Integrated soybean transcriptomics, metabolomics, and chemical genomics reveal the importance of the phenylpropanoid pathway and antifungal activity in resistance to the broad host range pathogen *Sclerotinia sclerotiorum*.

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Sclerotinia sclerotiorum, a predominately necrotrophic fungal pathogen with a broad host range, causes a significant yield limiting disease of soybean called Sclerotinia stem rot (SSR). Resistance mechanisms against SSR are poorly understood, thus hindering the commercial deployment of SSR resistant varieties. We used a multiomic approach utilizing RNA-sequencing, Gas chromatography-mass spectrometry-based metabolomics and chemical genomics in yeast to decipher the molecular mechanisms governing resistance to S. sclerotiorum in soybean. Transcripts and metabolites of two soybean recombinant inbred lines, one resistant, and one susceptible to S. sclerotiorum were analyzed in a time course experiment. The combined results show that resistance to S. sclerotiorum in soybean is associated in part with an early accumulation of JA-Ile ((+)-7-iso-Jasmonoyl-L-isoleucine), a bioactive jasmonate, increased ability to scavenge reactive oxygen species (ROS), and importantly, a reprogramming of the phenylpropanoid pathway leading to increased antifungal activities. Indeed, we noted that phenylpropanoid pathway intermediates such as, 4-hydroxybenzoate, ferulic acid and caffeic acid were highly accumulated in the resistant line. In vitro assays show that these metabolites and total stem extracts from the resistant line clearly affect S. sclerotiorum growth and development. Using chemical genomics in yeast, we further show that this antifungal activity targets ergosterol biosynthesis in the fungus, by disrupting enzymes involved in lipid and sterol biosynthesis. Overall, our results are consistent with a model where resistance to S. sclerotiorum in soybean coincides with an early recognition of the pathogen, leading to the modulation of the redox capacity of the host and the production of antifungal metabolites.
Resistance to plant fungal pathogens with predominately necrotrophic lifestyles is poorly understood. In this study, we use Sclerotinia sclerotiorum and soybean as a model system to identify key resistance components in this crop plant. We employed a variety of omics approaches in combination with functional studies to identify plant processes associated with resistance to S. sclerotiorum. Our results suggest that resistance to this pathogen is associated in part with an earlier induction of jasmonate signaling, increased ability to scavenge reactive oxygen species, and importantly, a reprogramming of the phenylpropanoid pathway resulting in increased antifungal activities. These findings provide specific plant targets that can exploited to confer resistance to S. sclerotiorum and potentially other pathogens with similar lifestyle.
INTRODUCTION

*Sclerotinia sclerotiorum* (Lib.) de Bary, is a plant fungal pathogen with a predominately necrotrophic lifestyle and worldwide distribution that is known to infect over 400 plant species (1). On soybean (*Glycine max* (L.) Merr.), it causes sclerotinia stem rot (SSR), a significant and challenging yield limiting disease. Data suggest that 1.6 billion kilograms of soybean are lost each year to SSR in the US alone, making it the second most damaging disease of soybean (2,3). Globally, SSR can cause yield reductions as high as 60% (4). Soybeans are an important source of plant proteins and oils globally (5–7), these products can also be significantly affected by SSR (8). Management strategies against SSR rely largely on chemical control (9), though cultural practices such as crop rotation, seeding rate and row spacing modifications have been used with limited success (3). While genetic resistance is by far more sustainable, our understanding of resistance mechanisms against this pathogen is limited, and current commercial varieties lack adequate levels of resistance to SSR.

In the past two decades, studies have increased our understanding of *S. sclerotiorum* pathogenic development. *S. sclerotiorum* is a prolific producer of cell wall degrading enzymes (CWDEs) that contribute to its pathogenic success (10). In addition to its lytic repertoire, the pathogenic success of *S. sclerotiorum* relies on the key virulence factor oxalic acid (OA). Mutants that are defective in OA production are weakly pathogenic (11–14). OA was proposed to contribute to pathogenesis by acidifying host tissues and sequestering calcium from host cell walls, facilitating the action of CWDEs. However, recent studies showed that *S. sclerotiorum* relies on OA to induce programmed cell death (PCD) in the host to its own benefit (11,15). OA induced PCD is dependent on the production of reactive oxygen species (ROS) in the plant (14,15). This
points to the importance of the ROS machinery in the pathogenic success of this pathogen. Indeed, we showed that *S. sclerotiorum*, via OA, co-opts soybean NADPH oxidases to increase host ROS, and that the silencing of specific NADPH oxidases confers enhanced resistance to this fungus (16). In the addition to OA, other virulence factors targeting host responses are known to contribute to the pathogenicity of *S. sclerotiorum*. The necrosis and ethylene-inducing like proteins (NLPs) SsNEP1 and SsNEP2 induce cell death in tobacco leaves, thus contributing to disease establishment (17). The secreted chorismate mutase (SsCM1) and integrin-like protein (SsITL) were proposed to target Arabidopsis defenses by affecting salycilic acid and jasmonate signaling, respectively (18,19). The ability of *S. sclerotiorum* to detoxify plant chemical defenses has also been discussed (20). Unfortunately, these studies have largely focused on the fungal side of this interaction, and only provide glimpses into the plant mechanisms governing resistance to this pathogen, mainly in model plants. In soybean, Bi-parental linkage mapping led to the discovery of many quantitative trait loci (QTL) for resistance to this pathogen. Remarkably, a total of 103 QTL on 18 of the 20 soybean chromosomes have been recorded in SoyBase (21) with minimal overlap between QTL reported by different studies (9,22–26). Despite these efforts, gene level and mechanistic details of soybean resistance mechanisms against *S. sclerotiorum* remain unknown.

Recent advances in Next-Generation RNA sequencing (RNAseq) allow for cost-efficient and powerful examination of global differences in the transcriptional response to environmental cues. The application of RNAseq approaches in soybean-*S. sclerotiorum* interaction studies will, most assuredly, contribute to the development of molecular genetic resources crucial for mechanistic and translational research. Transcriptomics were used to study the interaction of *S. sclerotiorum* with other non-model plant hosts, including canola (27–29), pea (30), and common bean (31). While informative, these studies were solely based on gene expression comparisons,
and thus may not provide a complete picture of the flow of biologic information. When coupled with other omics approaches, such as metabolomics and chemical genomics, RNAseq can provide a much greater understanding of the biology, by integrating genetic responses of a particular interaction to functional consequences. Herein, we apply multi-omics approaches to identify and validate resistance processes against *S. sclerotiorum* in soybean lines generated in our breeding program. The identification of these processes will not only increase our understanding of the soybean-*S. sclerotiorum* interaction but will also facilitate the introgression of resistance into soybean varieties.

Our recent breeding efforts led to the identification of several recombinant inbred lines (RILs) highly resistant to *S. sclerotiorum* (9), using the soybean line W04-1002 as the main source of resistance (32). After multiple generations of greenhouse selection, we have chosen two RILs originating from the same cross; one resistant (91-145), and one susceptible (91-44) to SSR to complete this study. A comparative analysis using a combination of RNAseq, metabolomics, and chemical genomics in yeast, shows that resistance in 91-145 is associated in part with an earlier induction of jasmonate signaling, increased ability to scavenge ROS, and importantly, a reprogramming of the phenylpropanoid pathway leading to increased antifungal activities. We further discuss and provide evidence for the importance of antifungal compounds during the resistance response against this pathogen and show that this antifungal activity targets ergosterol biosynthesis in the fungus. We propose that the modulation of the identified pathways through RNAi/gene editing or overexpression approaches may be used to introgress SSR resistance in commercial soybean germplasm and possibly other host crops.
RESULTS

Disease development in the resistant (91-145) and susceptible (91-44) soybean lines

To determine soybean processes involved in resistance to SSR, two recombinant inbred lines (RIL) of soybean showing a differential response to *S. sclerotiorum* were chosen for this study. The resistant and susceptible selections were classified based on our previous germplasm selection study (9). Both the resistant 91-145 (R) and the susceptible 91-44 (S) lines were developed utilizing the SSR resistant parental line W04-1002 (P1) and LN89-5717 (PI 5745542), an SSR-susceptible parental line having other desirable pathogen resistance traits (9). Plants were inoculated using the cut petiole inoculation method (16) and infection progression was monitored in R and S soybean lines over seven days. Initial symptoms of SSR began appearing on the main stem 48 hours post-inoculation (hpi) as light brown lesions surrounding the point of inoculation, which spread as the disease progressed. By 96 hpi, extensive tissue colonization occurred along the main stem in the S line, while small restricted lesions with a red coloration were observed at the inoculation site in the R line (Fig. 1). By day 7, it was apparent that the R line had largely restricted fungal growth on the main stem and the red coloration observed at the site of infection had become more prominent, whereas the *S. sclereotiorum* infection had girdled the main stem of the S line (Fig. 1).

Global transcriptome analysis was conducted on *S. sclereotiorum* infected soybean stem tissue collected from both R and S lines at 0, 24, 48, and 96 hpi. Metabolomic analysis comparing the R and S lines was also conducted, similarly, stem tissues were collected at 0, 48 and 72 hpi.

Mapping and overview of RNA sequencing data

To determine transcript levels in the R and S soybean lines over the course of infection, RNA sequencing was conducted on twenty-four stem samples, consisting of three independent biological replicates for each of the time points selected. Depending on the time point, 64.9 – 89.7
8 million raw reads were generated, with 95.7 - 96.6% of reads mapping to the host reference genomes of soybean and *S. sclerotiorum* at all timepoints. On average, 96% of the total reads mapped uniquely to the soybean reference genome in the uninfected plants of both lines. In the S line, 91.6, 91.8, and 68.9% of the reads mapped to the soybean genome at 24, 48 and 96 hpi, respectively. In the R line, 92.8, 91.9, and 88.2% of the reads mapped to the soybean genome at 24, 48 and 96 hpi, respectively (Table 1). At 96 hpi the reads mapping to the fungal genome in the S line (27%) is significantly higher than those in R line (8%), which is expected considering the extensive colonization of the soybean stem by *S. sclerotiorum* in the S line, particularly at the later stages of the infection process. 4.4 % to 3.2 % of reads at different time point of study did not map to any reference genome and was therefore excluded (data not shown).

**Differentially expressed genes (DEGs) of soybean during infection**

A comparative analysis of differentially expressed genes (DEGs) was performed during the course of infection in both the R and S lines, as well as between these two lines at the specified time points to investigate the mechanisms underlying resistance to *S. sclerotiorum*. In the R line, we observed the maximum number of DEGs at 48 hours following inoculation (Table S1 and Fig. 2A). In contrast, the maximum DEGs in the S line occurred 96 hours post inoculation (Table S1 and Fig. 2B). Interestingly, at 96 hpi, the number of DEGs in the S line (14,050) were markedly higher than the R line (2,442), suggesting that the resistant response may be contributing to the reduced differential expression of host genes and the return to homeostasis at the later stages of infection. In total, 16,462 unique DEGs were identified in our lines during the course of infection (Fig. 2C). Among these, 7,319 were differentially regulated in both R and S lines, while 920 were strictly associated with the R line and 8,223 were S line specific (Fig. 2C). These results indicate that *S. sclerotiorum* infection causes a dramatic change in gene expression in soybean (~ 19% of...
total annotated genes in *Glycine max* genome), and suggests substantial differences between the resistant and susceptible soybean lines used in this study in response to *S. sclerotiorum* challenge.

Pairwise comparisons of DEGs were performed between R and S soybean lines at the selected time points (Table S2 and Fig. 2D). At 24 and 48 hpi, 1,039 and 803 DEGs were identified between the two lines, respectively. However, the number of DEGs sharply increased to 2,087 at 96 hpi. This is consistent with the distinct patterns of gene expression in the S and R lines at the later stages of infection (Fig. 2 A, B). Among the identified DEGs, 447, 191, and 1541 genes were found to be time-point specific between the R and S lines at 24, 48 and 96 hpi, respectively, while 340 genes were differentially expressed at all time points following pathogen challenge (Table S2 and Fig. 2D). The putative functions of these genes were determined using Phytozome (33), National Center for Biotechnology Information (NCBI), and Soybase (21) databases (Table S2).

**Gene ontology enrichment and biological process analyses**

We focus the reminder of this manuscript on direct comparisons between resistant and susceptible soybean lines, to single out potential processes associated with resistance to *S. sclerotiorum* in soybean. Soybean gene locus IDs identified through the DEG analysis were used to perform gene ontology (GO) enrichment analysis using the Soybase gene model data mining and analysis tool (34). A false discovery rate (FDR) value of 0.05 was used to identify significantly regulated GO biological processes, and individual GO processes were considered in this analysis if they were significantly enriched in at least one of the time points used (Table S3 and Fig. 3). The under or overrepresentation of DEGs in each GO category was determined based on the relative frequencies of the GO terms associated with the genes (34). Our data show an overrepresentation of genes in GO terms related to signal transduction (i.e. kinase signaling, phosphorylation), plant defense responses (i.e. response to chitin, fungi), phenylpropanoid
pathway (i.e. chalcones, anthocyanins, flavonoids, salicylic acid), ROS scavenging, and the biosynthesis/regulation of phytohormones (i.e. jasmonic acid, salicylic acid, ethylene). DEGs underrepresented throughout the course of infection include gene related to nucleotide binding and zinc ion binding. While the involvement of categories such as ROS, defense responses, and phytohormone regulation are not surprising, our results point to complex mechanisms of gene regulation associated with resistance to *S. sclerotiorum*. The differential regulation of phenylpropanoid pathway genes is intriguing and will be the subject of further characterization below.

**Metabolite profiling of susceptible and resistant soybean lines in response to *S. sclerotiorum*.**

Transcriptomic analysis was complemented by metabolite profiling of stem samples collected from our resistant and susceptible soybean lines following *S. sclerotiorum* infection. Simple changes in transcript levels might not always correlate with biological activity, but changes in metabolite flux are quantifiable outcomes that can directly explain disease phenotypes. Gas chromatography-mass spectrometry (GC-MS) analysis was performed to broadly evaluate metabolite profiles in control and infected soybean stems at 48 and 72 hpi. Three independent biological replicates were used for each time point. A total of 360 metabolites were detected, but only 164 could be identified based on available databases (Table S4). Each metabolite was characterized by its distinct retention time and mass to charge ratio (m/z). All the 164 identified metabolites found in infected soybean were also detected in non-inoculated stems and therefore are likely of plant origin. Despite this, we note that as disease progresses, contributions from *S. sclerotiorum* cannot be ruled out.

Metabo Analyst 3.0 (35) was used for the analysis of the 164 identified metabolites. One-way ANOVA using Fisher's least significant difference method (Fisher's LSD) identified 80
metabolites, the accumulation of which was significantly (FDR <0.05) affected by *S. sclerotiorum* inoculation at 48 and 72 hpi in R-line compared to S-line (Table S5). These metabolites included polar compounds, such as nucleotides, amino acids, alcohols, organic acids, and carbohydrates, along with nonpolar compounds including fatty acids, and long-chain alcohols (Table S5). The multivariate analysis of identified metabolites was performed using Partial Least Squares - Discriminant Analysis (PLS-DA). The first (PC1) and second (PC2) principal components explain 52.7% of the variance. The analysis showed distinct metabolomic profiles at each time point during the course of infection, culminating with the largest segregation of metabolites between the R and S lines at 72 hpi (Fig. S1). The fold change of significantly regulated metabolites during this time course are indicated in Supplementary Table 5. Significantly regulated metabolites were assigned to distinct functional categories according to the chemical groups to which they belong (Table S5) and to specific plant pathways in which they may function (Table S6). Our data revealed several differentially regulated metabolic processes between the R and S lines (Fig. 4). However, those involved in phenylalanine metabolism are particularly interesting given the differential expression of phenylpropanoid pathway genes identified in transcriptomic analysis (Fig. 5A) and the important role of this pathway in plant defense. Indeed, the precursor of the phenylpropanoid pathway, phenylalanine, and downstream intermediate metabolites, such as benzoic acid, caffeic acid, and ferulic acid, are highly accumulated in the stem of the R line compared to the S line following *S. sclerotiorum* infection (Fig. 5B). Overall, the observed parallel between transcriptomics and metabolomics data with respect to the differential modulation of the phenylpropanoid pathway indicates its potential key participation in resistance to *S. sclerotiorum*.

Similar to transcriptomic data, the comparative metabolite profiles also implicated phytohormones in this interaction. Namely, the fatty acids linolenic acid (a precursor of jasmonic
acid) and cyanoalanine (an indicator of ethylene biosynthesis) (36,37) are both significantly induced at 48 and 72 hpi, in the R line (Fig. S3). Interestingly, the most highly upregulated metabolite in our R line is mucic acid with an ~86-fold higher accumulation (Table S5). Mucic acid, also referred to as galactaric acid, can be produced by the oxidation of d-galacturonic acid, the main component of pectin.

Galactose metabolism and the TCA cycle were the pathways most affected by this analysis, and the metabolites assigned to them were primarily carbohydrates and organic acids, respectively. Interestingly, of the metabolites downregulated in R plants in comparison to S plants, 81.8% (9/11) belonged to one of these two groups (Table S5). Although these metabolites relate to multiple pathways, their downregulation in R plants may be a strategy to reduce *S. sclerotiorum* access to preferable carbon sources. Of the top six potentially affected pathways, three of them (Glyoxylate and dicarboxylate metabolism; Alanine, aspartate and glutamate metabolism; the TCA cycle) contain both fumaric and succinic acid. These organic acids were downregulated at 48 and 72 hpi, respectively, and demonstrate the potentially broad impacts that changes in individual metabolites can have on plant biosynthesis and metabolism.

**Reprogramming of the phenylpropanoid pathway in resistance to *S. sclerotiorum***.

Many secondary metabolites derived from multiple branches of the phenylpropanoid pathway, including lignin, isoflavonoid-phytoalexins, and other phenolic compounds such as benzoic acid, have been proposed as important components of defense responses (38,39). In this study, we found differential expression of transcripts and metabolites related to the phenylpropanoid pathway between the R and S soybean lines. At the transcript level, we observed a downregulation of genes encoding phenylalanine ammonia-lyase (PAL), lignin biosynthetic enzymes, ferulate 5-hydroxylase (F5H), N-hydroxycinnamoyl/ benzoyltransferase (HCT),
Caffeoyl-CoA O-methyltransferase (CCoAOMT), cinnamoyl-CoA reductase (CCR), cinnamyl alcohol dehydrogenase (CAD), chalcone synthase (CHS), flavonol synthase (FLS), and flavonoid 4'-O-methyltransferase (FOMT) in R line compared to S line (Table 2 and Fig. 5A). Coincidentally, genes coding for enzymes involved in anthocyanin and phytoalexin biosynthesis were upregulated in the R line, these include, anthocyanidin synthase (ANS), anthocyanin acyltransferase (AT), dihydroflavonol reductase (DFR), isoflavone 7-O-glucoside-6''-O-malonyltransferase (IFMaT) isoflavone 7-O-methyltransferase (IOMT), and isoflavone reductase (IFR) (Table 2 and Fig. 5A). Transcript levels of select genes within these pathways were validated using RT-qPCR, thus confirming the RNA-Seq results (Fig. 6).

In accordance with the transcriptomics data, we observed a marked accumulation of the metabolites phenylalanine, a phenylpropanoid pathway precursor, and the lignin intermediates ferulic and caffeic acids in the R line compared to the S line (Fig. 5B). Benzoic acid, a salicylic acid precursor and known antimicrobial compound (40) also accumulated at higher levels in the Res line (Fig. 5B). Increased accumulation of ferulic and caffeic acids is likely due to the downregulation of F5H (Glyma.09G186400) and other downstream enzymes within the lignin pathway (Table 2 and Fig. 5A). We reasoned that the contribution of these ferulates to resistance may be due to their antifungal activity, and their ability to inhibit S. sclerotiorum growth was tested in vitro. A significant reduction of fungal growth was observed when S. sclerotiorum was grown on PDA with increasing concentrations of ferulic acid (Fig. S2). While caffeic acid did not significantly affect colony size, it clearly affected S. sclerotiorum growth patterns on PDA with the appearance of abnormal concentric ring growth patterns and premature sclerotia formation (Fig. S2).
While lignification is recognized as a disease resistance mechanism in plants, counterintuitively, our data suggest a reprogramming of the phenylpropanoid pathway away from lignin and towards the accumulation of lignin intermediates, anthocyanins, and phytoalexins in the resistance response against *S. sclerotiorum*. We propose that lignin intermediates, such as caffeic and ferulic acid, and possibly other compounds with antifungal activity are an important component of this response.

**ROS scavenging and antioxidant activities are associated with resistance to *S. sclerotiorum***

Reactive oxygen species (ROS) and ROS scavengers play active roles in redox status regulation in biotic stress (41,42). *S. sclerotiorum*, via oxalic acid, is known to upregulate host ROS levels to induce PCD and achieve pathogenic success (11,15,16). Our transcriptomics analysis shows a differential regulation of genes related to ROS scavenging, such as peroxidases, glutathione S-transferases (GSTs), ascorbate oxidases, and superoxide dismutase (SODs), when comparing the R and S soybean lines (Table S8 and Fig. 7). Three *GmGSTs* (Glyma.06G193400, Glyma.02G024600, Glyma.02G024800), two *GmSODs* (Glyma.12G081300, Glyma.12G178800) and five ascorbate oxidases or like proteins (Glyma.05G082700, Glyma.11G059200, Glyma.17G012300, Glyma.17G180400, Glyma.20G051900, Glyma.05G057400) were significantly upregulated in the R line compared to the S line as early as 24 hpi, suggesting a role in preventing oxidative damage imposed by *S. sclerotiorum* (Table S8 and Fig. 7). Peroxidases were also differentially regulated at 24 hpi, with five family members (Glyma.01G19250, Glyma.11G049600, Glyma.11G080300, Glyma.14G201700, Glyma.17G177800) significantly upregulated, however, many others were downregulated in the R line compared to S line (Table S8 and Fig. 7).
We mined our metabolomics data for differentially accumulated metabolites that may serve as ROS scavengers or antioxidants. Dehydroascorbic acid (DHA), the oxidized form of ascorbate, an important antioxidant, was specifically accumulated later in the infection time course (72 hpi), but not at the early stages in the R line (Fig. 7). Similarly, the proline derivative, trans-4-hydroxy-L-proline, a known osmoprotectant and antioxidant (43,44) is significantly accumulated in the R line at the later stages of the infection process (Fig. 7). Proline plays a major role as an antioxidant, owing to its ROS scavenging capacity (45,46). Overall, these results and the earlier observation of anthocyanin induction, point to a marked activation of ROS scavenging and antioxidant processes in the resistant response to *S. sclerotiorum*, presumably to counter to oxidative state imposed by this pathogen.

**Jasmonic acid signaling contributes to the resistant response to *S. sclerotiorum***.

During plant-pathogen interactions, phytohormones such as salicylic acid (SA), abscisic acid (ABA), ethylene (ET), and Jasmonic acid (JA), have been shown to regulate plant immune responses (47–50). ET and JA have generally been implicated in the activation of defense responses against necrotrophs (51). Our GO enrichment analysis highlighted DEGs between the R and S lines related to JA/ET biosynthesis and responses (Table S9 and Fig. 3). Metabolic profiling also identified phytohormone-related metabolites that were differentially accumulated during the course of infection between our lines, namely, linolenic acid, a precursor of jasmonic acid, and cyanoalanine, an indicator of ethylene biosynthesis (Fig. S3). Accordingly, we conducted a targeted GC-MS analysis to more accurately estimate the dynamic changes in SA, ABA, cinnamic acid (an intermediate of SA and the phenylpropanoid pathway), and JA precursors and derivatives (12-oxophytodienoic acid (OPDA), and (+)-7-iso-jasmonoyl-L-isoleucine (JA-Ile) in a time course experiment comparing the R and S soybean lines. Distinct patterns of phytohormone accumulation...
were identified during the course of infection (Fig. 8). The bioactive form of JA, Jasmonoyl-L-isoileucine (JA-Ile) but not JA, was significantly induced in the R line compared to the S line between 6 and 24 hpi, before decreasing at the later stages of our time course (Fig. 8). Interestingly, JA and JA-Ile levels drastically increased in the S line, albeit at a much later stage of infection (48 – 72 hpi). Thus, this late dramatic surge in JA and JA-Ile in the S line may be perceived as a delayed response to fungal colonization, perhaps after the establishment of infection (Fig. 7). In accordance, OPDA, a precursor of JA, accumulated at significantly higher levels in the S line at 48 – 72 hpi (Fig. 8). This pattern also explains the significant induction of JA biosynthetic transcripts at the later stages of infection observed in the S line (Table S9). Expectedly, SA accumulated to higher levels in the S line throughout the time course (Fig. 8). JA and SA responses are known to be antagonistic in many plant-pathogen interactions (52). In toto, phytohormone estimation in the S and R lines following *S. sclerotiorum* challenge suggests that resistance to this pathogen in soybean coincides with an early induction of JA signaling. The timing of this induction appears to be critical to the outcome of this interaction.

**Antifungal activity and mode of action of stem extract from the resistant soybean line**

Our earlier observation that soybean stems from the R line accumulated metabolites with antifungal activity, such as ferulic and caffeic acids, following *S. sclerotiorum* challenge was intriguing. This resistant response was also associated with the development of a prominent red coloration at the site of infection (Fig. 1). Considering the metabolomics data, we reasoned that total stem extract from the resistant line should also exhibit antifungal activity against *S. sclerotiorum*. To test this hypothesis, we prepared an ethanol extract of stem sections harvested from the R line, 10 days post inoculation, that we termed the red stem extract. Equal amount of
green stem extracts harvested from non-inoculated plants served as control. The final extracts were
diluted in DMSO following ethanol evaporation. *S. sclerotiorum* growth was assayed on potato
dextrose broth (PDB) amended with the red stem extract, green stem extract, or DMSO control for
48 hours. Fungal biomass as determined by mycelial fresh weight was markedly reduced (12-14
fold) in PDB cultures containing the red stem extract compared to PDB amended with the green
stem extract or DMSO (Fig. S4 B, C). These results confirm that the resistant response associated
with our R line clearly involves the accumulation of antifungal compounds that inhibit *S.
sclerotiorum* growth.

We next examined the mechanism by which the red stem extract inhibits fungal growth by
performing chemical genomic profiling in yeast. Chemical genomic profiling is built upon
barcoded strains of *Saccharomyces cerevisiae* mutants representing approximately 4000 gene
deletions. The use of unique barcodes for each mutant makes this approach compatible with high-
throughput screening of drugs using next-generation sequencing (53). Chemical genomic profiling
revealed that deletion mutants of genes involved in phospholipid and sterol biosynthesis were
significantly sensitive to the red stem extract (Fig. 9A). Mutants of *ERG6*, which encodes a protein
in the ergosterol biosynthetic pathway, had the greatest sensitivity to the extract, and this was a
highly significant response (p<1e-7). *ERG2*, which is also involved in ergosterol biosynthesis, was
also significantly sensitive (p<0.01). Mutants of *ARO7*, which encode a gene involved in amino
acid biosynthesis was significantly sensitive. While *ARO7* is not known to be directly involved in
lipid/sterol biosynthesis, it has many genetic interactions with lipid related genes (54). *CHO2* and
*OPI3* mutants were also significantly sensitive. Cho2p and Opi3p are both involved in
phosphatidylcholine biosynthesis. A deletion mutant of *PAH1* was the most significantly resistant
mutant. Pah1p is a phosphatase that regulates phospholipid synthesis. Deletion mutants of *PAH1*
have increased phospholipid, fatty acid and ergosterol ester content (55). Further, in two out of three replicates, the chemical genomic profile of the red stem extract had significant correlation (p<0.05) with the profile of fenpropimorph (56), an ergosterol biosynthesis inhibitor that targets \textit{ERG2} and \textit{ERG24} in yeast. Taken together, these data suggest that the red stem extract may exert toxicity by either disrupting enzymes involved in lipid/sterol biosynthesis, or alternatively physically binding membrane lipids and causing cell leakage.

\textbf{Red stem extract inhibits ergosterol biosynthesis in} \textit{S. sclerotiorum}

To test if the red stem extract causes rapid cell lysis like the antifungal compounds amphotericin B (binds ergosterol) and poacic acid (binds glucan), we performed a cell leakage assay. The red stem extract did not cause significant cell permeability (Fig. 9B). The antifungal drug fluconazole inhibits ergosterol biosynthesis but doesn’t cause rapid cell permeability as it targets the enzyme Erg11p rather than a membrane or cell wall structure. This result supports the hypothesis that compounds within the red stem extract target ergosterol biosynthetic enzymes. We thus tested if treatment of \textit{S. sclerotiorum} with the red stem extract alters ergosterol production. \textit{S. sclerotiorum} was grown in PDB amended with red stem extract, green stem extract, or DMSO, and ergosterol level was calculated by first calculating the total ergosterol-plus-24(28) DHE (dehydroergosterol) content and then subtracting from the total the amount of absorption due to 24(28) DHE only (57). Ergosterol content of \textit{S. sclerotiorum} mycelia grown in red stem extract was significantly reduced compared to control treatments (Fig. 9C). The results confirm that the red stem extract inhibits the growth of \textit{S. sclerotiorum} by targeting ergosterol biosynthesis in the fungus.
DISCUSSION

Resistance to fungal pathogens with a predominately necrotrophic lifestyle, such as *Sclerotinia sclerotiorum*, is not well understood due to the likely complex network of responses to these pathogens or their determinants. Uncovering key components of these defense responses is essential for the deployment of disease resistant crops. Omics approaches offer a unique opportunity to identify global cellular networks in plants in response to these pathogens. *S. sclerotiorum* is a broad host pathogen that infects over 400 species, mostly dicotyledonous plants. Its pathogenic success most assuredly relies on a broadly effective toolkit that allows it to infect multiple hosts, however, host-specific dialogue between a given host and the pathogen may also be of importance. In this study, we specifically examine soybean resistance mechanisms against *S. sclerotiorum* by comparing opposing outcomes in two soybean lines in response to this pathogen using a combination of transcriptomics, metabolomics, and chemical genomics. Several lines of evidence are consistent with the following conclusions: (i) *S. sclerotiorum* challenge induces drastic changes in gene expression and metabolite production in soybean; (ii) Resistance in soybean is associated with an early recognition of the pathogen and a rapid induction of JA signaling; (iii) The redox buffering capacity of the host is essential to counter the oxidative state imposed by *S. sclerotiorum*; (iv) A reprogramming of the phenylpropanoid pathway and upregulation of antifungal metabolites are observed during the resistant response to *S. sclerotiorum*; (v) The antifungal activity associated with resistance targets ergosterol biosynthesis in the pathogen. Overall, comprehensive genetic, biochemical, and transcriptomic analyses allowed us to uncover a novel resistance mechanism connecting the upregulation of antifungal activity to a successful defense response against *S. sclerotiorum* and highlights the importance of early recognition and redox regulation in resistance to this pathogen.
The importance of ROS in plant immunity and other plant processes, including abiotic stress responses, growth and development, is well documented (58). In plant immunity, ROS can function not only as antimicrobials and in plant cell wall reinforcement, but also as signaling molecules activating additional defense responses (59). The implication of host ROS in plant defenses, including the hypersensitive response (HR) and pathogen-associated molecular pattern (PAMP)-triggered immunity following pathogen recognition is well documented (60,61). However, ROS are also produced during compatible interactions, thus facilitating host colonization of certain fungal pathogens (11,14,16,62). *S. sclerotiorum* is known to induce apoptotic-like PCD via oxalic acid, a process that requires ROS upregulation in the host (15). Indeed, we have recently shown that this pathogen hijacks soybean NADPH oxidases to increase ROS levels leading to tissue death and the establishment of disease, and that the silencing of specific host NADPH oxidases confers enhanced resistance to this pathogen (16). This study confirms the importance of ROS in this pathosystem and suggests ROS scavenging and antioxidant activity as viable resistance mechanisms in soybean. Indeed, we noted the significant upregulation of genes related to ROS scavenging such as peroxidases, glutathione S-transferases, ascorbate oxidases, and superoxide dismutase, in association with the successful defense response against *S. sclerotiorum*. Similar trends were reported in other host and non-host plant species in association with *S. sclerotiorum*, with strong upregulation of genes encoding ROS scavenging enzymes and higher antioxidant enzymatic activities coinciding with resistance to this pathogen (63–66). We also noted the accumulation of antioxidant metabolites dehydroascorbic acid (DHA) and trans-4-hydroxy-L-proline. DHA is converted into ascorbic acid (AA), which is known for its redox buffering capacity and ROS detoxification (67,68). The utilization of ascorbic acid as an antioxidant in cells causes its oxidation back to dehydroascorbic acid (69). Thus, the low levels of
DHA in our resistant line at the onset of infection can be explained by higher ascorbic acid levels at this stage. Once oxidized, AA is converted to DHA, which accumulated at the later stages of our time course. Similarly, the proline derivative, trans-4-hydroxy-L-proline, was markedly increased in our resistance response, and is a known osmoprotectant and antioxidant (43,44). The accumulation of proline in plants has been implicated in stress tolerance by maintaining osmotic balance, stabilizing membranes, and modulating ROS levels (43). Overall, our accumulating evidence suggests that the antioxidant capacity of soybean plays a critical role in its ability to resist *S. sclerotiorum*, a pathogen that induces ROS and cell death in the host to achieve pathogenic success.

An important question is what causes this stark difference in ROS buffering capacity between the R and S soybean lines despite common genetic components? We propose that the early recognition of the pathogen in the R line leads to a timely response that includes the activation of the antioxidant machinery within the host. This is corroborated by our phytohormone analysis that shows a rapid induction of the bioactive form of jasmonic acid (JA), Jasmonoyl-L-isoleucine (JA-Ile) as early as 6 hpi in the resistance response, JA-Ile levels decreased at the later stages of infection, presumably once the infection was under control. In contrast, in the susceptible response, JA and JA-Ile levels remained low early, but drastically increased at the later stages of our time course, which can be conceived as a delayed and unsuccessful response to fend off an already established infection. A large body of work on plant defenses described the implication of specific hormone pathways depending on pathogen lifestyle (47–50,52). Plant defenses involving JA typically inhibit fungal necrotrophs (52), and mutants specifically impaired in JA-Ile accumulation show enhanced susceptibility to such pathogens (70–72). Our results are consistent with the model where upon *S. sclerotiorum* challenge, JA is rapidly biosynthesized from linolenic acid and
subsequently catalyzed to JA-Ile, the active form of JA. Our metabolomics analysis also showed that linolenic acid is hyper accumulated early in the resistance response. JA-Ile is known to act upon the F-box protein COI1 (Coronatine-insensitive protein 1) leading to the targeting of JAZ (Jasmonate-Zim Domain) proteins for degradation, thus liberating JAZ repressed transcription factors involved in defense (72,73). Thus, while JA signaling appears to be activated in both resistant and susceptible responses, the timing of this induction is key to the resistance outcome in *S. sclerotiorum*-soybean interaction. The early induction likely leads the timely activation of defense components culminating in the arrest of fungal growth and colonization. The reciprocal antagonism between JA and SA signaling pathways is often discussed, and SA signaling is expected to have a negative effect on resistance to pathogens with a predominately necrotrophic lifestyle (52). Our data indicates a higher and sustained levels of SA in the susceptible line throughout our time course, thus in line with the notion that these signaling pathways are antagonistic. SA is also associated with elevated ROS levels and cell death induction, to the benefit of necrotrophic pathogens (74). However, the exact role of SA in this interaction will require further investigation knowing that the crosstalk between JA and SA signaling is complex, and synergistic interactions have also been reported (50).

JA signaling have been linked to the alteration of plant secondary metabolites, including alkaloids, terpenoids, flavonoids, phenolic compounds, and phytoalexins (75–79). Secondary metabolites derived from multiple branches of the phenylpropanoid pathway, including lignins, isoflavonoid-phytoalexins, and other phenolic compounds have also been proposed as important components of defense responses (38,39). The integration of our transcriptomic and metabolic data revealed that the phenylpropanoid pathway is differentially regulated in our soybean lines in response to *S. sclerotiorum* challenge. Specifically, our data suggest a reprogramming of the
phenylpropanoid pathway with its flux diverted from lignin to lignin intermediates, anthocyanins, and phytoalexins in the resistance response against *S. sclerotiorum*. While these results will require further confirmation by the absolute quantification of phenylpropanoid pathway components, the accumulation of upstream metabolites such as ferulic acid, caffeic acid, and benzoic acid in the resistance line support a reduced flow towards lignins. The observed accumulation of ferulic acid is also consistent with decreased lignification considering its role as a nucleation site for lignin polymerization (80). These results may seem counterintuitive considering that lignin biosynthesis is associated with cell wall fortification as a mechanism of disease resistance (81). However, in support of our results, a negative correlation between soybean stem lignin content and resistance to *S. sclerotiorum* has previously been reported (32). Flux changes within the phenylpropanoid pathway in maize have been discussed in response to the biotrophic fungal pathogen *Ustilago maydis* (82,83). Interestingly, *U. maydis*, via the secreted effector Tin2, diverts the flow of the phenylpropanoid pathway away from lignins by increasing anthocyanin biosynthesis to facilitate infection. In the absence of this effector, lignin biosynthesis is enhanced presumably to limit fungal colonization (83). In contrast to our results, resistance to *U. maydis* appears to be associated with the activation of the lignin branch of the phenylpropanoid pathway. However, these observations are in line with the limited lytic repertoire and the biotrophic lifestyle of *U. maydis*. Against necrotrophic fungal pathogens, anthocyanin may act as antioxidants by scavenging ROS and limiting the induction of cell death required by these pathogens. Indeed, the ROS scavenging capacity of anthocyanins has been proposed to provide protection against necrotrophic pathogens such as *Botrytis cinerea* (84) and *Erwinia carotovora* (85). We propose that this mechanism may also confer resistance against *S. sclerotiorum* in soybean.
The reported antimicrobial activities of ferulic (86), caffeic (87), and their significant accumulation in the resistance response against S. sclerotiorum prompted us to consider antifungal activity as a component of the soybean defense response against this pathogen. Indeed, total red stem extracts from our resistant line following S. sclerotiorum challenge clearly inhibited fungal growth in vitro. This antifungal response was seemingly absent from healthy soybean plants, activated only in response to S. sclerotiorum. Furthermore, we were able to show using chemical genomics in yeast that factors within this antifungal activity target ergosterol biosynthesis in the fungus. Ergosterol, a lipid found in the cellular membranes of fungi, is important to the regulation of membrane fluidity and permeability. It is therefore conceivable that plants have evolved means to target this key component of fungal membranes. For example, saponins, which affect membrane integrity by targeting 3β-hydroxyl sterols, were shown to be required for resistance against Gaeumannomyces graminis var. tritici and several Fusarium spp (88–90). Plant antimicrobial peptides and coumarin, a product of the phenylpropanoid pathway, were also proposed to target fungal ergosterol (91,92). The specific metabolites responsible for these antifungal activities in our resistant line are currently unknown. While we provided evidence that both ferulic and caffeic acids affected S. sclerotiorum growth, their chemical genomic profile did not match that of the red stem extract. Thus, we propose that other unknown compounds target ergosterol biosynthesis and contribute to resistance to this pathogen. The identification of these compounds through high-resolution mass spectrometry may lead to the discovery of novel bioactive metabolites and help devise specific strategies to introgress resistance to fungal pathogens in crop plants.
METHODS

**Plant material and pathogen inoculation**

Two recombinants inbred lines of soybean (*Glycine max*), 91-145 and 91-144, were used in this study. Both the resistant 91-145 (R) and the susceptible 91-44 (S) lines were developed utilizing W04-1002 (P1), a SSR resistant parental line, and LN89-5717 (PI 5745542), a SSR-susceptible parental line demonstrating other desirable pathogen resistance traits (9). Soybean seedlings and plants were maintained in the greenhouse or growth chamber at 24 ± 2°C with 16-h light/8-h dark photoperiod cycle. Plants were supplemented fertilizer (Miracle-Gro) every two weeks.

SSR infection was performed using a wild type strain of *S. sclerotiorum* (1980) grown at room temperature on potato dextrose agar (PDA) as described by Godoy et al. 1990. Four-week-old soybean plants were infected with *S. sclerotiorum* by petiole inoculation, using an agar plug of actively growing fungal hyphae. Plant tissue was sampled by cutting horizontally above and below (1.5 cm) the node of the inoculated petiole with a clean straight-edge razor (16). Tissue samples were then immediately frozen in liquid nitrogen prior to RNA extraction and metabolomic analysis. Samples from non-inoculated stem tissues were also collected as a control. The experimental design was completely randomized and consisted of three biological replicates for each of the treatments. Each biological replicate consisted of stem segments (~3 cm, first internode) from 2 different plants.

**RNA Extraction and library preparation**

Total RNA was extracted from soybean stem tissues using a modified Trizol protocol (Invitrogen Corp., Carlsbad, CA, USA). Briefly, collected tissue from each sample was finely ground in liquid nitrogen. For each 100 mg of tissue, 1 ml of chilled Trizol was added. Samples were centrifuged...
at 12k rpm for 5 min at 4°C. Supernatant was discarded and 200 µl of chilled chloroform was added, and vortexted at high speed for 15 sec. Samples were centrifuged again at 12k rpm for 15 min at 4°C. The aqueous phase was mixed with 0.8x isopropanol and left at room temperature for 10 minutes, followed by centrifugation at 12k rpm at 4°C. Supernatant was discarded, and the pellet was washed with 75% ethanol. Pellet was air dried for 10 minutes at room temperature and resuspended in 20µl of nuclease free water followed by incubation at 55°C for 10 minutes. Samples were cleaned using the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany). RNA concentration and purity were determined by Nanodrop (Thermo Fisher Scientific, Wilmington, DE) and sample quality was assessed using an Agilent Bioanalyzer 2100 and an RNA 6000 Nano Kit (Agilent Technologies, Santa Clara, CA). The RNAseq experiment included three biological replicates per treatment.

Library preparation was performed at the University of Wisconsin – Madison Biotechnology Centre (Madison, WI, USA). Individually indexed libraries were prepared using the TruSeq RNA Sample Preparation v2 kit according to the manufacturer’s instructions (Illumina, San Diego CA, USA). Library concentrations were quantified with the Qubit HS DNA kit (Thermo Fisher Scientific, Wilmington, DE). The size and quality of the libraries were evaluated with an Agilent Bioanalyzer 2100 and an Agilent DNA 1000 kit (Agilent Technologies, Santa Clara, CA) and the libraries were sequenced using Illumina HiSeq2500 (1X100bp) (Illumina, San Diego CA, USA).

**Quality check and sequence analysis**

Illumina raw read data quality was verified with fast QC. The soybean and *S. sclereotiorum* genome sequences were acquired from Phytozome v12.1 (https://phytozome.jgi.doe.gov/pz/portal.html#! bulk?org=Org_Gmax) and the Broad institute
Raw sequence reads were mapped to both genomes using the Subjunc aligner from Subread (94). Alignments were compared to the gene annotation GFF files for both organisms (Soybean: Gmax_275_Wm82.a2.v1.gene.gff3 (93), S. sclerotiorum: sclerotinia_sclerotiorum_2_transcripts.gtf (10)) and raw counts for each gene were generated using the feature Counts tool from subread. The raw counts data were normalized using voom from the R Limma package, then used to generate differential expression (log$_2$FC) values (95,96). DEGs were generated from the comparison of inoculated soybeans of both lines at different time points to their respective uninoculated control (FDR<0.05; log$_2$FC>1 or <-1).

**Gene annotation and gene ontology enrichment analysis**

Differentially expressed genes were annotated using soybean genome gene annotations (Annotation_Gmax_275_Wm82.a2.v1.gene.gff3) from Phytozome (https://phytozome.jgi.doe.gov/pz/portal.html#!bulk?org=Org_Gmax) while soybean chloroplast (https://www.ncbi.nlm.nih.gov/nuccore/91214122) and mitochondrion (https://www.ncbi.nlm.nih.gov/nuccore/476507670) sequences were used from NCBI. The statistically significant DEGs (p-value<0.01; log$_2$FC>1 or <-1) of the R v S comparison were used to identify enriched Gene ontology (GO) terms using the Soybase GO term enrichment tool (https://www.soybase.org/goslimgraphic_v2/dashboard.php) (34).

**Metabolites estimation and analysis**

Four-week-old soybean plants of susceptible and resistant lines were petiole inoculated as described above. Four stems (3 cm) for each treatment per biological replicate were harvested 48 and 72 hours post inoculation and non-inoculated stems were used as a control. Collected stem
samples were immediately frozen in liquid nitrogen and kept at −80°C until used. Gas chromatographic mass spectrometry (GC-MS) analysis was performed by West Coast Metabolomics (Univ. California, Davis). For metabolite estimation, stem tissue from each treatment was finely ground with liquid nitrogen and extracted with 1 ml of chilled 5:2:2 MeOH: CHCl₃: H₂O. The aqueous phase was transferred into a new tube and a 500 µl aliquot was dried under vacuum and the residue was derivatized in a final volume of 100 µl. Detailed methods of metabolite derivatization, separation, and detection are described in Fiehn et al., 2016 (97). Briefly, samples were injected (0.5 µl, split less injection) into a Pegasus IV GC (Leco Corp., St Joseph, MI) equipped with a 30 m x 0.25 mm i.d. fused-silica capillary column bound with 0.25 µm Rtx-5Sil MS stationary phase (Restek Corporation, Bellefonte, PA). The injector temperature was 50°C ramped to 250°C by 12°C s⁻¹. A helium mobile phase was applied at a flow rate of 1 ml min⁻¹. Column temperature was 50°C for 1 min, ramped to 330°C by 20°C min⁻¹ and held constant for 5 min. The column effluent was introduced into the ion source of a Pegasus IV TOF MS (Leco Corp., St Joseph, MI) with transfer line temperature set to 230°C and ion source set to 250°C. Ions were generated with a -70 eV and 1800 V ionization energy. Masses (80-500 m/z) were acquired at a rate of 17 spectra s⁻¹. ChromaTOF 2.32 software (Leco Corp) was used for automatic peak detection and deconvolution using a 3 s peak width. Peaks with signal/noise below 5:1 were rejected. Metabolites were quantified by peak height for the quantification ion. Metabolites were annotated with the BinBase 2.0 algorithm (98).

Statistical analyses were conducted in MetaboAnalyst 3.0 (35). Multivariate and univariate statistics were performed on generalized log transformed peak heights. Metabolites with FDR <0.05 were considered differentially regulated. Metabolite pathway analysis was done using the MetaboAnalyst 3.0 Pathway Analysis tool (35).
Reverse Transcriptase–quantitative PCR (RT-qPCR)

The internodal region of the infected petiole (including symptomatic and non-symptomatic tissue) was used for RNA isolation. Stems (3 cm) were harvested and immediately frozen in liquid nitrogen. RNA was isolated using the above-mentioned protocol and then treated with RNase-free DNaseI (NEB Inc., Ipswich, MA, USA). The RNA was reverse transcribed using the AMV First-Strand cDNA Synthesis Kit (NEB Inc., Ipswitch, MA) and oligo-dT primer according to the manufacturer’s instructions. RT-qPCR was performed using a SensiFAST SYBR No ROX Kit (Bioline USA Inc., Taunton, MA, USA). Each reaction consisted of 5 µL of SensiFAST SYBR No-ROX Mix, 1µL of 1 : 10-fold diluted template cDNAs, and 0.4 µL of 10 µM gene-specific forward and reverse primers in a final volume of 10 µL. Primers were designed using Primer3 software (99,100) for the amplification of gene fragments that were approximately 100 – 200 bp in length and with an annealing temperature of 60°C (Table S7). The primer specificity was checked in silico against the NCBI database through the Primer-BLAST tool (http://www.ncbi.nlm.nih.gov). RT-qPCR was performed on a CFX96 real-time PCR system (Bio-Rad, Hercules, CA). The run conditions were: 2 min of initial denaturation at 95°C; 95°C for 5 s, 58°C for 10 s and 72°C for 20 s (40 cycles). The relative expression of genes was calculated using the $2^{-\Delta\Delta C_t}$ method (101) with the soybean gene GmCon15S (102) as an endogenous control. Three biological replicates were used for each sample.

Targeted GC-MS analysis of CA, SA, JA, JA-Ile, cis-OPDA, and ABA

Four-week-old soybean plants of susceptible and resistant lines were petiole inoculated with actively growing agar plugs of *S. sclerotiorum*. Four stems (3 cm) for each biological replicate were harvested at 6, 12, 24, 48, and 72 hpi. Uninoculated stems were also collected for
the estimation of the basal concentration of the phytohormones. These collected stem samples were immediately frozen with liquid nitrogen and kept at −80°C until used. Stem tissues from each treatment were finely ground with liquid nitrogen and 100 mg of the ground tissue was added to 500 µl of phytohormone extraction buffer (1-propanol/water/HCl [2:1:0.002 vol/vol/vol]) and 10 µl of 5 µM solution of deuterated internal standards: d-ABA ([2H6](+)-cis,trans-ABA; [Olchem]), d-IAA ([2H5] indole-3- acetic acid, Olchem), d-JA (2,4,4-d3; acetyl-2,2-d2 JA; CDN Isotopes), and d-SA (d6-SA, Sigma) and analyzed using GC-MS (103–105). The simultaneous detection of several hormones was accomplished using the methods described in Muller et al 2011 with modifications (106). The analysis utilized an Ascentis Express C-18 Column (3 cm × 2.1 mm, 2.7 µm) connected to an API 3200 LC-electrospray ionization-tandem mass spectrometry (MS/MS) with multiple reaction mentoring (MRM). The injection volume was 2 µl and had a 600 µl/min mobile phase consisting of Solution A (0.05% acetic acid in water) and Solution B (0.05% acetic acid in acetonitrile) with a gradient consisting of (time – %B): 0.3 – 1%, 2 – 45%, 5 – 100%, 8 – 100%, 9 – 1%, 11 – stop. Three biological replicates of each treatment were performed.

Plate inhibition assay of *S. sclerotiorum*

The plate growth inhibition assay of *S. sclerotiorum* was done on solid PDA culture plates containing 0, 250, 500, or 1000 µg/mL of ferulic or caffeic acid. Three replicates were used for each treatment. Plates were inoculated with an actively growing plug of *S. sclerotiorum* and grown at 25 °C for either 48 hours (ferulic acid) or 7 days (caffeic acid) prior to assessment.

Compound extraction from soybean stem
Five hundred mg of infected red stem or unaffected green stem was mixed 1:1 w/v in 100% ethanol at 80°C for 1 h. Samples were resuspended with 100 µl of DMSO and used for *S. sclerotiorum* inhibition assay, chemical genomics, and cell permeability assays.

**Chemical genomic analysis**

Chemical genomic analysis of the red stem extract was performed using the non-essential yeast deletion mutant collection as described previously (53). Briefly, triplicate 200 µL cultures of the pooled deletion collection were exposed to a 1:10 dilution of the red stem extract and allowed to grow for 48 h. Genomic DNA was extracted using the Invitrogen Purelink 96-well genomic extraction kit (Invitrogen, Carlsbad, CA, USA). Mutant specific barcodes were amplified using indexed primers. Samples were sequenced on a HiSeq2500 (1X50bp) (Illumina, San Diego CA, USA) rapid run and reads were processed using BEAN-counter (107) and EdgeR (108).

**Cell permeability assay**

To quantify the membrane damage caused by the red stem extract a FungaLight™ Cell Viability assay (Invitrogen L34952) and Guava Flow Cytometer (Millipore, USA) was used as described previously (109). Amphotericin B (100 µg/mL) and poacic acid (100 µg/mL) were included as positive controls. Fluconazole (1 mg/mL) was also included as a control, given its ability to inhibit ergosterol biosynthesis without causing rapid cell permeability. We exposed 200 µL log phase populations of yeast cells in YPD media to the control drugs, red stem extract (0.01, 0.005, and 0.0025%), the control green stem extract at (0.01, 0.005, and 0.0025%), and a 1% DMSO control.
(n=3) for 4 h at 30°C. The cells were then stained with the FungaLight™ kit and immediately analyzed by flow cytometry.

**Biomass and ergosterol estimation of S. sclerotiorum**

The biomass of *S. sclerotiorum* was measured by growing the fungus in potato dextrose broth (PDB). Freshly grown PDA cultures were scraped and then washed twice with water at 4000 rpm (4°C) before being resuspended in water. Equal amounts of resuspended *S. sclerotiorum* were inoculated into 250 mL conical flasks containing 30 mL PDB. For each 30 ml of PDB, 300µl of red stem extract, green stem extract, or DMSO were added and incubated over a period of 48 hours. To estimate the wet weight, the mycelia were filtered on a pre-weighed Miracloth (Darmstadt, Germany). Ergosterol level was estimated as described by Yarden et. al 2014 (57) and Arthington-Skaggs et al 1999 (110). Briefly, mycelia of the *S. sclerotiorum* treated with either red stem extract, green stem extract, or DMSO as described above were washed with distilled water. One gram of mycelium for each strain was resuspended in 3 ml of 25% alcoholic potassium hydroxide solution and vortexed vigorously. Mycelia were then incubated at 85°C for 1 h and sterols extracted by adding 4 ml of n-heptane solution (25% sterile distilled water and 75% n-heptane). The heptane layer was transferred to a clean glass tube and spectrophotometric readings taken between 230 and 300 nm. Four biological replicates were used for each treatment. Ergosterol content was calculated as a percentage/g of wet weight using the following equations: % ergosterol + % 24(28)DHE = 

\[
\left(\frac{A_{290.5}}{A_{290}}\right) \times F, \quad \% \ 24(28)\text{DHE} = \left(\frac{A_{518}}{A_{230}}\right) \times F,
\]

where F is the factor for dilution in ethanol and 290 and 518 are the E values determined for crystalline ergosterol and 24(28) dehydroergosterol (DHE), respectively.
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Table 1. Summary of the sequencing metrics of the RNA-seq.

| Time points (hours) | Total reads | Mapping to *Glycine Max* | Mapping to *S. sclerotiorum* |
|---------------------|-------------|--------------------------|-----------------------------|
| Control             | 79,559,096  | 76,737,931 (96.5%)       | 0 (0.0%)                    |
| 24 hpi              | 81,259,978  | 74,457,988 (91.6%)       | 3,607,921 (4.4%)            |
| Susceptible line (S)| 48 hpi      | 74,344,340               | 68,265,965 (91.8%)          | 2,941,195 (4.0%)            |
|                     | 96 hpi      | 69,266,099               | 47,765,044 (68.9%)          | 18,712,374 (27.0%)          |
| Control             | 89,753,272  | 85,133,076 (96.6%)       | 0 (0.0%)                    |
| 24 hpi              | 64,923,300  | 60,261,793 (92.8%)       | 2,286,347 (3.5%)            |
| Resistant line (R)  | 48 hpi      | 67,520,185               | 62,099,244 (91.9%)          | 2,513,457 (3.8%)            |
|                     | 96 hpi      | 72,951,207               | 64,399,890 (88.2%)          | 5,691,551 (7.8%)            |
### Table 2. A list of differentially expressed genes (DEGs) within the soybean phenylpropanoid pathway.

| Gene locus     | 24 hpi<sup>a</sup> | 48 hpi<sup>a</sup> | 96 hpi<sup>a</sup> | Description                                      |
|---------------|--------------------|--------------------|--------------------|--------------------------------------------------|
| Glyma.03G181700 | -                  | -                  | -1.08              | Phenylalanine ammonia-lyase                       |
| Glyma.03G181600 | -                  | -0.60              | -1.20              | Phenylalanine ammonia-lyase                       |
| Glyma.02G039300 | -                  | -                  | -1.28              | Phenylalanine ammonia-lyase                       |
| Glyma.09G186400 | -                  | -                  | -1.05              | Ferulate 5-hydroxylase                           |
| Glyma.04G040000 | -                  | -                  | -2.08              | N-hydroxycinnamoyl transferase                    |
| Glyma.01G004200 | -                  | -                  | -1.32              | Caffeoyl-CoA O-methyltransferase                  |
| Glyma.05G147000 | -                  | -                  | -1.54              | Caffeoyl-CoA O-methyltransferase                  |
| Glyma.19G007200 | -                  | -                  | -1.12              | Cinnamoyl-CoA reductase                           |
| Glyma.19G006900 | -                  | -                  | -1.13              | Cinnamoyl-CoA reductase                           |
| Glyma.02G130400 | -                  | -                  | -1.05              | Chalcone synthase                                |
| Glyma.09G075200 | -                  | -                  | -1.13              | Chalcone synthase                                |
| Glyma.01G091400 | -                  | -                  | -1.23              | Chalcone synthase                                |
| Glyma.08G110500 | -1.26              | -1.48              | -2.05              | Chalcone synthase                                |
| Glyma.08G110700 | -1.25              | -1.27              | -2.06              | Chalcone synthase                                |
| Glyma.08G109200 | -1.3               | -1.58              | -2.13              | Chalcone synthase                                |
| Glyma.08G110400 | -1.38              | -1.64              | -2.18              | Chalcone synthase                                |
| Glyma.08G110900 | -1.12              | -1.49              | -2.18              | Chalcone synthase                                |
| Glyma.08G110300 | -1.13              | -1.5               | -2.21              | Chalcone synthase                                |
| Glyma.16G219400 | -                  | -                  | -2.23              | NAD(P)H-dependent 6'-deoxychalcone synthase      |
| Glyma.08G109300 | -1.17              | -1.53              | -2.24              | Chalcone synthase                                |
| Glyma.08G109400 | -1.3               | -1.65              | -2.42              | Chalcone synthase                                |
| Glyma.02G048700 | -                  | -                  | 1.49               | Chalcone-flavanone isomerase                      |
| Glyma.07G150900 | -                  | -                  | -2.50              | Flavonol synthase                                |
| Glyma.18G201900 | -                  | -                  | -3.77              | Flavonol synthase                                |
| Glyma.18G267800 | -                  | -5.27              | -9.45              | Flavonoid 4'-O-methyltransferase                  |
| Glyma.11G164700 | -                  | -                  | 2.99               | Dihydroflavonol reductase                         |
| Glyma.12G238200 | -0.88              | -0.98              | -2.07              | Dihydroflavonol reductase                         |
| Glyma.11G027700 | -                  | -                  | 3.27               | Anthocyanidin synthase                            |
| Glyma.01G214200 | -                  | -                  | 2.95               | Anthocyanidin synthase                            |
| Glyma.02G226000 | -                  | -                  | 1.46               | Anthocyanidin 3-O-glucosyltransferase             |
| Glyma.18G271600 | 3.31               | -                  | -                  | Anthocyanin acyltransferase                       |
| Glyma.13G302300 | -                  | -                  | 2.49               | Anthocyanin acyltransferase                       |
| Glyma.13G302500 | -1.10              | -                  | -                  | Anthocyanin 5-aromatic acyltransferase           |
| Glyma.14G034100 | -5.85              | -3.69              | -2.93              | Anthocyanin 5-aromatic acyltransferase           |
| Glyma.19G030800 | -                  | -                  | 4.58               | Malonyl-CoA:isoflavone 7-O-glucoside-6''-O-       |
| Glyma.11G256500 | -                  | -                  | 4.61               | Malonyltransferase                               |
| Accession   | Log₂ FC | Fold Change | Enzyme Name                           |
|------------|---------|-------------|---------------------------------------|
| Glyma.09G094400 | -1.01   | -1.01       | Isoflavone-7-O-methyltransferase      |
| Glyma.06G286600 | 2.69    | 2.69        | Isoflavone 7-O-methyltransferase      |
| Glyma.13G173300 | -1.80   | -1.80       | Isoflavone 4'-O-methyltransferase     |
| Glyma.13G173600 | -1.83   | -1.83       | Isoflavone 4'-O-methyltransferase     |

^a Fold changes (Log₂ FC) relative to S line.
Figure Legends

Figure 1. **Symptom development in susceptible and resistant soybeans.** Disease symptoms observed following petiole inoculation with an agar plug containing actively growing mycelia of *S. sclerotiorum* at 24, 48, 72, 96 hours post-inoculation (hpi) and 7 days post inoculation (dpi). (A) Susceptible (S) line. (B) Resistant (R) line. At 7 dpi in the R line, red coloration at point of inoculation (red node) is prominently visible.

Figure 2. **Differentially expressed genes (DEGs) identification in R and S line after *S. sclerotiorum* infection.** Venn diagram showing (A) DEGs in R line at 24, 48 and 96 hpi compared to control (non-inoculated) sample, (B) DEGs in S line at 24, 48 and 96 hpi compared to control (non-inoculated) sample, (C) In total, 921 and 8223 DEGs were unique to the R and S line, respectively, while 7319 were identified in both the lines, (D) DEGs in R line compared to S line at 24, 48 and 96 hpi.

Figure 3. **Significantly enriched Gene ontology (GO) biological processes in R line compared to S line at different time-points following infection with *S. Sclerotiorum*.** The y-axis represents significantly enriched GO processes which were enriched (FDR <0.05) at at-least one time point of infection (24, 48 and 96 hpi). The x-axis indicates the total number of genes annotated to each GO process. Orange sections represent downregulated genes while blue sections represent upregulated genes.

Figure 4. **Pathway analysis of metabolites differentially detected between the R and S lines at 48 and/or 72 hpi.** The y-axis represents the annotated metabolites in each pathway. Blue sections represent the differentially detected metabolites, whereas green sections represent the remaining annotated metabolites in the pathway. Percentages denote the portion of the pathway found to be differentially detected. Only pathways with at least 10% of their annotated metabolites showing significant regulation were included.
Figure 5. DEGs and metabolites involved in the phenylpropanoid pathway in R and S soybean lines following S. sclerotiorum infection. (A) Enzymes are indicated in uppercase letters. Gene names indicated in red or green represent significantly upregulated or downregulated genes in the R line compared to the S line, respectively. The metabolite name in red indicates significantly upregulated phenylpropanoid pathway intermediates in the R line compared to the S line. PAL (phenylalanine ammonia-lyase); BA2H (benzoic acid 2-hydroxylase); C4H (cinnamate 4-hydroxylase); COMT (caffeic O-methyltransferase); F5H (ferulic acid 5-hydroxylase); 4CL (4-coumarate: CoA ligase); C3H (p-coumarate 3 hydroxylase); HCT (N-hydroxycinnamoyl transferase); CCoAOMT (Caffeoyl-CoA O-methyltransferase); F5H (flavanone 5-hydroxylase); CCR (Cinnamoyl-CoA reductase); CAD (Cinnamoyl alcohol dehydrogenase); CHS (chalcone synthase); CHI (chalcone isomerase); CHR (chalcone reductase); FOMT (Flavonoid 4'-O-methyltransferase); FLS (Flavonol synthase); F3H (flavanone 3-hydroxylase); DHFR (dihydroflavonol-4-reductase); ANS (anthocyanidin synthase also called LDOX, leucoanthocyanidin dioxygenase); ANS (Anthocyanidin synthase); UFGT (UDP-flavonoid glucosyltransferase); AT (Anthocyanin acyltransferase); IFS (Isoflavone synthase); IFMaT (Isoflavone 7-O-glucoside-6''-O-malonyltransferase); IOMT (Isoflavone methyltransferase); this figure was adapted from Baxter and Stewart, 2013 (111) and Ferrer et al 2008 (111) (B) Estimation of the phenylpropanoid pathway intermediate metabolites (peak height) phenylalanine, benzoic acid, 3, 4 - dihydroxycinnamic acid (caffeic acid) and ferulic acid in R and S line at 48 and 72 hpi using GC – MS. Data are presented as means ± standard deviation (SD) from three independent biologically replicate with each replicate containing four pooled stem samples. * Indicates a significant difference at p-value < 0.05 (T-Test).

Figure 6. Confirmation of expression profiles of select phenylpropanoid pathway genes using qRT-PCR. (A) PAL (Phenylalanine ammonia-lyase, Glyma.02G309300); (B) HCT (N-hydroxycinnamoyl transferase, Glyma.04G040000); (C) CCoAOMT (Caffeoyl-CoA O-methyltransferase, Glyma.05G147000); (D) CCR (Cinnamoyl-CoA reductase, Glyma.19G006900); (E) CHS (Chalcone synthase, Glyma.08G110400); (F) DHFR (Dihydroflavonol reductase, Glyma.12G238200); (G) ANS (Anthocyanidin synthase, Glyma.11G027700); (H) IOMT (Isoflavone 7-O-methyltransferase, Glyma.11G256500) and (I) AT (Anthocyanin acyltransferase, Glyma.18G271600). The fold changes in expression values for
qRT-PCR were calculated by comparing the expression values of genes in inoculated vs. non-infected soybean stem tissues using the $2^{-\Delta\Delta Ct}$ method. GmCons15 was used as endogenous control. The absolute fold changes were converted to Log$_2$FC. Data are presented as means ± standard deviation (SD) from three independent experiments.

**Figure 7. Reactive oxygen species (ROS) scavenging machinery.** (A) Heat map of ROS scavenging (Glutathione S-transferase, Peroxidase, L-ascorbate oxidase, and Superoxide dismutase) and antioxidant genes (Proline-rich protein) induced in the R line compared to the S line during S. sclerotiorum infection at 24, 48 and 96 hpi. (B) Differential accumulation of ROS related metabolites trans-4-hydroxy-L-proline and dehydroascorbic acid during S. sclerotiorum infection at 48 and 72 hpi. Data are presented as means ± standard deviation (SD) from three independent experiment. * Indicates a significantly difference at p-value < 0.05 (t-test).

**Figure 8. Estimation of phytohormones and a phenylpropanoid pathway precursor (Cinnamic acid, CA) in R and S lines following S. sclerotiorum infection.** (A) Cinnamic Acid (CA), (B) Salicylic Acid (SA), (C) Abscisic Acid (ABA), (D) Jasmonic Acid (JA), (E) JA precursor 12-oxophytodienoic acid (OPDA) and (F) bioactive JA derivative (+)-7-iso-jasmonoyl-L-isoleucine (JA-Ile). The bars represent the standard deviation (n = 3). * Indicates a significant difference between the R and S lines at p-value < 0.05 (One-way ANOVA).

**Figure 9. Chemical genomic profiling of the red stem extract in yeast mutants and ergosterol biosynthesis.** (A) Plot showing growth differences between yeast deletion mutants exposed to the red stem extract. Dots represent mutants which were significantly sensitive (blue) or resistant (yellow) to the fungicidal activity of the extract. Underlined mutants are implicated in phospholipid and sterol biosynthesis (B) cell leakage assay to test the permeability of S. sclerotiorum treated with DMSO, fluconazole (FCZ), amphotericin B (AMB), poacic acid, and different concentrations of green (Cont) and red (Red) stem extract, (C) ergosterol estimation from S. sclerotiorum treated with DMSO, green stem extract and red stem extract. Vertical bars show standard deviation of means of three replicates. * Indicates a significant difference at p-value < 0.05 (t-test).
Figure 10. Cellular model summarizing *S. sclerotiorum* resistance mechanisms in soybean.

Black line = Induction/suppression of a process, Red line = Secretion/release, Blue line = Bioconversion, Dashed orange line = Translocation of a metabolite, purple asterisk (*) = Downregulation in resistant response, Green asterisk (*) = Upregulation in resistant response, JA = Jasmonic Acid, JA-Ile = Jasmonic Acid-Isoleucine.

**Supplementary Figure Legends**

Supplementary Figure 1. Partial least squares-discriminate analysis (PLS-DA) score plots of metabolic profiles in soybean R and S line. The first (PC1) and second (PC2) principal components explain 52.7% of the variance. Control samples are R0 and S0. *S. sclerotiorum* infected samples at 24, 48 and 96 hpi for the R lines are represented as R24, R48, and R96, respectively. *S. sclerotiorum* infected samples at 24, 48 and 96 hpi for the S line are represented as S24, S48, and S96, respectively. Numbers 1, 2, and 3 represents three independent biological replicates.

Supplementary Figure 2. Effect of Ferulic and Caffeic acids on *S. sclerotiorum* growth. Ferulic acid (A) inhibits the growth of *S. sclerotiorum*. Representative photographs were taken 48 hours post inoculation. Caffeic acid (B) effects normal development of *S. sclerotiorum*. Representative photographs were taken 5 days post inoculation. DMSO (Dimethyl sulfoxide) is the solvent control.

Supplementary Figure 3. (A) Increased accumulation of the jasmonic acid precursor linolenic acid in the R line compared to the S line, (B) Increased accumulation of cyanoalanine (an indicator of ethylene biosynthesis) in the R line compared to the S line. The bars represent the standard deviation (n = 3). * Indicates a significantly difference at p-value < 0.05 (t-test).

Supplementary Figure 4. (A) Red and green stem extract of a R line plant infected with *S. sclerotiorum* and a S line plant mock inoculated. Extraction was performed 10 dpi, (B) Fungal biomass after growth in PDB cultures containing the red stem extract, DMSO, or green stem
extract, (C) weight of fungal biomass in PDB cultures containing the red stem extract, green stem extract, or DMSO.

**List of supplementary tables**

Supplementary Table 1. Differentially expressed genes (DEGs) in R and S lines following *S. sclerotiorum* infection at 24, 48, and 96 hpi compared to control. (A-F) Comparisons between each time-point for both lines and their respective controls.

Supplementary Table 2. Differentially expressed genes in the R line compared to the S line following *S. sclerotiorum* infection at 24, 48, and 96 hpi. (A-C) Time-point comparisons between R and S.

Supplementary Table 3. GO enrichment of significant biological processes generated from differentially regulated genes in the R line compared to the S line. (A-C) GO processes identified through the comparison of the R and S lines at 24, 48, and 96 hpi, (D) Summary of significantly regulated genes belonging to each process at each time-point. Processes were included if they had a FDR<0.05 in at-least one timepoint.

Supplementary Table 4. Estimated gas chromatography – mass spectrometry (GC-MS) peak intensity list of all the metabolites.

Supplementary Table 5. Significantly regulated metabolites in the R line compared to the S line following *S. sclerotiorum* infection at 24, 48, and 96 hpi. Blue = upregulated. Red = downregulated.

Supplementary Table 6. Metabolic pathways assigned to significantly regulated metabolites from comparison of R and S lines at 48 and 72 hpi. (A) Percentage of all annotated metabolites within each pathway which were found to be significantly regulated in this study. Upregulated = Green. Downregulated = Red, (B) Fold changes of the individual metabolites assigned to each pathway.
Values >1 demonstrate upregulation. Value <1 demonstrate downregulation. Values =1 demonstrate no change.

Supplementary Table 7. Primer list for qRT-PCR of phenylpropanoid genes

Supplementary Table 8. Differentially expressed genes encoding putative reactive oxygen species (ROS) scavenging and antioxidant genes in the R line compared to the S line following S. sclerotiorum infection at 24, 48, and 96 hpi.

Supplementary Table 9. Differentially expressed genes encoding putative jasmonic acid (JA) and ethylene (ET) biosynthetic and response genes in the R line compared to the S line following S. sclerotiorum infection at 24, 48, and 96 hpi.
