Abstract

Fusarium virguliforme causes the serious disease sudden death syndrome (SDS) in soybean. Host resistance to this pathogen is partial and is encoded by a large number of quantitative trait loci, each conditioning small effects. Breeding SDS resistance is therefore challenging and identification of single-gene encoded novel resistance mechanisms is becoming a priority to fight this devastating fungal pathogen. In this transcriptomic study we identified a few putative soybean defense genes, expression of which is suppressed during F. virguliforme infection. The F. virguliforme infection-suppressed genes were broadly classified into four major classes. The steady state transcript levels of many of these genes were suppressed to undetectable levels immediately following F. virguliforme infection. One of these classes contains two novel genes encoding ankyrin repeat-containing proteins. Expression of one of these genes, GmARP1, during F. virguliforme infection enhances SDS resistance among the transgenic soybean plants. Our data suggest that GmARP1 is a novel defense gene and the pathogen presumably suppress its expression to establish compatible interaction.

Introduction

Soybean [Glycine max (L.) Merr.] is an economically important crop. Sudden death syndrome (SDS) is one of the most serious soybean diseases and a major cause of soybean yield losses in...
the United States as well as in South American countries [1–3]. In North America, it is caused by the soil-borne fungus, *Fusarium virguliforme* O'Donnell and T. Aoki (formerly *F. solani* (Mart.) Sacc. f. sp. *glycines*); whereas in South America, it is caused by four *Fusarium* spp., *F. virguliforme*, *F. tucumaniae*, *F. brasiliense*, and *F. cuneirostrum* [4,5]. Of the four *F. spp.*, *F. tucumaniae* is the major causal agent of SDS in South America [4]. *F. virguliforme* is asexually propagated, whereas *F. tucumaniae* is sexually propagated. Recently, it has been shown that *F. tucumaniae* carries two idiomorphs at the MAT locus, whereas *F. virguliforme* carries only one [6].

*F. virguliforme* is a hemi-biotrophic fungus that remains in soil. It attacks roots and produces root rot symptoms [7,8]. The pathogen has never been detected in the aboveground diseased plants. In infected roots, it produces fungal toxins including FvTox1 that cause foliar SDS [9–13]. Additional candidate toxins have been detected in xylem sap of *F. virguliforme*-infected soybean plants [14].

Gene expression profiling using RNA sequencing has facilitated understanding the molecular basis of plant-pathogen interactions. Such studies have revealed interesting novel genes and pathways modulated following pathogen infections, and their myriad of responses to overcome pathogen attacks [15–18].

Transcriptome profiles of soybean and other crops infected with pathogens have brought new insights in our understanding of host-pathogen interactions. For instance, Moy and colleagues [19] reported that defense and pathogenesis-related protein genes were strongly induced while lipoxygenases and peroxidases genes were strongly repressed during infection of soybean with the oomycete pathogen *Phytophthora sojae*. Following inoculation of soybean with *F. virguliforme*, many defense-related genes were up-regulated in the partially resistant soybean recombinant inbred line 23 (RIL23), whereas these genes were either unchanged or down-regulated in the SDS susceptible cultivar, ‘Essex’ [20]. Defense-related genes have been shown to be induced in both resistant and susceptible soybean cultivars following *F. virguliforme* infection [21].

In the US, although SDS was first detected in Arkansas only in 1971, it has now spread throughout the soybean growing areas of the North Central United States and Canada [22–25] and is becoming a serious threat to soybean production. The disease has been reported to cause soybean yield losses valued over 100 million dollars [2]. Options for managing SDS are limited. Use of resistant cultivars has been the most effective method of managing this disease. Unfortunately, SDS resistance is partial and governed by a large number of QTL, each contributing a small effect [26–29]. To date, more than 40 QTL for SDS resistance have been reported [30]. As a result, development of SDS resistant soybean lines by combining a large number of QTL by hybridization is not trivial; and therefore, identification of novel single major genes conferring SDS resistance is becoming essential. Unfortunately, it is very unlikely that there are any natural major genes in managing this emerging disease problem. Therefore, development of transgenic soybean lines with manipulated expression of candidate or known defense genes is becoming very urgent for controlling SDS. Earlier it has been demonstrated that transgenic approaches can effectively reduce the yield losses caused by pathogens [31–34].

We hypothesize that pathogens suppress defense-related genes to overcome potent host defense mechanisms to establish in host cells, multiply and spread. To our knowledge, no attempt has been made to alter the expression of down-regulated putative host defense genes to enhance disease resistance in transgenic plants. This study was undertaken primarily to uncover candidate defense-related genes repressed during *F. virguliforme* infection and to determine if altered expression of such a gene can enhance resistance against *F. virguliforme*. We examined the expression profile of soybean genes in roots of young etiolated seedlings infected with *F. virguliforme* conidial spore suspensions or treated with sterile water. We
observed that following inoculation with *F. virguliforme*, transcripts of more genes were up-regulated than down-regulated. We altered the expression of one member of a family of two down-regulated genes, GmARP1 and GmARP2, encoding ankyrin repeat-containing proteins during *F. virguliforme* infection in transgenic soybean plants. Several independent transgenic soybean plants showing induced expression of GmARP1 exhibited enhanced SDS resistance. Our study suggests that (i) *F. virguliforme* somehow suppresses defense-related genes to cause susceptibility and (ii) GmARP1 encoding ankyrin repeats containing protein is a defense gene.

**Materials and Methods**

**Plant materials, treatments, and growth conditions**

For RNA-sequencing (RNA-seq) experiment, soybean seeds of cultivar 'Williams 82' were sown in vermiculite and grown under the dark for 10 days according to Bhattacharyya and Ward [35]. Etiolated seedlings were inoculated with either water (water treatment) or *F. virguliforme* conidial spores (infection) at a concentration of 10^7 spores ml^−1. Root samples were harvested at different time-points, 3, 5, 10 and 24 days following inoculation with *F. virguliforme* or treatment with sterile water [10]. We grouped the samples in four categories: S1, a pooled sample of equal amounts RNAs isolated from roots collected 3 and 5 days following water treatment; S2, a pooled sample of equal amounts RNAs isolated from roots collected 10 and 24 days following water treatment; S3, a pooled sample of equal amounts RNAs isolated from roots collected 3 and 5 days following inoculation with *F. virguliforme*; S4, a pooled sample of equal amounts RNAs isolated from roots collected 10 and 24 days following inoculation with *F. virguliforme*. For three independent RT-PCR experiments, RNA samples were prepared from roots harvested 8 h, 12 h, 1 d, 3 d, and 5 d following either treatment with water or inoculation with *F. virguliforme*.

**DNA isolation, plasmid vector construction, and soybean transformation**

Three root specific and infection inducible promoters (Prom) and the GmARP1 gene (S1 and S2 Figs) were amplified from soybean cv. Williams 82 DNA. Prom 1 (*Glyma18g47390*) was discovered in our lab (B.B. Sahu and M.K. Bhattacharyya, unpublished); Prom 2 (*Glyma10g31210*) and Prom 3 (*Glyma20g36300*) are two root specific promoters, reported earlier (http://www.oardc.ohio-state.edu/SURE/GmROOT/GmRoot.htm). Genomic DNA was isolated using a modified CTAB extraction method [36] adapted from Doyle and Doyle [37]. Promoter sequences were amplified using the following pairs of primers: Prom1F-Prom1R; Prom2F-Prom2R; Prom3F-Prom3R (S1 Table). The sequence of GmARP1 was amplified using the primers GmARP1G-F and GmARP1G-R. In these primers, sequences in bold font indicate cloning sites. The binary vector pTF102 [38] (S3 Fig) was used to create three GmARP1 transgenes: Prom1-GmARP1, Prom2-GmARP1, and Prom3-GmARP1 as follows. First, the CaMV 35S promoter was removed from pTF102 by digesting with XbaI and replaced it with any of the three new promoters. The restriction site for cloning GmARP1 (BstXI) was inserted at the 3’-end of the promoter primers. Next, we excised the GUS gene and CaMV 35S terminator (containing the Poly(A+) signal) by digesting with BstXI and the HindIII. The CaMV 35S terminator was reinserted with addition of the BstXI restriction site at the 5’-end-specific PCR primer, and cloned in the BstXI and HindIII sites. Finally, the created vectors were digested with BstXI and the GmARP1 sequence including 84 nucleotides upstream of the ATG start codon and 45 nucleotides beyond the TAA stop codon was inserted. The constructs were cloned in to *Escherichia coli* strain DH10B and sequenced to confirm their identity. The constructs were then transferred by electroporation into *Agrobacterium tumefaciens* strain EH101 for transformation of Williams 82 at the Plant Transformation Facility, Iowa State University. R_0 transgenic
soybean plants carrying GmARP1 transgenes were maintained in a greenhouse. R1 seeds were harvested for further characterization in growth chambers and then under field conditions.

Infection assays of transgenic soybean plants

Evaluation of transgenic plants in growth chambers. For inoculation of transgenic soybean plants, F. virguliforme Mont-1 was grown on 1/3 potato dextrose agar (PDA) plates for three to five weeks. We prepared the inocula on sorghum grains and mixed with a 1:1 mixture of sand and soil in a 1:20 inoculum:soil ratio for sowing soybean seeds [39]. To assess the responses of the R1 progenies to F. virguliforme infection, we conducted three independent inoculation experiments as follows. We evaluated responses of 15 to 30 R1 progenies to F. virguliforme infection by sowing 3 seeds in a 237-ml Styrofoam cup containing the inocula mixed soil and sand mixture. The cups were then placed in growth chamber maintained at 22–23°C and 16 h light and 8 h dark. The light intensity was 350 μE/m²/s. The plants were watered daily.

Foliar symptoms were scored 4 weeks following planting in a 1 to 7 scale, modified from previously published protocols [40–42]. Plants were considered resistant if they showed symptoms of scores 1 and 2 with symptoms of slight yellowing. The plants were classified as susceptible when disease scores were 3 to 7 characterized by severe chlorosis to necrosis. For molecular analysis, roots of infected plants were harvested and frozen in liquid nitrogen. Chlrophyll contents in leaves of infected plants were used as a measure of foliar symptoms. Extraction and estimation of chlorophyll contents were conducted according to [43]. Extent of root rots was visually evaluated and root resistance to the pathogen was calculated in percentage of healthy roots with no obvious blackening caused by necrosis and rotting.

Field evaluation of transgenic plants with SDS pathogen. A field test of transgenic soybean plants was carried out in the Hinds Research Farm, Iowa State University located in north of Ames, Iowa between June 11 and October 30, 2015. Each transgenic line carrying GmARP1 transgenes were grown in two replications along with the SDS resistant cultivar, MN1606, and the SDS susceptible transgene recipient line, Williams 82. Seeds of individual genotypes were mixed with F. virguliforme NE305S inoculum grown on sorghum grains during planting with a push planter. At 1 to 2-trifoliate stage, all transgenic lines were sprayed with basta herbicide (glufosinate at a 250 mg/L concentration) mixed with 0.1% Tween 20 twice with an interval of two days (S2 Table). DNA samples were harvested from twelve plants that showed resistance to the herbicide. The plants were heavily irrigated in the last week of August that followed by heavy rains. SDS symptoms appeared following heavy rainfall and flood. Individual plants were scored on September 11th, 22nd, 30th, and October 7th, based on a scale of 1 to 9, with 1 being symptomless to 9 for severe symptoms with death of soybean plants (S3 Table) (www.siu.edu/~soybean).

RNA extraction, RNA sequencing, sequence assembly and alignment of reads to Glycine max reference genome

Total RNA samples were extracted from root tissues using the SV Total RNA isolation system (Promega, Madison, WI, USA) following the protocol provided by the manufacturer. The amount and the quality of RNAs in each sample were determined using a spectrophotometer and running on formaldehyde agarose gels, respectively. RNA sequencing was conducted on an Illumina HiSeq 2500 instrument at the DNA Facility, Iowa State University. The sequences were first processed for quality check using FASTX tool-kit. They were then indexed on the soybean reference genome using the open source Bowtie 2 tool [44]. The processed files were aligned to corresponding predicted high confidence coding sequences of the Glycine max
reference genome to calculate RPKM values using Bowtie program and generated SAM (Sequence Alignment/Map) output files for each condition using unix script command [45]. For GO annotation, sequences of differentially expressed genes (DEG) were extracted from Soyabase.org through scripts and Phytozome [46]. The assigned biological function to the DEG was categorized further based on their molecular functions, biological processes and cellular component.

Semi-quantitative RT-PCR amplification

cDNA synthesis was conducted using the M-MLV reverse transcriptase following the instructions of the manufacturer from two μg of total RNAs in each sample (Promega, Inc., Madison, WI, USA). Approximately 200 to 500 bp cDNA fragments were amplified by PCR using gene specific primers for five soybean genes. PCR was conducted for 25 cycles using the following condition: Step 1, 94°C for 2 min; Step 2 94°C for 30 sec; Step 3, at annealing temperature of 60°C for 30 sec; Step 4, extension for 1 min at 72°C; Step 5, repeated cycles 2 through 4 for 24 more times; Step 6, final extension step of 10 min at 72°C. For the three independent RT-PCR experiments of Fusarium-infected and water-treated roots, gene specific primers of each of the four selected genes were used to determine their transcript levels in infected and non-infected roots (S4 Table). Expression of soybean levels was quantified by analyzing the scanned gels carrying electrophoresed RT-PCR products with the ImageJ program (http://imagej.nih.gov/ij/) [47].

For expression analysis of GmARP1 transgenes among transgenic plants, GmARP1-specific forward (GmARP1-RT-F) and reverse primer specific to the poly(A+) signal of transgenes (RT-pTF102-R) were used to determine the expression levels of the GmARP1 transgenes (S1 Table).

Transgene copy number analysis by qPCR

Genomic DNA was extracted from young leaves of 12 transgenic plants for each line. We used approximately 50 mg of lyophilized leaf tissues for DNA extraction at the Iowa State University DNA Facility using the fully automated system, Autogen Autogenprep 740 DNA extraction robot (AutoGen, MA, USA). DNA quantity in each sample was determined using a nanodrop spectrophotometer, and diluted to 20 ng per μl for qPCR reaction.

qPCR was conducted on a Biomark HD system using the 192.24 Taqman CNV protocol (Fluidigm, South San Francisco, CA, USA). Two Taqman assays were designed, the bar gene (target) and the reference gene (an endogenous single copy gene, Glyma.05G014200). Reporter/quencher dyes were FAM/MGB-NFQ for bar and VIC/TAMRA for the reference gene. Data were analyzed using a Biomark HD data collection software and the copy number for the bar gene was calculated.

Results

Identification of differentially expressed soybean genes following F. virguliforme infection

Ten day-old seedlings of cultivar Williams 82 were either treated with water (water treatment) or infected with F. virguliforme isolates. In order to monitor the expression of genes during infection, roots tissues were harvested at different time periods: (i) S1, early time period (ETP) of pooled root samples, 3 and 5 days following water treatment; (ii) S2, late time period (LTP) of pooled root samples, 10 and 25 days following water treatment; (iii) S3, ETP of pooled root samples, 3 and 5 days following F. virguliforme infection; (iv) S4, LTP of pooled root samples,
10 and 24 days following *F. virguliforme* infection. Total RNA samples were extracted from the root tissues and sequenced using Illumina HiSeq 2500 (Illumina, San Diego, CA) and deposited in GEO (accession GSE86201).

The deep-transcript sequencing experiment was conducted only once. We therefore, considered the genes showing at least 10-fold or more changes in transcript levels between infected and control tissues as the differentially expressed genes (DEGs). Furthermore, we considered only those genes as DEGs that have shown to contain at least five sequence reads or fragments per kilobase pair exon sequences in at least one of the treatments considered for comparison. RPKM (reads per kilobase of exon model per million mapped reads) values for individual genes were calculated to normalize the expression levels of individual genes and were used in calculating the fold changes. To validate the transcriptomic data, we conducted three independent biological replications of an RT-PCR experiment for four soybean genes that were repressed following *F. virguliforme* infection.

Pairwise comparison of the expression levels of soybean genes in inoculated roots with those of corresponding water treated roots during ETP- or LTP revealed 314 DEGs that showed \( \geq 10 \)-fold change (FC). In identifying DEGs, we considered only those genes that showed to contain at least 5 sequence reads per kilobase pair exon sequences in at least one of the treatments considered for comparison. We detected transcripts for 54,305 of the predicted soybean genes [48,49]. We found more DIGs in roots of ETP than that in LTP (Fig 1A). In infected roots of both ETP and LTP, there were more up-regulated genes than the down-regulated ones (Fig 1). During ETP, 289 genes were differentially expressed between infected and water-treated roots with FC \( \geq 10 \). The majority of these DIGs (238; 82%) were induced; only 54 genes (18%) were repressed in the infected roots of ETP as compared to the water treated root tissues (Fig 1B and 1C; Table 1; S1 and S2 Datasets). In the infected root tissues of LTP, of the 77 DIGs, 54 (70%) were up-regulated and 23 (30%) were down-regulated (Fig 1B and 1C; Table 2, S3 Dataset).

**Functional classification of soybean genes induced in roots following *F. virguliforme* infection**

We used public transcriptomic databases such as SoyBase [50](http://soybase.org/) and Phytozome (phytozome.jgi.doe.gov), PFAM, and National Center for Biotechnology Information...
Table 1. Genes down-regulated (with FC >10) in soybean roots during early time period following infection with *F. virguliforme*. The full list is reported in S2 Dataset.

| Locus ID   | RPKM S1 | RPKM S3 | Fold change | P-value     | Functional Annotation                                      |
|------------|---------|---------|-------------|-------------|-----------------------------------------------------------|
| Glyma.01g171600 | 12.38   | 0.05   | 254.8       | 1.12E-06    | SAM dependent carboxyl methyltransferase                  |
| Glyma.02g054200 | 37.71   | 0.17   | 225.5       | 9.33E-15    | SAM dependent carboxyl methyltransferase                  |
| Glyma.07g098500 | 19.19   | 0.13   | 144.4       | 6.39E-08    | copper transport protein atox1-related                    |
| Glyma.08g163600 | 1.29    | 0.02   | 74.4        | 2.13E-04    | pre-mRNA processing protein prp39-related                 |
| Glyma.09g022800 | 193.8   | 2.63   | 73.7        | 7.33E-15    | peroxidase                                                |
| Glyma.10g185900 | 8.01    | 0.11   | 71.5        | 2.34E-09    | sieve element occlusion protein                           |
| Glyma.09g024300 | 1199.19 | 17.48  | 68.6        | 2.51E-06    | glycosyl hydrolases family 18                            |
| Glyma.09g024200 | 1967.96 | 22.9   | 65.4        | 3.13E-05    | glycosyl hydrolases family 18                            |
| Glyma.09g024400 | 1102.06 | 16.91  | 65.2        | 3.53E-07    | glycosyl hydrolases family 18                            |
| Glyma.12g12470* | 10.06   | 0.16   | 62.1        | 3.05E-10    | ankyrin repeat-containing                                 |
| Glyma.10g266500 | 1.04    | 0.02   | 60.5        | 4.50E-04    | plant protein of unknown function                         |
| Glyma.03g02810* | 56.25   | 1.01   | 55.8        | 1.27E-13    | glycosid hydrolase                                       |
| Glyma.03g025000 | 53.91   | 0.97   | 55.5        | 3.73E-13    | glycosyl hydrolases family 18                            |
| Glyma.03g02818* | 32.43   | 0.58   | 55.5        | 5.20E-13    | unknown                                                  |
| Glyma.03g02843* | 45.41   | 0.82   | 55.2        | 1.66E-13    | unknown                                                  |
| Glyma.06g294400 | 7.77    | 0.16   | 47.7        | 1.73E-09    | ankyrin repeat-containing                                 |
| Glyma.13g113100 | 11.65   | 0.27   | 43.8        | 6.61E-10    | flavin-containing monoxygenase                           |
| Glyma.03g024800 | 64.4    | 1.53   | 42          | 1.09E-11    | glycosyl hydrolases family 18                            |
| Glyma.07g204900 | 6.09    | 0.16   | 39          | 6.94E-07    | lipase (class 3); alpha/beta-hydrolases superfAMILY protein |
| Glyma.14g032000 | 13.31   | 0.34   | 38.7        | 1.23E-04    | unknown                                                  |
| Glyma.06g013200 | 29.27   | 0.82   | 35.7        | 8.37E-12    | protein of unknown function (DUF2775)                    |
| Glyma.03g116300 | 108.09  | 3.21   | 33.7        | 2.86E-10    | ubiquitin specific protease family C19-related            |
| Glyma.06g300100 | 4.26    | 0.14   | 31.3        | 1.30E-05    | transcription factor; MYB-related                         |
| Glyma.01g115500 | 40.93   | 1.35   | 30.4        | 3.42E-08    | unknown                                                  |
| Glyma.10g944800 | 6.29    | 0.22   | 29          | 2.13E-05    | unknown                                                  |
| Glyma.14g051600 | 18.92   | 0.84   | 22.6        | 2.54E-07    | copper transport protein atox1-related                    |
| Glyma.13g205400 | 73.71   | 3.36   | 21.9        | 2.33E-07    | AT1G11655                                                |
| Glyma.14g061800 | 2.07    | 0.11   | 19.6        | 2.28E-04    | HXXXD-type acyl-transferase family protein               |
| Glyma.18g182800 | 6.03    | 0.32   | 18.6        | 5.34E-06    | 2-deoxyglucose-6-phosphate phosphatase 2                 |
| Glyma.18g39500* | 1.59    | 0.09   | 18.4        | 1.90E-05    | NADPH oxidase; flavin adenine dinucleotide binding        |
| Glyma.14g05621* | 4.93    | 0.3    | 16.3        | 2.87E-04    | copper transport protein atox1-related                    |
| Glyma.01g115700 | 4.63    | 0.29   | 15.9        | 4.71E-06    | plant protein of unknown function                        |
| Glyma.03g061200 | 18.63   | 1.2    | 15.5        | 1.60E-08    | plant protein of unknown function (DUF247)                |
| Glyma.07g18225* | 24.51   | 1.59   | 15.4        | 6.82E-06    | SGNH hydrolase-type esterase superfamily protein          |
| Glyma.11g216700 | 6.2     | 0.4    | 15.4        | 6.85E-05    | unknown                                                  |
| Glyma.03g141900 | 20.64   | 1.37   | 15.1        | 5.40E-06    | unknown                                                  |
| Glyma.16g197300 | 11.65   | 0.8    | 14.6        | 2.34E-04    | unknown                                                  |
| Glyma.16g197400 | 11.65   | 0.8    | 14.6        | 2.34E-04    | unknown                                                  |
| Glyma.08g189600 | 1.31    | 0.09   | 14.5        | 1.73E-04    | lipoxigenase                                             |
| Glyma.02g083500 | 351.22  | 26.37  | 13.3        | 4.78E-06    | extensin                                                 |
| Glyma.10g25800* | 10.86   | 0.87   | 12.5        | 9.47E-06    | serine-threonine protein kinase; disease resistance/LRR family |
| Glyma.19g144500 | 28.35   | 2.31   | 12.3        | 1.52E-04    | unknown                                                  |
| Glyma.06g149600 | 4.36    | 0.36   | 12.2        | 2.94E-04    | nodulin MtN21 /EamA-like transporter family protein       |
| Glyma.09g023000 | 542.08  | 44.79  | 12.1        | 4.68E-04    | peroxidase                                               |
| Glyma.0466s00200 | 63.01   | 5.4    | 11.7        | 5.39E-08    | unknown                                                  |

(Continued)
(NCBI) for assignment of the gene models associated with the infection-induced genes, and also for their functional annotations. Functional classification of these genes based on their putative molecular functions indicated that a majority of the genes (62 genes during ETP and 11 during LTP) had putative oxygen binding properties (Fig 2). A large number of infection-

### Table 1. (Continued)

| Locus ID       | RPKM  | Fold change | P-value  | Functional Annotation              |
|---------------|-------|-------------|----------|-----------------------------------|
| Glyma.09g099900 | 10.49 | 0.93        | 11.2     | 3.07E-05 serine/threonine protein kinase |
| Glyma.20g173800 | 13.19 | 1.2         | 11       | 1.25E-04 protein-tyrosine phosphatase 1 |
| Glyma.12g225700 | 6.51  | 0.59        | 11       | 4.85E-04 unknown                  |
| Glyma.20g215500 | 32.66 | 3.02        | 10.8     | 3.71E-08 unknown                  |
| Glyma.01g002300 | 3.63  | 0.34        | 10.7     | 1.63E-04 cation transport protein |
| Glyma.08g285400 | 6.34  | 0.61        | 10.4     | 2.88E-05 glycosyl hydrolase family 28 |

S1, roots tissues 3 and 5 days following water treatment; S3, root tissues 3 and 5 days following infection.

* Predicted genes in the old soybean genome sequence version [Glyma.Wm82.a1.v1.1 (Gmax1.01)].

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### Table 2. Genes down-regulated at late time period in soybean roots infected with *F. virguliforme*.

| Locus ID       | RPKM  | Fold change | P-value  | Functional Annotation              |
|---------------|-------|-------------|----------|-----------------------------------|
| Glyma.06g300000 | 27.9  | 0.1         | 187.4    | 5.81E-05 MYB-related; myb binding domain |
| Glyma.02g054200 | 33.6  | 0.4         | 93.7     | 4.79E-05 SAM dependent carboxyl methyltransferase |
| Glyma.07g234100 | 187.1 | 3.5         | 53.9     | 1.24E-05 uncharacterized protein |
| Glyma.15g082200 | 170.5 | 4           | 42.6     | 8.53E-07 cysteine proteinase cathepsin F |
| Glyma.16g038100 | 943.1 | 23.5        | 40.1     | 4.11E-06 uncharacterized protein |
| Glyma13g11969*  | 662.7 | 20.4        | 32.5     | 3.43E-06 uncharacterized protein |
| Glyma.09g022800 | 93.9  | 3.2         | 29.2     | 6.42E-06 peroxidase                 |
| Glyma.09g163800 | 434.4 | 15.3        | 28.3     | 6.47E-06 trypsin and protease inhibitor; endopeptidase inhibitor |
| Glyma.03g024400 | 664.4 | 23.5        | 28.3     | 3.92E-05 hydrolase activity         |
| Glyma.10g232100 | 2129.7| 77.8        | 27.4     | 8.57E-05 uncharacterized protein |
| Glyma.06g298700 | 639.1 | 23.6        | 27       | 1.72E-05 wound-induced protein; wound-responsive |
| Glyma.02g303200 | 211.4 | 8.1         | 26       | 7.51E-06 uncharacterized protein |
| Glyma.03g024300 | 635.2 | 25.7        | 24.7     | 9.18E-05 hydrolase activity         |
| Glyma17g03850*  | 32802 | 1367.3      | 24       | 6.10E-05 uncharacterized protein |
| Glyma.16g178000 | 262.1 | 11          | 23.8     | 1.21E-05 lipid-transfer/copper transport protein atox1-related |
| Glyma.11g224900 | 472.4 | 20.4        | 23.2     | 1.20E-05 uncharacterized protein |
| Glyma.17g039400 | 2205.5| 96.7        | 22.8     | 2.26E-05 uncharacterized protein |
| Glyma.13g282200 | 117.7 | 5.4         | 21.7     | 2.30E-05 wound-induced protein; wound-responsive |
| Glyma.13g282400 | 82.9  | 3.9         | 21       | 3.73E-05 wound-induced protein; wound-responsive |
| Glyma13g42850*  | 1375.5| 69.2        | 19.9     | 5.80E-05 uncharacterized protein |
| Glyma.12g048000 | 393.6 | 21.3        | 18.4     | 5.90E-05 uncharacterized protein |
| Glyma.01g210500 | 50.7  | 2.8         | 18.3     | 3.52E-05 oligopeptide transporter-related |
| Glyma.03g082100 | 197.6 | 13.1        | 15.1     | 9.50E-05 metallothion binding       |

S2, roots tissues 10 and 24 days following water treatment; S4, root tissues 10 and 24 days following infection.

* Predicted genes in the old soybean genome sequence version [Glyma.Wm82.a1.v1.1 (Gmax1.01)].

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Fig 2. Classes of genes up-regulated in roots infected with *F. virguliforme* as compared to the roots treated with water. A. A total of 238 genes were induced (with a FC $\geq 10$) in the pooled RNA sample of roots harvested 3 and 5 days following *F. virguliforme* infection as compared to the water control; and were classified based on their putative molecular functions. B. A total of 54 genes were induced in pooled RNA sample of roots harvested 10 and 25 days following *F. virguliforme* infection as compared to the water control; and were classified based on their putative molecular functions.

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induced genes (60 genes during ETP and 32 in LTP) encode proteins with unknown molecular functions.

We investigated the infection-induced genes for their putative biological processes (Fig 3). Again a large portion of genes induced during ETP (82%) and in LTP (92%) encodes proteins with unknown functions. The next category of genes induced during ETP (Fig 3A) includes 10 genes (4%) involved in carbohydrate metabolic processes. In addition, eight genes (3%) involved in the signal transduction, six (2.5%) in cell death, four (2%) responsive to stress and stimuli, four (2%) are transporters, and four (2%) involved in the lipid metabolism processes. During LTP, only four genes (8%) were assigned with a biological process and are likely involved in the carbohydrate metabolic process, signal transduction, and cell differentiation (Fig 3B).

We conducted GO analyses of the up-regulated genes for determining the putative sub-cellular locations of their encoded proteins. A large number of the genes (24%; 58 genes), up-regulated in infected-roots during ETP, most likely encode cell wall proteins (Fig 4A). Among the rest, 34 genes (14%) encode membrane bound proteins, 26 genes (11%) encode plasma membrane associated, 20 genes (8%) encode cytoplasmic proteins, and 14 genes (6%) encode endoplasmic reticulum proteins (Fig 4A). Similarly, the majority of the genes up-regulated in infected roots during LTP, 29 genes (37%) encode extracellular proteins and nine genes (12%) encode cell wall proteins (Fig 4B).

We observed that 48 of the 54 genes induced in the infected roots of LTP were induced also during ETP (Fig 1B; S1 and S3 Datasets). Functionally, 10 of the 50 genes up-regulated at both ETP and LTP belong to the cytochrome P450 CYP2 subfamily protein involved in the synthesis of defense metabolites [51]. Seven of these genes encode RmlC-like cupins with a nutrient reservoir activity; five genes encode peroxidases probably associated with signaling [52]; four genes encode receptor-like kinases (RLKs) presumably to modulate the induction of immunity [53]; three genes are members of the thaumatin family of pathogenesis-related protein that were shown to be induced during F. virguliforme infection [21], and three encode chitinase-related proteins involved in plant defense mechanisms [54,55]. Among the genes induced only in infected roots during LTP include: (i) a cupin-like gene, (ii) two defense/pathogenesis-related genes, and (iii) a gene with unknown function (S3 Dataset).

Functional classification of soybean genes repressed in roots infected with F. virguliforme

We observed that steady-state transcript levels of 51 soybean genes were decreased (with a FC ≥10) in F. virguliforme-infected roots as compared to that in the water control roots (Fig 1; Tables 1 and 2; S2 Dataset). We investigated the possible function of these infection-repressed genes for (i) molecular functions, (ii) biological processes and (iii) cellular locations through gene ontology (GO) analyses and results are presented in Figs 5–7.

GO analyses for molecular functions revealed that a large proportion of the down-regulated soybean genes do not show any identity to previously characterized genes and thus could be novel genes. For example, 40 genes (78%) with reduced transcript levels during ETP and 19 genes (78%) during LTP did not show homology to any functionally characterized genes (Fig 5). Infection-suppressed soybean genes encode proteins, most of which possess hydrolase and transferase activities at ETP (10% and 4% respectively) and lipid binding and transporter activities at LTP (11% for each category). Some of the repressed genes encode transferases, catalytic proteins, and transcription factors (Fig 5).

Reclassification of the down-regulated genes through GO analyses for biological processes again revealed that majority of the genes (83% in ETP and 96% in LTP) did not show identify
Fig 3. Classes of genes up-regulated in roots infected with *F. virguliforme* as compared to the roots treated with water. A. A total of 238 genes were induced (with a FC $\geq 10$) in the pooled RNA sample of roots harvested 3 and 5 days following *F. virguliforme* infection as compared to the water control. Induced genes were
to any genes with known biological processes. Down-regulated soybean genes with known biological processes during ETP include four (8%) genes for primary metabolic processes (lipid and carbohydrate metabolism), three (6%) for signal transduction, two (4%) for growth and cell death (Fig 6).

To shed light on the possible function of the down-regulated genes, we conducted GO analyses for putative sub-cellular location. Surprisingly, a significant proportion (15%) of the down-regulated genes encode plasma membrane proteins especially during ETP. These proteins may be involved in regulating signaling process during infection (Fig 7). Again, a large number of the genes (20 [39%] genes during and 14 [64%] in LTP) were found not to show similarity to any known functionally characterized genes.

Of the 51 genes repressed in soybean roots during ETP, only 15 with FC $\geq$ 15 continued to be suppressed during LTP following infection (Fig 1C; Tables 1 and 2, S2 Dataset). These genes include Glyma.02g054200 encoding a SAM (salicylic acid methyltransferase) dependent carboxyl methyltransferase, Glyma.15g082200 encoding a cysteine protease Cathepsin F, Glyma.13g282200 and Glyma.13g282400 encoding wound-induced proteins, Glyma.16g178000 encoding a lipid transfer protein, and Glyma.06g294400 and Glyma.12g111500 encoding ankyrin-repeat containing proteins.

Validation of transcriptomic data for a few selected genes repressed during SDS infection

To validate down-regulation of a few selected genes during infection, we performed a semi-quantitative reverse transcriptase polymerase chain reaction (RT-PCR) for one member from each of the four selected classes of down-regulated genes (Table 1). They are: (i) Glyma.01g171600 encoding an uncharacterized salicylate o-methyltransferase (SAM)-like protein, we termed GmSAM1, most likely involved in the conversion of salicylic acid to the volatile methyl salicylate, a plant defense signal [56] for defending attack of necrotrophic pathogens including F. virguliforme; (ii) Glyma12g12470 encoding an uncharacterized ankyrin repeat-containing protein, we termed it as GmARP1; (iii) Glyma.10g185900 encodes a sieve element-occlusion protein (SEO)-like protein (http://www.uniprot.org; http://www.phytozome.net); and (iv) Glyma.10g094800 encoding an uncharacterized, putative transmembrane protein [46] (http://www.phytozome.net).

RNA samples of soybean roots harvested 8 and 12 h and 1, 2, 3 and 5 d post inoculation with F. virguliforme or treated with water were considered for RT-PCR. The results of the RT-PCR analysis confirmed the observation of the RNA seq experiment for the four selected genes of interest (Fig 8). RT-PCR data also suggested that the down-regulation of three of the four selected genes, Glyma.01g171600, Glyma.10g094800, and Glyma12g12470, is very rapid with little or no detectible transcript levels observed 8 h post inoculation.

Alteration in expression of a F. virguliforme-repressed gene GmARP1 showed foliar SDS resistance

We hypothesized that F. virguliforme down-regulates transcription of some of the defense-related genes to cause susceptibility in soybean. To test this hypothesis, we induced the expression of GmARP1 (Glyma12g12470) in transgenic soybean plants following F. virguliforme
Fig 4. Classes of genes up-regulated in roots infected with *F. virguliforme* as compared to the roots treated with water. A. A total of 238 genes were induced (with a FC ≥10) in pooled RNA sample of roots harvested 3 and 5 days following *F. virguliforme* infection as compared to the water control. Induced genes were
Glyma.12g111500 and Glyma.06g294400 are two highly similar genes that encode ankyrin repeats containing proteins with 76% identity (S4 and S5 Figs), but with no identity to any functionally characterized ankyrin repeat-containing genes including GmNPR1 [57]. GmARP1 protein contains 218 residues and slightly larger than all known ankyrin repeat-containing proteins including ANK superfamily with 71–199 residues; ANK_2 with 81–171; ANK_4 with 145–199 residues.

We used three F. virguliforme-infection inducible promoters to induce the expression of GmARP1 during F. virguliforme infection. Three fusion genes were generated by ligating GmARP1 individually to these three promoters (S1A Fig) and used to transform soybean cultivar ‘Williams 82.’ R₀ plants were grown in a greenhouse and analyzed for integration of GmARP1 transgenes by genomic PCR (S1B Fig).

R₁ progenies of individual transformants (R₀) carrying any of the three fusion GmARP1 genes were inoculated with F. virguliforme Mont-1 isolate in a growth chamber (Fig 9A–9E). Approximately 30 to 50% of the R₁ plants showed no symptoms to slight yellowing with disease scores 1 and 2. The rest showed severe disease symptoms with interveinal to severe chlorosis and necrosis with disease scores 3 to 7 (Fig 9B). On the contrary, susceptible R₁ plants presumably lacking a functional transgene exhibited typical SDS foliar symptoms from the second week of infection and were severely diseased by the end of the fourth week following inoculation, with severe chlorosis and necrosis (disease scores 3 to 7), and reduced chlorophyll content as compared to the resistant R₁ progeny plants (Fig 9C). Moreover, resistant transgenic plants showed increased root weight as compared to the susceptible R₁ progenies implying root resistance of those resistant R₁ progenies to the pathogen (Fig 9D) and is supported by the levels of root resistance observed among the R₁ progenies (Fig 9E). Molecular characterization of the infected R₁ plants indicated that expression of foliar and root SDS resistance among the R₁ progenies of multiple transformants was associated with the expression of GmARP1 transgenes (Fig 9F). No transcript of the endogenous GmARP1 gene was detected during F. virguliforme infection.

We evaluated whether GmARP1 transgenes could provide enhanced SDS resistance in transgenic plants under field conditions. We observed that 70 to 100% of the transgenic R₁ plants descended from six independent transgenic R₀ plants showed enhanced SDS resistance under field conditions and did not develop SDS symptoms. Non-transgenic Williams 82 control plants developed severe SDS symptoms including chlorosis and necrosis of leaves that caused total defoliation of plants before maturity stage (Fig 10A and 10C; S6 Fig). The transgene copy number assay conducted using qPCR revealed that all six transgenic lines contain at least one copy of the transgene, sufficient to enhance resistance against F. virguliforme (Table 3; S4 Dataset). The line Prom2-ARP1-9 carrying single copy Prom2-ARP1 transgene showed 100% resistant R₁ progenies; whereas, Prom2-ARP1-9 line carrying on the average 4 copies of the same Prom2-ARP1 transgene showed the lowest number SDS resistant plants (70%).

**Discussion**

Study of the steady state transcriptomes using next-generation sequencing is a powerful method for revealing candidate genes that may play roles in host-pathogen interactions. Alterations of the expression of defense-related genes have been reported in plants that are infected by fungi, particularly by *Fusaria* species [18,58]. Transcriptomic studies have revealed that...
Fig 5. Classes of genes down-regulated in roots infected with *F. virguliforme* as compared to the roots treated with water. A. A total of 51 genes were repressed (with a FC ≥ 10) in pooled RNA sample of roots harvested 3 and 5 days following *F. virguliforme* infection as compared to the water control; and were classified based on their putative molecular functions. A total of 23 genes were repressed (with a FC ≥ 10) in pooled RNA sample of roots harvested 10 and 25 days following *F. virguliforme* infection as compared to the water control; and were classified based on their putative molecular functions.

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Fig 6. Classes of genes down-regulated in roots infected with *F. virguliforme* as compared to the roots treated with water. A. A total of 51 genes were repressed (with a FC ≥10) in pooled RNA sample of roots harvested 3 to 5 days following *F. virguliforme* infection as compared to the water control; and were classified based on their putative biological processes. B. A total of 23 genes were repressed in pooled RNA sample of roots harvested 10 and 25 days following *F. virguliforme* infection as compared to the water control; and were classified based on their putative biological processes.

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more host genes are induced and fewer repressed following infection with fungal pathogens [19,21,58,59]. The majority of the induced genes are defense related, and it has been shown that overexpression of defense genes induced during infection in transgenic plants could enhance resistance to pathogens [32,60].

Here we report GO analyses of differentially expressed soybean genes identified by comparing transcriptomes of soybean roots following *F. virguliforme* infection with those of water treated soybean roots. A large number of the genes induced in soybean roots infected with *F. virguliforme* encode cell wall and plasma membrane proteins (Fig 4; S1 Dataset) and are presumably involved in generating protective barriers against the invading pathogens. A majority of proteins encoded by these genes have binding activities. It is known that proteins and enzymes that bind other proteins and metabolites are implicated in various metabolic functions. For instance, DNA binding proteins have been shown to recognize specific promoter sequences of the target defense genes for regulating defense mechanisms [61,62].

A set of infection-induced genes are involved in carbohydrate metabolism. Besides being primary metabolites in plant cells, sugars are also known as signals for plant induced responses to pathogen attacks [63,64]. Many of the genes induced in roots following *F. virguliforme* infection encode enzymes with catalytic or kinase activities. Some of the kinases are receptor kinases for receiving and transmitting signals [65] and are induced in roots during fungal attack [33,54]. Our data suggest that defense-related transcripts are induced following infection presumably to defend soybean against the *F. virguliforme* attack.

In the Arabidopsis-*F. oxysporum* interaction a large number of genes were repressed as opposed to induction as observed in most plant-pathogen interactions [66]. Despite a large number of soybean genes were up-regulated upon *F. virguliforme* infection, in our analysis of the soybean-*F. virguliforme* interaction revealed that 70 soybean genes were suppressed following infection. Of these, only four genes were down-regulated in both ETP and LTP following *F. virguliforme* infection (Table 1; S2 Dataset).

Among the genes down-regulated following *F. virguliforme* infection, seven soybean genes encode glycosyl hydrolases (Table 1). Of these three genes Glyma.03g024400, Glyma.03g024300, and Glyma.03g024200 encoding the uncharacterized glycosyl hydrolase family 18 were highly expressed during ETP of the water treated root tissues (RPKM > 1,102). The expression of these genes was strongly suppressed upon *F. virguliforme* infection (FC > 65) (Table 1). Members of the glycosyl hydrolase 18 with high identity to some pathogen related (PR) genes are not only implicated in metabolic processes of cell wall, but also considered to have defense and signaling functions [67–70]. The glycosyl hydrolases encoded by Glyma.03g024400, Glyma.03g024300, and Glyma.03g024200 are 100% identical and contain a GH18_chitinase-like domain; therefore most likely they function in plant defense against pathogens with chitin molecules [71,72]. Four additional soybean genes, Glyma03g02810, Glyma.03g025000 Glyma.03g024800, and Glyma.08g285400 encoding glycosyl hydrolases with moderate expression levels (FC from 6 to 56), were also repressed during ETP following *F. virguliforme* infection.

Transcript levels of two genes Glyma.09g022800 and Glyma.09g023000 encoding uncharacterized peroxidases were strongly down-regulated following *F. virguliforme* infection (Table 1).
The two proteins are 64% identical and contain a secretory peroxidase domain. Some peroxidases genes are also shown to be PR genes [73] and involved in plant defense against pathogens. 

In our transcriptomic study, we included transcripts of roots of seedlings 3 and 5 days following F. virguliforme infection or water treatment for ETP. In the confirmatory RT-PCR experiment, we included roots of seedlings 8, 12 and 24 hours in addition to 3 and 5 days following infection or water treatment. Transcripts of all four genes included in the RT-PCR study were not detectable after 12 h following F. virguliforme infection (Fig 8; Table 1). Of these four genes, Glyma.10g185900 (Fig 6A and 6B; Table 1; S2 Dataset) encodes a sieve element-occlusion protein (SEO), which is a structural protein implicated to play a role in plant defense [74,75]. Our data suggest that somehow F. virguliforme suppressed the expression of this gene to avoid any barrier arisen from this protein in the roots. 

More than half of plant genes encode proteins of unknown functions [76]. We observed down-regulation of many genes encoding proteins of unknown functions during both ETP and LTP (FC > 10; Tables 1 and 2). During LTP, four genes have RPKM values in water treated roots over 1,000 (1,375 to 32,802) and are strongly repressed (FC ≥ 20) in F. virguliforme infected roots. The most highly expressed, Glyma17g03850 (RPKM in water treated roots 32,802 and repressed after infection to FC = 24.0), is an orphan gene of Glycine max [77] (http://www.greenphyl.org/) with no annotated functional domain. This gene is only co-expressed with Glyma.10g216000 (correlation 0.87, www.phytozome.com) that encodes a gibberellin-regulated protein 2 involved in gibberellic acid mediated signaling pathway [78]. Similarly, infection-suppressed gene Glyma.10g232100 (RPKM 2,130 in water treated roots vs. FC 27 following infection) encoding an unknown protein also co-expressed with the gibberellin-regulated protein 2. Finally, Glyma.17g039400 (RPKM 2,206 in water treated roots vs. FC 24 following infection). It is highly co-expressed with a fasciclin-like arabinogalactan protein gene (correlation 0.88, www.phytozome.com) implicated in the cell-wall composition [79]. Thus, these genes with unknown functions could be involved in plant immunity. Future studies on these four genes might yield new insights about the soybean- F. virguliforme interaction. 

Aknyrin repeat-containing proteins widely exist in plants [80]. We identified five GmARP1 homologs in G. soja and Phaseolus vulgaris exhibiting ≥ 62% identities among them, specifically in the ankyrin domain (S7 Fig). Two to more than 20 ankyrin-repeats involved in protein-protein interactions could be present in ankyrin-repeat containing proteins [81–83]. Ankyrin-repeats are involved in many metabolic processes including defense against pathogens and signaling [81,84–86]. The ankyrin repeat-containing ACD6 is implicated in salicylic acid signaling in Arabidopsis [87]. The ankyrin repeat-containing protein CaKR1 in Capsicum annuum is responsive to both abiotic and biotic stresses [88]. Ankyrin repeat-containing protein NPR1 modulates plant immunity cooperatively with transcription factors [89]. In Arabidopsis, BDA1 containing ankyrin repeats has been shown to regulate immunity [90]. Similarly,
Mou and colleagues [60] reported that overexpression of the ankyrin repeat-containing protein gene OsPIANK1 enhanced immunity against *Magnaporthe oryzae* in transgenic rice. *GmARP1*
Glyma12g12470 has been shown to have a protein binding function and putatively localized to plasma membrane [80]. Here we have shown that the steady state transcript level of this gene is not detectable immediately after *F. virguliforme* infection (Fig 8). Expression of this gene using infection-inducible promoters in transgenic plants enhanced resistance to the SDS pathogen, indicating that like other known ankyrin repeat-containing protein genes, GmARP1 is involved in soybean defense against this fungal pathogen.

**Conclusions**

Our transcriptomic study revealed many genes that are induced, whereas expression of only a few genes including a previously uncharacterized GmARP1 gene is strongly suppressed by *F. virguliforme* infection. Many of the genes that are suppressed during infection might be defense-related and, are somehow suppressed by pathogens to establish the compatible interaction or to cause susceptibility. This hypothesis was tested by expressing GmARP1 during *F. virguliforme* infection in transgenic soybean lines. We observed that expression of GmARP1 led to induction of foliar SDS resistance. How GmARP1 regulates immunity against SDS pathogen is yet to be determined. It is also unknown how the gene is down-regulated by *F. virguliforme* infection. Our transgenic study suggests that altered expression of pathogen-repressed host genes could be a suitable strategy in engineering disease resistance in crop plants.
Table 3. Average GmARP1 transgene copy numbers among the transgenic lines.

| Genotype      | Transgene copy number | Number of plants | Average copy number (± SE) |
|---------------|------------------------|------------------|---------------------------|
| Prom1-ARP1-5  | 1                      | 7                | 2 (0.06)                  |
|               | 2                      | 4                |                           |
|               | 3                      | 4                |                           |
|               | 1                      | 3                |                           |
|               | 3                      | 2                |                           |
| Prom2-ARP1-3  | 4                      | 3                | 4 (0.18)                  |
|               | 5                      | 1                |                           |
|               | 6                      | 1                |                           |
|               | 7                      | 2                |                           |
|               | 2                      | 7                |                           |
| Prom2-ARP1-7  | 3                      | 1                | 3 (0.14)                  |
|               | 4                      | 2                |                           |
|               | 5                      | 2                |                           |
|               | 1                      | 9                |                           |
| Prom2-ARP1-9  | 2                      | 2                | 1 (0.05)                  |
|               | 3                      | 1                |                           |
| Prom3-ARP1-3  | 2                      | 10               | 2 (0.02)                  |
|               | 3                      | 2                |                           |
|               | 1                      | 7                |                           |
| Prom3-ARP1-11 | 2                      | 4                | 2 (0.06)                  |
|               | 3                      | 1                |                           |

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Supporting Information

S1 Fig. Generation of GmARP1 transgenes and R₀ transgenic soybean plants.

(SDOCX)

S2 Fig. GmARP1 Sequences used in the transgenic study.

(SDOCX)

S3 Fig. Development of a binary construct for transformation of soybean.

(SDOCX)

S4 Fig. Screenshots of the GmARP1 gene in the new and the old versions of genome sequence assemblies (www.soybase.org).

(SDOCX)

S5 Fig. Sequence analysis of two ankyrin-repeat containing proteins downregulated following F. virguliforme infection.

(SDOCX)

S6 Fig. Phenotype of transgenic soybean plants carrying GmARP1 fusion genes in a field trial.

(SDOCX)

S7 Fig. Sequence analysis of five ankyrin-repeat containing GmARP1 (Glyma12g12470) homologs.

(SDOCX)

S8 Fig. RT-PCR analysis of three independent replicates for the four selected genes.

(SDOCX)
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