymorphic regions. Together, these findings provide evidence for polygenic *trans*-regulation of gene expression in *B. cinerea* interactions that then coalesces around specific transcriptional modules to influence virulence.

Polygenic modules and pleiotropy in cross-species eQTL

Previous pathogen eQTL studies identified qualitative patterns whereby each host expression profile was explained by a single major-effect pathogen locus (Guo *et al.* 2017) or each pathogen eQTL connected to a specific host network (Wu *et al.* 2015). In contrast, cotranscriptome GWA with *B. cinerea* identified a more complex picture, with numerous *trans*-eQTL hotspots altering multiple transcriptome modules in either the host or the pathogen. This suggests that the

polygenic architecture of the pathogen may partially function by influencing these defined modules rather than working as thousands of individual genes, each independently targeting the host. It remains to be ascertained if this system functions to create robustness in these connections in the face of changes to the pathogen or host genetics, or if this is an indication of existence of a discrete set of interaction mechanisms between the host and the pathogen.

Candidate causal loci encode diverse mechanisms

Querying the putative function of the candidate loci underlying the different *trans*-eQTL hotspots identified an array of potential molecular mechanisms. While one might assume that transcription factors are the most likely genes for containing genetic variation that would lead to *trans*-eQTL hotspots, we instead found enrichment for enzyme-encoding genes among these loci. Eight enzyme-encoding genes contained the SNPs for four of the *B. cinerea* *trans*-eQTL and four of the *A. thaliana* *trans*-eQTL hotspots. Interestingly, these enzymes have potential activities in sugar release from the plant cell wall, or reactions involving sugar phosphates (Table 1). In addition to enzymatic activity, four of the *Botrytis*

Table 1 Annotation of the genes identified from *B. cinerea* and *A. thaliana* eQTL hotspots

Hotspot gene	Hotspot SNP	<i>Botrytis</i> transcripts	<i>Arabidopsis</i> transcripts	Name	Gene function	Virulence correlation	Virulence GWA	<i>Arabidopsis</i> GO overrepresentation
Bcin01g01610	629157	0	219		Glucose/ribitol dehydrogenase	NA		Amylopectin, glycogen, chlorophyll, chloroplast, photosynthesis
Bcin02g02480	910858	31	6			$P = 0.008$		NA
Bcin02g02850	1032242	32	0		Fructosamine-3-kinase	NA		NA
Bcin03g00960	336467	1	122		GTP cyclohydrolase I	NA		Carotenoid biosynthesis, chloroplast organization
Bcin03g05020	1695103	24	15			NA		NA
Bcin04g00830	310509	0	144		NACHT nucleoside triphosphatase	NA		Nucleic acid metabolism
Bcin04g04700	1633072	0	634		Heterokaryon incompatibility	NA		Photosynthesis, light, translation
Bcin04g05160	1791561	3	114			NA		Chitin response
Bcin05g02780	1015184	22	2			NA		NA
Bcin06g05680	1952544	0	157		WLM	$P < 0.001$		NA
Bcin06g05790	1988179	25	0			NA		NA
Bcin08g05340	2014228	27	0		F-box domain	NA		NA
Bcin09g03390	1235841	73	6		6-Phosphogluconate dehydrogenase	NA		NA
Bcin09g06590	2330312	106	56		SNF2-related; Helicase, superfamily 1/2	NA		Metabolism
Bcin09g06590	2334368	23	0		SNF2-related; Helicase, superfamily 1/2	NA		NA
Bcin10g00940	383007	23	1			$P < 0.001$		NA
Bcin10g05900	2268522	0	117		Winged helix-turn-helix transcription factor	NA		Water stress
Bcin12g00330	115491	37	1	Bcpat1	Topoisomerase-II-associated protein PAT1	$P < 0.001$		NA
Bcin12g00330	115511	64	3	Bcpat1	Topoisomerase-II-associated protein PAT1	$P < 0.001$		NA
Bcin12g02130	758420	0	110			NA		Response to stimulus
Bcin12g02130	760499	2	265			NA		Jasmonic acid, fungal response, microbe defense, biotic stress
Bcin12g02340	842369	1	449	Bccds1	Phosphatidate cytidyltransferase	$P < 0.001$		Primary metabolism, amino acid biosynthesis, salt stress, biotic response
Bcin13g02930	1026752	0	123		SET domain	NA		Transport
Bcin16g00010	76596	23	2		SsuA/THI5-like	NA	Yes	NA
Bcin16g01950	787512	0	240	Bccwh41	Glycoside hydrolase, family 63	NA		Photosynthesis

Each row identifies a significant eQTL hotspot SNP associated with transcript variations in *B. cinerea* or in *A. thaliana*. Gene functions are from BotPortal, *Arabidopsis* GO overrepresentation are from PANTHER. The table lists if the gene's transcript is correlated with virulence or if the SNP was associated with virulence via GWA (Atwell *et al.* 2018 preprint; Zhang *et al.* 2019).

trans-eQTL candidate genes encoded transcriptional regulators. *Bcin10g05900*, a putative winged helix transcription factor (TF), is predicted to have pathway-specific effects, while the other three are putative general transcriptional regulators, including *Bcin12g00330*, a putative

topoisomerase-II-associated protein PAT1, and *Bcin09g06590*, a putative helicase (Table 1). Interestingly, the putative winged helix TF *trans*-eQTL hotspot alters the expression of *A. thaliana* genes associated with water deprivation responses. This suggests that this putative winged helix TF may influence a specific

virulence factor that influences this *A. thaliana* network. Interestingly, while the candidate genes connect to processes that likely influence virulence, none of them have been explicitly shown to influence virulence in *B. cinerea*. Future work is necessary to test these loci and discover the mechanisms by which they influence virulence and the host/pathogen cotranscriptome.

Complications in detection of cis-acting loci

The vast majority of eQTL studies identify a strong signature of cis-acting loci. However, in *B. cinerea*, the main detected pattern was trans-eQTL, with few identified cis-eQTL. A deeper investigation suggested that this may be due to genetic factors that complicate the ability to identify cis-acting SNPs. *B. cinerea* has high haplotype diversity, and in three gene clusters investigated, potential rare cis-acting variants fell below the MAF cutoff for GWA, potentially leading to false-negative detection errors (Tabangin *et al.* 2009). Further, a number of deletions were found in this locus, which complicates the detection of a cis-eQTL signature by introducing non-SNP variation unused by current GWA algorithms. Additional cis-acting variants may also be hidden by undetected transposon variation (Porquier *et al.* 2019).

A full understanding of the pattern of potential cis-acting loci in *B. cinerea* would require a detailed investigation of structural variation by incorporating long-read sequencing in a larger population size. Additionally, the GWA algorithms would need to be written to allow for simultaneous use of both SNP and presence/absence polymorphism data; one option is to code deletions as an additional state for each genotyped variant (Wang *et al.* 2018a). It should be noted that these same difficulties would also create undetected trans-eQTL. This does suggest that there is likely a significant fraction of undetected cis-eQTLs within *B. cinerea*, caused by the high polymorphism rate within this species.

To summarize, previous work in the *A. thaliana*–*B. cinerea* pathosystem established connections between host polymorphisms and lesion growth, between gene expression and lesion size, and between transcriptomes of the host and pathogen (Corwin *et al.* 2016b; Zhang *et al.* 2017; Zhang *et al.* 2019). This study establishes a foundation to study how genetic variation in the pathogen can manipulate the host transcriptome to influence disease progression. In the *Arabidopsis*–*Botrytis* pathosystem, this connection had a preponderance of trans-acting polymorphisms with mainly moderate to small effects, suggesting that a polygenic architecture underlies the transcriptome variation, similar to the polygenic architecture observed for virulence. Using previously defined transcriptome modules showed that there may be a modular structure to these effects, with specific pathogen SNPs linking to specific modules in either the host or the pathogen. However, future validation work will be required to test the directionality and mechanism of this cross-talk. Similar work in other systems will help to build

our functional knowledge of cross-kingdom communication between host and pathogen.

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