RESEARCH ARTICLE

The transcriptional reprogramming and functional identification of WRKY family members in pepper’s response to Phytophthora capsici infection

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Abstract

Background: Plant transcription factors (TFs) are key transcriptional regulators to manipulate the regulatory network of host immunity. However, the globally transcriptional reprogramming of plant TF families in response to pathogens, especially between the resistant and susceptible host plants, remains largely unknown.

Results: Here, we performed time-series RNA-seq from a resistant pepper line CM334 and a susceptible pepper line EC01 upon challenged with Phytophthora capsici, and enrichment analysis indicated that WRKY family most significantly enriched in both CM334 and EC01. Interestingly, we found that nearly half of the WRKY family members were significantly up-regulated, whereas none of them were down-regulated in the two lines. These induced WRKY genes were greatly overlapped between CM334 and EC01. More strikingly, most of these induced WRKY genes were expressed in time-order patterns, and could be mainly divided into three subgroups: early response (3 h-up), mid response (24 h-up) and mid-late response (ML-up) genes. Moreover, it was found that the responses of these ML-up genes were several hours delayed in EC01. Furthermore, a total of 19 induced WRKY genes were selected for functional identification by virus-induced gene silencing. The result revealed that silencing of CaWRKY03–6, CaWRKY03–7, CaWRKY06–5 or CaWRKY10–4 significantly increase the susceptibility to P. capsici both in CM334 and EC01, indicating that they might contribute to pepper’s basal defense against P. capsici; while silencing of CaWRKY08–4 and CaWRKY01–10 significantly impaired the disease resistance in CM334 but not in EC01, suggesting that these two WRKY genes are prominent modulators specifically in the resistant pepper plants.

Conclusions: These results considerably extend our understanding of WRKY gene family in pepper’s resistance against P. capsici and provide potential applications for genetic improvement against phytophthora blight.

Keywords: Pepper, Phytophthora, WRKY, Transcription factor, Transcriptome, Transcriptional reprogramming, Disease resistance

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**Background**

During co-evolution with diverse pathogens, plants have evolved a highly sophisticated and effective innate immune system to protect themselves against the pathogenic invaders. This system consists of two primary layers [1, 2], the first layer of plant immunity is triggered upon perception of highly conserved pathogen-associated molecular patterns (PAMPs) via plant pattern recognition receptors (PRRs) and is termed as PAMP-triggered immunity (PTI). PTI can be attenuated or blocked by effectors that are secreted into host cells by some adapted pathogens. The remaining weakened plant immunity during such compatible interactions is defined as basal defense, which is also activated in susceptible plants; however, it is not sufficient to prevent disease propagation [3]. The second layer of plant immunity is triggered by host receptors encoded by resistance (R) genes upon recognizing pathogen-delivered effectors either directly or indirectly and is termed as effector-triggered immunity (ETI), which brings out a more robust defense response and often accompanied by hypersensitive response [2]. Despite with variations in the magnitude and duration of immune responses, PTI and ETI share some common signaling components such as reactive oxygen species (ROS), MAPK cascades, phytohormones [4–7]. These defense signaling are generally integrated and relayed into appropriate immune outputs by the action of various transcription factors (TFs).

Both PTI and ETI are largely regulated at transcriptional level with the action of various plant TFs constituting transcriptional networks [8]. Over the past few decades, a large number of plant TFs, particularly in the model plants Arabidopsis and rice, have been functionally characterized to play important roles in modulating defense response [9, 10]. Accumulating data indicate that some plant TF families such as AP2/ERF, bHLH, bZIP, NAC and WRKY are key regulators in the defense processes [4, 9]. However, previous studies focused primarily on the functional characterization of individual TFs in host immune response, a genome-wide and systematic comparative analysis of certain plant TF families, especially between the resistant and susceptible host plants, will be valuable for elucidating their regulatory relationships during the pathogen infection.

Pepper (Capsicum annum), an economically important crop worldwide. Phytophthora blight of pepper is a devastating disease caused by the oomycete pathogen Phytophthora capsici [11, 12]. This pathogen can infect all parts of the pepper plant, including the roots, leaves and fruits [11, 13]. The disease frequently reaches epidemic levels and causes huge yield losses in pepper production regions. In plant-Phytophthora interaction system, recently several WRKY TFs have been identified to play important roles in plant defense against Phytophthora species. For example, WRKY TFs from Nicotiana benthamiana could be phosphorylated by MAPK and regulate immunity to P. infestans mediated by RBOHB-dependent ROS burst [14]. In Glycine max, GmWRKY31 and GmWRKY40 were identified in resistance to P. sojae [15, 16]. In Solanum tuberosum, StWRKY1 and StWRKY8 regulate phytoecdysteroid and benzylisoquinoline alkaloid pathway conferring resistance to P. infestans, respectively [17, 18]. In Solanum pimpinellifolium, eight WRKY TFs were identified to be involved in response to P. infestans infection by transcriptome analysis, and SpWRKY3 was found to act as a positive regulator in tomato resistance to late blight disease [19]. Overexpression of SpWRKY1 in tobacco and tomato conferred increased resistance to P. nicotianae and P. infestans, respectively [20–22]. Loss and gain of function analysis also indicated that SpWRKY6 acts as a positive regulator in tomato resistance to P. infestans infection [23]. Importantly, these results also indicate that a subset of WRKY TFs might be involved in response to a single pathogen infection. In Capsicum annum, although phytophthora blight caused by P. capsici is one of the most important diseases worldwide, the globally transcriptional reprogramming and functional identification of WRKY family members in defense against the pathogen remain largely unknown.

Herein, we performed time-series RNA-seq from a resistant pepper line CM334 and a susceptible pepper line EC01 upon challenged with P. capsici. The objective of this study was to identify key TF families and their family members involved in pepper defense against P. capsici infection and provide new insights into plant defense signaling regulation.

**Results**

**High-throughput RNA sequencing and DEG analysis**

To genome-wide investigate transcriptional regulation mechanism of pepper in response to P. capsici infection, time-series RNA-seq data from the resistant line CM334 and the susceptible line EC01 at 0, 3, 6, 12, 24, 48, and 72 h post inoculation (hpi) were analyzed. As shown in Fig. 1a, serious disease symptoms (wilt phenotype) were observed in the susceptible pepper line EC01 when inoculated with the virulent P. capsici strain JX1 but not in the resistant line CM334, which was in agreement with the result of the previous report [24]. Illumina-based next-generation sequencing was performed from pepper roots infected with P. capsici. In total, 42 samples (2 genotypes × 7 time points × 3 biological replicates) were collected for library construction. Approximately 80.2 million and 84.6 million raw reads were generated respectively from each sample of CM334 and EC01 (Additional file 1: Dataset S1). After filtered with low-quality reads, approximately 79.6 million and 84.0 million clean
reads were obtained from each sample of CM334 and EC01, respectively. On the average, more than 69.9 million (87.8%) and 68.0 million (80.1%) unique mapped reads from each sample of CM334 and EC01 were respectively aligned to the pepper CM334 genome version PEP (v1.6). Following alignment to each gene model, we normalized the number of mapped reads to fragments per kilobase million (FPKM). We identified differentially expressed genes (DEGs, with fold change > 2 and FDR ≤ 0.01) between the inoculated and mock-inoculated samples. Approximately 4075 DEGs with 2530 up-regulated and 1545 down-regulated genes were identified in the resistant pepper line CM334; while in the susceptible pepper line EC01, 437 TF genes (with 249 up-regulated and 188 down-regulated TF genes), which account for ~26.2% of all identified TF genes, were significantly altered during the infection (Fig. 1c). These results indicated that there are dramatic transcriptional changes both in CM334 and EC01 during P. capsici infection, while with more DEGs and differential expressed TFs in EC01.

Previous reports have suggested that some plant TF families such as AP2/ERF, bHLH, bZIP, NAC and WRKY are key regulators in defense response [4, 9]. To gain insights into the 58 TF families in response to P. capsici infection, we performed enrichment analysis using the identified TF DEGs in the two lines. Among all these TF families, it was found that WRKY family was the most significantly enriched both in CM334 and EC01 (Fig. 2 and Additional file 2: Figure S1). In-depth
analysis showed that nearly half of the WRKY genes were significantly induced in the resistant line CM334 (36, ~50.0%) and the susceptible line EC01 (34, ~47.2%). More strikingly, none of the WRKY genes were significantly down-regulated in both lines during the infection (Table 1 and Fig. 1d). Our comparative transcriptomic analysis suggested that WRKY may play a more critical role in modulating the host transcriptional immune response to *P. capsici*, and thus we then focused primarily on the WRKY family in the subsequent study.

**Transcriptome analysis of WRKY family members in pepper during *P. capsici* infection**

A total of 72 putative CaWRKY genes that contained the conserved WRKY domain were identified according to the pepper CM334 genome version PEP (v1.6). Due to their low homology with AtWRKYs from *Arabidopsis* and also avoid naming confusion [26, 27], we designated all these CaWRKY genes from CaWRKY01–CaWRKY12 according to their location on chromosomes. The chromosomal distribution of these CaWRKY genes was shown in Fig. 3.

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**Table 1** Global analysis of TF families in pepper’s response to *P. capsici* infection

|       | WRKY (72) | bZIP (53) | bHLH (129) | NAC (96) | MYB (107) | AP2/ERF (143) | All TFs (1665) |
|-------|------------|------------|-------------|----------|------------|---------------|----------------|
| CM334-up | 36 (50.0%) | 1 (1.9%)   | 10 (7.8%)   | 8 (8.3%) | 12 (11.2%) | 22 (15.4%)    | 137 (8.2%)     |
| CM334-down | 0 (0.0%)  | 5 (9.4%)   | 9 (7.0%)    | 1 (1.0%) | 3 (2.8%)   | 6 (4.2%)      | 121 (7.3%)     |
| EC01-up  | 34 (47.2%) | 8 (15.1%)  | 20 (15.5%)  | 20 (20.8%) | 17 (15.9%) | 36 (25.2%)    | 249 (15.0%)    |
| EC01-down | 0 (0.0%)  | 11 (20.8%) | 18 (14.0%)  | 1 (1.0%) | 8 (7.5%)   | 12 (8.4%)     | 188 (11.3%)    |
about these CaWRKY genes, including gene loci accession number in PEP (v1.6), WRKYGOK heptapeptide stretch, zinc-finger motif type and gene classification, was listed in Additional file 3: Dataset S2. The nucleotide and protein sequences of CaWRKY members were listed in Additional file 4: Dataset S3. The phylogenetic relationship between these 72 CaWRKY and 71 AtWRKYS was analyzed by multiple sequence alignment (Fig. 3). The result indicated that only 15 CaWRKYs exhibit a high similarity with their Arabidopsis WRKY orthologs.

Base on the transcript abundance and dynamic changes of gene expression, these CaWRKY family
members could be classified into three groups (Figs. 4 and 5a, b); group I, which included 17 and 19 CaWRKY genes in CM334 and EC01 respectively, were not or very low expressed (FPKM < 1.0); group II, which included 19 CaWRKY genes in both lines, were more-or-less constitutively expressed (FPKM > 1, fold < 2); group III, which were significantly up-regulated during the infection (FPKM > 1, fold > 2). The Z-scores of RNA-seq data sets were used for analysis of gene expression patterns using the ggplot2 package (https://cran.r-project.org/web/packages/ggplot2/) and construction of heat maps using the pheatmap package (https://cran.r-project.org/web/packages/pheatmap/).

![Heat map of WRKY family members in pepper’s response to P. capsici infection](image)

**Fig. 4** The heat map of WRKY family members in pepper’s response to *P. capsici* infection. Based on the transcript abundance and dynamic changes of gene expression, the 72 CaWRKY family members could be mainly classified into three groups: group I, were not or very low expressed (FPKM < 1); group II, more-or-less constitutively expressed (FPKM > 1, fold < 2); group III, significantly up-regulated during the infection (FPKM > 1, fold > 2). The Z-scores of RNA-seq data sets were used for analysis of gene expression patterns using the ggplot2 package (https://cran.r-project.org/web/packages/ggplot2/) and construction of heat maps using the pheatmap package (https://cran.r-project.org/web/packages/pheatmap/).

EC01 respectively, were significantly up-regulated during the infection. It was worth noting that the repertoires of these induced CaWRKY genes were greatly overlapped in CM334 and EC01, excepting that two CaWRKYs (CaWRKY08–4 and CaWRKY01–10) were resistant line specific up-regulated.

More interestingly, most of these induced CaWRKY genes could be mainly separated into three subgroups:
early response (3 h-up), mid response (24 h-up), and mid-late response (ML-up) genes (Fig. 5c, d and Additional file 5: Dataset S4). Among the 36 up-regulated CaWRKY genes in CM334, it was found that 18 of them had peaks of transcriptional induction at 3 hpi, 6 of them showed the highest transcripts level at 24 hpi, and 8 of them were mid-late response genes which sustain transcriptional induction at 12 ~ 72 hpi. Among the 34 up-regulated CaWRKY genes in EC01, 15 of them exhibited the highest transcripts level at 3 hpi, 5 of them showed the highest transcripts level at 24 hpi, and 9 of them were ML-up genes but with several hours delay. Our time-resolved transcriptome analysis suggested that 3 and 24 hpi are two of critical time points in the transcriptional reprogramming of WRKY family members in pepper’s response to *P. capsici* infection.

**Functional identification of WRKY family members in pepper defense against *P. capsici***

To further characterize their contributions in pepper defense against *P. capsici* infection, 19 of these induced CaWRKY genes with differential expression patterns were selected for functional identification. As shown in Fig. 6, all the tested genes validated by qRT-PCR analysis were most consistent with the results of RNA-seq data. We then performed knockdown experiments both in the susceptible line EC01 and the resistant line CM334 using the tobacco rattle virus (TRV) induced gene silencing system [28, 29]. The gene silencing efficiency of each CaWRKY TF was determined by qRT-PCR analysis at its highest induction time points after *P. capsici* inoculation. The results revealed that the silencing efficiency was greater than 65% for all the tested genes in EC01 and CM334, except for *CaWRKY08–4* and *CaWRKY01–10* (Fig. 7a, c and Additional file 6: Figure S2). Owing to up-regulated specifically in CM334 upon *P. capsici* infection, the transcript abundances of *CaWRKY08–4* and *CaWRKY01–10* were too low to detectable in EC01 after gene silencing.

After inoculation with *P. capsici*, the pathogen was restricted to small, localised lesions on TRV:0 treated CM334 leaves (no wilting); whereas, it expanded greatly and displayed much larger lesions on TRV:0 treated EC01 leaves, and most of the inoculated EC01 leaves displayed wilt phenotype within 3 days post
inoculation. For these TRV treated leaves targeting the selected CaWRKY genes, we found that most of the overlap induced CaWRKY TFs contribute with various degrees to pepper defense against *P. capsici* (Fig. 7b, d and Additional file 7: Figure S3). Among them, silencing of *CaWRKY03*–6, *CaWRKY03*–7, *CaWRKY06*–5 and *CaWRKY10*–4 significantly impaired the disease resistance both in CM334 and EC01. While for *CaWRKY08*–4 and *CaWRKY01*–10, silencing of them significantly impaired their resistance to the pathogen in CM334 but not in EC01, indicating that these two WRKY genes are prominent modulators specifically in the resistant pepper plants.

**Discussion**

It has been suggested that plant defense against pathogen attacks are regulated largely by a complicated transcriptional network [30], and plant TFs might be key modulators in plant immune response [9, 10]. Recently, several WRKY TFs from different plant species have been demonstrated to participate in plant defense against *Phytophthora* spp. [14–23]. However, despite that phytophthora blight frequently cause serious loss in pepper production, our knowledge on the roles of CaWRKY TFs in pepper immunity against *P. capsici* is very limited. In the present study, dynamic profiles of WRKY family genes between the resistant line CM334
and the susceptible line EC01 were comparatively assayed, and these CaWRKY genes up-regulated significantly against *P. capsici* was functionally investigated.

Within the approximately 1665 TFs classified into 58 families in pepper genome [25], a total of 258 and 437 TF DEGs were identified in the present study to be altered in their transcript levels in the resistant line CM334 and susceptible line EC01 upon *P. capsici* infection, respectively (Fig. 1c). The data from enrichment analysis showed that WRKY family was most enriched in both CM334 and EC01 (Fig. 2), implying that WRKY TFs might play more critical roles in modulating the host transcriptional immune against *P. capsici* infection. More in-depth analysis indicated that 36 CaWRKY genes in CM334 (~50.0%) and 34 in EC01 (~47.2%) were up-regulated, whereas none of them were down-regulated during the defense response (Table 1 and Fig. 1d). Similarly, a previous study showed that 27 of AtWRKY genes from *Arabidopsis* were induced upon flg22 elicitation at 2 h post treatment, while only two

![Fig. 7](image-url) The functional identification of WRKY family members in pepper’s defense against *P. capsici* infection by VIGS system. Nineteen of these induced CaWRKY genes with differential expression patterns were selected for virus-induced gene silencing (VIGS). The VIGS efficiency was determined by qRT-PCR analysis at its highest induction time points after *P. capsici* inoculation in (a) EC01 and (c) CM334. Expression levels were normalized with *CaActin*, and expressed as mean fold changes relative to TRV0-treated leaves, which were set as 1. The disease lesions were measured from detached leaves (n = 5) at 2.5 days post inoculation from (b) EC01 and (d) CM334. Asterisks indicate statistically significant differences compared with the TRV0 empty vector controls by the least significant difference (LSD) test (*P < 0.05; **P < 0.01). This experiment was repeated twice with similar results.
AtWRKY genes were down-regulated [31]. Naveed et al. also revealed that most of the TFs from Carrizo citrange, such as ERF, bZIP and DOF, showed mixed trend of up- and down-regulation, whereas most of the WRKY TFs were up-regulated during the \textit{P. parasitica} infection [32]. To our knowledge, there are only two reports about the response of CaWRKYs to \textit{P. capsici} infection so far [24, 26]. Lu et al. selected seven CaWRKY genes (4 of them belong to 3 h-up genes, one is constitutively expressed, one is very low expressed both in CM334 and EC01; and one belong to ML-up genes in EC01 and induced at 6 hpi in CM334 in our RNA-seq data) and identified their expression levels in response to \textit{P. capsici} at 1 day post inoculation by qRT-PCR analysis. This report revealed that six CaWRKY genes were induced after \textit{P. capsici} inoculation, whereas one CaWRKY gene did not display any significant change in CM334. In EC, after \textit{P. capsici} inoculation, one CaWRKY gene was up-regulated, while other six CaWRKY genes remained unchanged or slightly down-regulated. Recently, Zheng et al. indicated that at least 10 CaWRKY genes from pepper cultivar Zunla-1 were induced after \textit{P. capsici} inoculation using the RNA-seq data with four time points (0, 1, 2 and 3 days post inoculation), whereas three CaWRKY genes were down-regulated during the infection. In general, these reports show the similar trend in CaWRKY TFs against \textit{P. capsici} infection with our findings, but also display minor deference and this may be due to use different pepper lines, \textit{P. capsici} strains, and samples collected with different time points. Moreover, our data revealed that the repertoires of these induced CaWRKY genes were highly overlapped between CM334 and EC01. In line with several previous studies [4–6], it indicated that both the resistant and susceptible plants share a large number of WRKY TFs as common signaling components to modulate immune response.

According to their difference in transcription patterns, these induced WRKY genes can be mainly divided into three subgroups: early response (3 h-up), mid response (24 h-up) and mid-late response (ML-up) genes (Fig. 5). This time-order expression patterns suggest that 3 and 24 hpi are two critical time points in the transcriptional reprogramming of WRKY family members in pepper’s response to \textit{P. capsici}. In consistent with the pervious researches [11, 33, 34], we speculated that this may be closely related to the pathogenic process of \textit{P. capsici} during the colonization. At 2–4 hpi, the zoospores of \textit{P. capsici} shed their flagella, encyst and adhere to the plant surface; at this time point, PAMPs might be perceived by plant membrane-localized PRRs to trigger PTI or basal defense, leading to the first round of WRKY TFs induction [11, 31]. At ~24 hpi, the hyphae can penetrate the plant host cells and form haustorial structure; at this time point, pathogen derived effectors might be delivered into plant host cells and perceived by some lower-evolved intracellular host receptors that trigger defense signaling, leading to the second round of WRKY TFs induction [11, 35, 36]. On further ingress, due to devoid of the corresponding advanced intracellular host receptors (here termed R proteins) in the susceptible host plants, hyphae spread in a large number of susceptible host cells and reached vascular tissue, leading to wilt phenotype. While in the resistant host plants, effectors could be recognized by R proteins, the expansion of hyphae in the epidermal cells is restricted, and the vascular tissue colonization is absent [6, 11, 33, 34], which was in accordance with our observations of the disease phenotypes in CM334 and EC01 inoculated with \textit{P. capsici} (Figs. 1a and 7).

Although a large overlap was found in these induced CaWRKY genes between the resistant and susceptible host plants, there are also some CaWRKY genes with different expression profiles between CM334 and EC01. Comparative transcriptomic analysis indicated that the responses of ML-up CaWRKY genes in the resistant host plants are several hours earlier than those in the susceptible host plants (Fig. 5c and d). In particular, \textit{CaWRKY08–4} and \textit{CaWRKY01–10} were exclusively up-regulated in CM334, while very low expressed in EC01 (Figs. 4 and 6). Our knockdown experiments revealed that most of the overlap induced WRKY TFs contribute to basal defense in the resistant and the susceptible pepper plants against \textit{P. capsici} (Fig. 7). Although some of these induced CaWRKY genes, such as \textit{CaWRKY02–4} and \textit{CaWRKY02–8}, were not significantly contributed to the disease resistance, it could not rule out the possibility of their functional redundancy in the defense signaling. Silencing of \textit{CaWRKY08–4} or \textit{CaWRKY01–10} significantly impaired its resistance to the pathogen in CM334 but not in EC01, indicating that these two CaWRKY genes are prominent modulators specifically in the resistant pepper plants. In fact, some pathogens could deliver effectors to target multiple defense-promoting WRKY TFs, causing loss of WRKY-DNA binding and trans-activating functions needed for defense gene expression and disease resistance [37–39]. Whether \textit{Phytophthora} uses this particular strategy to inhibit the transcripts or functions of WRKY TFs (such as \textit{CaWRKY08–4} and \textit{CaWRKY01–10}) in the susceptible host plants, is required to further investigate. It is noted that the large numbers of TFs including WRKYs in response to pathogen infection might act in a complex regulatory network rather than in a linear manner [31, 40, 41]. Further identification of their
upstream signaling components (such as cis-elements and trans-factors) and downstream target genes might provide new insights into the molecular mechanism underlying pepper resistance to \textit{P. capsici}.

**Conclusions**

In this study, we performed time-series RNA-seq from the resistant and susceptible pepper plants upon challenged with \textit{P. capsici}, and revealed their transcriptional similarities and differences of WRKY family members in response to the pathogen. We also performed knock-down experiments by VIGS to functionally investigate their roles in disease resistance. Collectively, the data presented here considerably extend our understanding of WRKY family members in plant defense response, and also provide potential applications for genetic improvement against phytophthora blight.

**Methods**

**Plant materials, pathogen and culture conditions**

The seeds of \textit{P. capsici}-resistant pepper landrace line ‘Criollo de Morelos 334’ (CM334) and susceptible cultivar ‘Early Calwonder 01’ (EC01) were sown in a soil mix [peat moss: perlite, 2:1 (v/v)] in plastic pots, and were placed in a growth room under a condition of 25 °C, 60–70 mmol photons m$^{-2}$ s$^{-1}$, a relative humidity of 70%, and a 16-h light/8-h dark photoperiod [29]. A highly virulent \textit{P. capsici} stain JX1 was isolated by our laboratory and cultured as described previously [42]. Briefly, the \textit{P. capsici} stain was cultured on 10% (v/v) V8 agar medium, and then transferred to 10% (v/v) V8 liquid medium for 3 days at 25 °C in the dark. The mycelia of \textit{P. capsici} were washed intermittently with sterilized H$_2$O for three times to induce zoospore release [42].

**Transcriptome analysis**

To obtain RNA-seq data from \textit{P. capsici}-infected pepper tissues, four-week-old (at 5 true leaf stage) soilless cultivated pepper lines CM334 and EC01 were grown with a Holland solution under a 16-h light/8-h dark cycle at 25 °C prior to inoculation. The zoospores were counted using a hemocytometer and their density was adjusted to approximately 5 × 10$^5$ zoospores/mL. Pepper roots were immersed with the zoospore suspension to ensure the root surface can be adhered by enough zoospores, and then respectively harvested at 0, 3, 6, 12, 24, 48, 72 h after pathogen inoculation. RNA samples extracted from three biological replicates of each treatment were used for library construction. These constructed libraries (PE150) were then sequenced by the Illumina HiSeq2000 (Illumina Inc., San Diego, USA). After filtered with low-quality reads by BBTools (https://jgi.doe.gov/data-and-tools/bbtools), the clean reads were then aligned to the pepper CM334 genome version PEP (v1.6) (http://peppergenome.snu.ac.kr/) using HISAT2 program (https://github.com/DaehwanKimLab/hisat2). Following alignment to each gene model, we normalized the number of mapped reads to FPKM. The Z-scores of RNA-seq data sets were used for analysis of gene expression patterns using ggplot2 package (https://cran.r-project.org/web/packages/ggplot2/) and construction of heat maps using the pheatmap package (https://cran.r-project.org/web/packages/pheatmap/). Enrichment analysis was performed by ggplot2 package (http://had.co.nz/ggplot2/).

**Identification of CaWRKY family members**

The pepper annotated genome and protein sequences were downloaded from the CM334 genome version PEP (v1.6) (http://peppergenome.snu.ac.kr/). The WRKY domain (PF03106) was obtained from PFAM database (http://pfam.sanger.ac.uk/), and was used to identify putative CaWRKY proteins by HMMER 3.0 software program (http://hmmer.janelia.org/) according to the HMMR User’s Guide. The non-redundant CaWRKY protein sequences were further confirmed using SMART program (http://smart.embl-heidelberg.de/).

**Chromosomal location and phylogenetic tree analysis**

All identified CaWRKYs were mapped to the ‘CM334’ chromosomes in the pepper genome database using BLASTn. The MapInspect software (http://mapinspect.software.informer.com/) was used to map the gene locus on chromosomes. \textit{Arabidopsis thaliana} WRKY (AtWRKY) protein sequences were downloaded from TAIR (https://www.arabidopsis.org/browse/genefamily/WRKY.jsp). We performed a multiple alignment of the 72 CaWRKY and 71 AtWRKY full-length protein sequences using ClustalX2 (http://www.clustal.org/clustalx2/). The alignment result was used to construct a phylogenetic tree using the neighbor-joining method of PhyML software (http://www.atgc-montpellier.fr/phyml/).

**RNA extraction and qRT-PCR analysis**

Total RNA was extracted from \textit{P. capsici}-infected pepper tissues at the indicated time points as described previously [42]. In brief, total RNA was extracted by a PureLink RNA mini kit (Invitrogen, Carlsbad, CA, USA) and treated with RNase-free DNase I (Takara Bio, Kusatsu, Japan). Then, the first-strand cDNA was reversely transcribed by Superscript II reverse transcriptase (Takara Bio, Japan). To determine the relative transcription levels of selected genes, real-time PCR was performed with specific primers (Additional file 8: Table S1) according to the manufacturer’s instructions for the BIO-RAD Real-time PCR system (Foster City, CA, USA) and the SYBR Premix Ex Taq II system (TaKaRa). Three independent biological replicates of each treatment were performed.
The data were analyzed using the Livak method and calculated as a normalized relative expression level \(2^{-\Delta\Delta CT}\) [43]. The pepper housekeeping gene CaActin was served as an endogenous control [44].

The VIGS vectors construction

To construct virus-induced gene silencing (VIGS) vectors of the selected CaWRKY genes, each of the specific silencing fragment was determined by BLAST analysis using the VIGS tool in the Sol Genomic Network (SGN) website [45], and no off-target gene (which share no more than 19 bp matching fragment) was detected in the pepper genome cDNA database. The silencing fragments were amplified by PCR from cDNA of CM334 using PrimeSTAR GXL DNA Polymerase (Takara Bio Inc., Otsu, Japan) with the primers listed in Additional file 8: Table S1. The purified fragments were cloned into the entry vector pDONR207, and then cloned into the TRV silencing vector pTRV2 by Gateway® technology (Invitrogen).

VIGS of CaWRKY genes in pepper plants

For silencing of each CaWRKY gene in pepper plants, we employed the TRV-based VIGS system according to our previous studies [28, 29]. Briefly, the A. tumefaciens strains GV3101 containing pTRV1 and pTRV2: CaWRKY were resuspended in the induction medium (OD600 = 0.8) and mixed thoroughly at 1:1 (v/v) ratio, and then infiltrated into the two cotyledons of 2-week-old pepper plants. pTRV2:0 (empty vector) was served as a negative control. pTRV2: CaPDS, which silences pepper phytoene desaturase (PDS) gene to induce photobleaching phenotype, was used as an indicator control (Additional file 9: Figure S3). The agro-infiltrated pepper plants were kept in an incubator in darkness at 16 °C for 56 h, and then grown in the growth room under normal conditions as described above for 3–4 weeks [28, 29, 46].

P. capsici infection assays

For infection assays, the third and fourth detached leaves from the top of each TRV treated pepper plant were inoculated with the highly virulent P. capsici stain JX1 zoospores, respectively [42]. To distinguish the disease phenotypes among different VIGS plants, especially for the susceptible pepper line EC01, each detached leaf was inoculated with ~ 100 P. capsici zoospores (a relative low concentration) under low disease-pressure conditions. After P. capsici inoculation, the VIGS efficiency was determined by qRT-PCR analysis using the fourth detached leaves at the highest induction time points of each CaWRKY gene, and the third detached leaves were kept at high humidity in the dark at 25 °C for 2–3 days prior to measure disease lesions. This experiment was repeated twice, each time with five replicates.

Supplementary information

Additional file 1: Dataset S1. Summary of RNA-seq libraries of pepper samples infected with P. capsici.

Additional file 2: Figure S1. Supplementary enrichment analysis of TF DEGs between CM334 and EC01 after P. capsici infection. (A) Enrichment analysis was performed using all identified TF DEGs in CM334 and EC01. (B) Enrichment analysis was performed using the down-regulated TF DEGs in the two lines. The enrichment analysis was performed by ggplot2 package (http://pad.co.nz/ggplot2/).

Additional file 3 Dataset S2. The characterization of CaWRKY family members.

Additional file 4: Dataset S3. The nucleotide and protein sequences of CaWRKY family members.

Additional file 5: Dataset S4. The classification of WRKY family members in pepper’s response to P. capsici infection.

Additional file 6: Dataset S5. Summary of silencing efficiency of the selected CaWRKY genes in VIGS pepper plants. The VIGS efficiency was determined by qRT-PCR analysis at its highest induction time points after P. capsici inoculation in (A) EC01 and (B) CM334. The expression levels were normalized with CaActin, and expressed as mean fold changes relative to TRV0-treated leaves, which were set as 1.

Additional file 7: Figure S3. Phenotypes of representative detached leaves from other CaWRKY-silenced pepper plants. Disease symptoms of detached leaves in pepper lines (A) EC01 and (B) CM334 at 2.5 days post-inoculation with P. capsici.

Additional file 8: Table S1. Primers used in this study.

Additional file 9: Figure S4. A POS-silenced control in the VIGS experiment.

Abbreviations

DEGs: Differentially expressed genes; FDR: False discovery rate; FPKM: Fragments per kilobase million; hpi: Hours post inoculation; qRT-PCR: Quantitative real-time polymerase chain reaction; TFs: Transcription factors; TRV: Tobacco rattle virus; VIGS: Virus-induced gene silencing

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Consent to publication

Not applicable.

Authors’ contributions
WC and SH conceived and designed the research; WC, YJ, ML, JG and CJ performed the experiments; WC, YJ, JP, WT, JH, DG and SH analyzed the data; WC and SH wrote the paper. All of the authors read and approved the final manuscript.

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Availability of data and materials

All data generated in this study are included in the paper and in the supporting information files. The RNA-Seq data used in the current study are
available in the NCBI Sequence Read Archive (SRA) under accession number: PRJNA627546 (https://www.ncbi.nlm.nih.gov/sra/PRJNA627546).

Ethics approval and consent to participate
Not applicable.

Competition interests
The authors declare that they have no competing interests.

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