Inhibition of multiple defense responsive pathways by CaWRKY70 transcription factor promotes susceptibility in chickpea under Fusarium oxysporum stress condition

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Abstract

Background: Suppression and activation of plant defense genes is comprehensively regulated by WRKY family transcription factors. Chickpea, the non-model crop legume suffers from wilt caused by Fusarium oxysporum f. sp. ciceri Race1 (Foc1), defense response mechanisms of which are poorly understood. Here, we attempted to show interaction between WRKY70 and several downstream signaling components involved in susceptibility/resistance response in chickpea upon challenge with Foc1.

Results: In the present study, we found Cicer arietinum L. WRKY70 (CaWRKY70) negatively governs multiple defense responsive pathways, including Systemic Acquired Resistance (SAR) activation in chickpea upon Foc1 infection. CaWRKY70 is found to be significantly accumulated at shoot tissues of susceptible (JG62) chickpea under Foc1 stress and salicylic acid (SA) application. CaWRKY70 overexpression promotes susceptibility in resistant chickpea (WR315) plants to Foc1 infection. Transgenic plants upon Foc1 inoculation demonstrated suppression of not only endogenous SA concentrations but expression of genes involved in SA signaling. CaWRKY70 overexpressing chickpea roots exhibited higher ion-leakage and Foc1 biomass accumulation compared to control transgenic (VC) plants. CaWRKY70 overexpression suppresses H2O2 production and resultant reactive oxygen species (ROS) induced cell death in Foc1 infected chickpea roots, stem and leaves. Being the nuclear targeted protein, CaWRKY70 suppresses CaMPK9-CaWRKY40 signaling in chickpea through its direct and indirect negative regulatory activities. Protein-protein interaction study revealed CaWRKY70 and CaRPP2-like CC-NB-ARC-LRR protein suppresses hyper-immune signaling in chickpea. Together, our study provides novel insights into mechanisms of suppression of the multiple defense signaling components in chickpea by CaWRKY70 under Foc1 stress.

Conclusion: CaWRKY70 mediated defense suppression unveils networking between several immune signaling events negatively affecting downstream resistance mechanisms in chickpea under Foc1 stress.

Keywords: Immune response, Protein-protein interaction, Reactive oxygen species (ROS), R-protein signaling, Systemic acquired resistance (SAR), Transcriptional regulation
Background

Plant defense against pathogens are rapidly conveyed through cell surface receptors or by the intracellular immune receptors. Cell surface receptors usually recognize specific pathogen or microbe associated molecular patterns (i.e., PAMPs or MAMPs) and elicits Pattern Triggered Immunity (PTI). By contrast, intracellular receptors bind PTI suppressing effector proteins released by the pathogens which induce strong immune response, known as Effector Triggered Immunity (ETI) [1, 2]. WRKY transcription factors (TFs) are indispensable regulators of both PTI and ETI to wide variety of pathogens. The members of large multigene family transcription factor comprise WRKY domain (WRKY GQK) and zinc finger motif (CX4-7CX22-23HXC/C) that binds at TGAC core of W-box containing DNA [3, 4]. There are 74 WRKY family members present in Arabidopsis thaliana, which have been classified into three major groups (I, II and III) based on the number and position of WRKY domains and features of the zinc finger motif [5].

Transcriptional regulation of plant defense related gene expression by WRKY proteins are crucial to enable the induction of host immunity. Binding of WRKY70 TF at promoters of SA and JA signaling pathway genes, such as NPR1, PR2, PR10, VSP1 and VSP2 are associated with positive regulation of plant defense signaling [6–9]. AtWRKY33 overexpression leads to enhanced resistance against necrotrophic fungal pathogens, Botrytis cinerea and Alternaria brassicicola, although, plants showed susceptibility to Pseudomonas syringae infection [10]. WRKY28 and WRKY46 play co-transcriptional regulators of ISOCHORISMATE SYNTHASE1 (ICS1) gene expression and SA biosynthesis which mount defense response against biotrophic pathogens [11]. SA accumulation severely affects both PTI and ETI [12, 13]. Activation of Systemic Acquired Resistance (SAR) in the pathogen free distal tissues is also dependent on SA accumulation and signaling that trigger resistance against a large variety of pathogens, including viruses, bacteria and fungi [14–16]. In Arabidopsis, SA INDUCTION-DEFICIENT2 (SID2), ENHANCED DISEASE SUSCEPTIBILITY5 (EDS5), and NONEXPRESSOR OF PR GENES1 (NPR1) control SA production and signaling on pathogen challenge [17].

SID2 encodes an isochorismate synthase enzyme that converts chorismate to isochorismate [18]. Pathogen induced expression of SID2, and concomitant SA accumulation is regulated by SYSTEMIC ACQUIRED RESISTANCE DEFICIENT1 (SARD1) [19]. SARD1 positively regulates ICS1 gene expression that promotes pathogen-inducible SA accumulation in Arabidopsis [19, 20]. AtWRKY70 binds at promoter and inhibits SARD1 expression, which lowers the endogenous SA levels [20]. AtWRKY70 also functions as transcriptional regulator of JA/ ET induced gene expression and Induced Systemic Resistance (ISR) triggered by Bacillus cereus AR156 [21]. The apparent positive or negative effects of AtWRKY70 on transcription may thus provide the mechanistic basis for regulation of SA induced defense gene expression during local and systemic resistance in Arabidopsis.

ROS are the primary inducer for plant defense signaling that can trigger activation of mitogen activated protein kinase (MPK) cascade during plant-pathogen interplay [22, 23]. ROS production leads to upregulation of genes involved in SA- and JA/ ET- signaling pathway [24]. Furthermore, SA and ROS together play crucial roles in hypersensitive response (HR) triggered cell death signaling during SAR development in Arabidopsis [25]. Respiratory Burst Oxidase Homologs (RBOHs), a plasma membrane bound NADPH oxidase contribute ROS production in Arabidopsis thaliana and Nicotiana benthamiana [26, 27]. WRKYs are the transcriptional regulator of ROS production in these plants. WRKYs regulate the expression of AtRBOHDF and AtRBOHDF that mediate ETI-induced ROS bursts [26]. WRKY8 triggers NbRBOHB expression and HR induced cell death in N. benthamiana [27]. Treatment of Arabidopsis leaves with H2O2, a primary ROS candidate also upregulates the expression of many WRKY genes [28]. Thus, WRKY genes expression and ROS production are coordinately regulated at transcriptional level that prompts the activation of multiple defense signaling pathways like, hormonal crosstalk, ROS signaling, MAPK signaling, and HR associated cell death.

HR develops only when an appropriate Avr (avirulent) protein interacts with its cognate R (resistance) protein in planta [29, 30]. Effector proteins often target WRKYs in order to manipulate plant immunity. It is a well-known fact that WRKYs and R proteins serve common regulators of resistance signaling pathways to several plant-pathogen interactions. Arabidopsis Resistance to Ralstonia solanacearum 1 (RRS1) carries an extra integrated WRKY domain at its C-terminal end. This type of extended WRKY module perceives PopP2 effector protein and protects acetylation of other WRKYs upon instigating strong immune responses to the bacterial pathogen R. solanacearum [31]. It is important that RRS1 with its single WRKY domain can induce transcriptional reprogramming during ETI. WRKY70 also contributes to Recognition of Peronospora Parasitica 4 (RPP4)-mediated resistance against Hyaloperonospora parasitica [32]. Our recent study has established that Foc1 resistance in chickpea is dependent on the interaction between RPP2-like CC-NB-ARC-LRR protein and CaWRKY64 [30].

The present study has been focussed on chickpea-Fusarium interaction since, a smaller number of reports are currently available on legume-fungus interactions and detailed molecular regulations are undoubtedly obscured. Chickpea (Cicer arietinum L.) is the world’s third most important pulse crop and a rich source of plant-
derived edible protein. Chickpea production has been severely affected by wilt-causing hemi-biotrophic fungus Fusarium oxysporum f. sp. ciceri Race1 (Foc1) [33]. Amongst the eight different pathogenic races of Fusarium oxysporum f. sp. ciceri, Race1 is known to have cosmopolitan distribution causing significant yield losses. Foc1 infection accounting 10–15% annual crop loss and reaches 90–100% during favourable season [34, 35]. Foc1 invades chickpea through roots and grows to shoots where it colonizes the xylem vessels at root-stem interface region. Increasing fungal biomass blocks water supply to the aerial shoots, which results in massive vascular wilting [34, 35]. Foc1 resistance in chickpea is hard to achieve by usual breeding approaches due to limited genetic resources and elevated autogamy [36]. We used wilt-susceptible JG62 and wilt-resistant WR315 chickpea accessions to unveil the immunomodulatory role of CaWRKY70 protein on Foc1 infection [36, 37]. Our study shows that CaWRKY70 transcription factor promotes susceptibility in chickpea upon Foc1 infection. CaWRKY70 inhibits SA concentrations and signaling in non-inoculated distal shoot tissues of transgenic chickpea. Transcripts measurement data suggests CaWRKY70 functions as negative regulator for subsets of immune-responsive genes that control defense responses in chickpea, including PR genes. CaWRKY70 also suppresses endogenous ROS levels and R-protein induced ectopic cell death. Together, we establish that CaWRKY70 negatively impacts defense signaling and SAR development in chickpea under Foc1 stress.

**Results**

**Systemic expression pattern of Cicer arietinum L. WRKY70 (CaWRKY70) under SA induction and Foc1 infection**

CaWRKY70 expression in different chickpea tissues were determined by quantitative real-time PCR (qRT-PCR) analysis. CaWRKY70 mRNA expression was detected in all tissues including root, shoot, flower and pod. CaWRKY70 transcript accumulation was higher at shoot tissues compared to flower, root and pod in non-infected plants. CaWRKY70 expression was detected to be ~ 2.5-fold higher at shoot tissues than the roots (Fig. 1a). Differential salicylic

![Fig. 1](image-url)
acid (SA) accumulation induces SAR activation at distal shoot tissues of susceptible and resistant chickpea plants upon Foc1 infection [38]. To investigate whether SA signal-influences CaWRKY70 expression in chickpea, we measured CaWRKY70 transcript abundance in shoot tissues of susceptible and resistant chickpea at 6 h of SA treatment. Result suggests that SA treatment promotes significantly higher accumulation of CaWRKY70 transcript at shoot tissues of susceptible chickpea over control treatment. However, resistant plant shows less induction of CaWRKY70 transcript upon exogenous SA application (Fig. 1b). Other inducers like, ABA and JA failed to stimulate CaWRKY70 expression neither in susceptible nor in the resistant chickpea shoots. Therefore, it may be suggested that differential expression of CaWRKY70 in chickpea is mediated through SA response under Foc1 infection. Arabidopsis WRKY70 is an important WRKY member that has been shown to regulate SAR activation against biotrophic pathogens [9, 39]. To ascertain whether CaWRKY70 is associated with the systemic defense responses of chickpea, we sought to determine its mRNA expression at shoot tissues under control treatment and Foc1 infection. Susceptible and resistant chickpea plants subjected to Foc1 infection at 1, 2, 3, 4 and 7 days were used for RNA isolation, cDNA preparation and qRT-PCR analyses. CaWRKY70 expression at different time-points as compared to the 0 dpi control treatment in susceptible JG62 and resistant WR315 were plotted (Fig. 1c). CaWRKY70 fold change levels were normalized to a value of 1 at 0 dpi in JG62 and WR315, respectively. CaWRKY70 transcript was found to be induced at shoot tissues of susceptible chickpea on challenge with Foc1. On the contrary, resistant chickpea plants were unable to stimulate CaWRKY70 expression at shoot tissues under Foc1 stress. Time-dependent data revealed that CaWRKY70 transcript was found to be ~100-fold upregulated at 3 dpi in susceptible chickpea shoots over control treatment (Fig. 1c). However, the inclusion of mock treatment to each time point would have been useful to determine the developmental stage specific CaWRKY70 expression in both susceptible and resistant chickpea upon Foc1 infected conditions. To investigate CaWRKY70 protein accumulation, total proteins were extracted from control and Foc1 infected susceptible and resistant chickpea shoot tissues. Next, immunoblotting experiment was performed using anti-CaWRKY70 polyclonal antibody. Result shows induction of ~35 kDa CaWRKY70 protein band in non-inoculated systemic shoot tissues of susceptible plants upon Foc1 infection, whereas resistant plants failed to stimulate such protein accumulation (Fig. 1d). The protein level was higher at shoot tissues of susceptible plant at 4 dpi on Foc1 challenge, suggesting systemic accumulation of CaWRKY70 in susceptible chickpea.

CaWRKY70 is a nuclear targeted protein
To test subcellular localization of CaWRKY70 protein, we examined its localization in chickpea leaf tissue. We fused yellow fluorescence protein (YFP) at its C-terminus. CaWRKY70 protein fused to YFP was expressed in onion epidermal cells by Agrobacterium mediated transient transformation. Contrarily, an empty construct expressing control YFP was tested. Onion epidermal cells were stained with nuclear marker 4′,6-diamidino-2-phenylindole (DAPI). Confocal microscopic analyses showed that CaWRKY70-YFP was localized in nucleus. In contrast, localization of control YFP protein was observed throughout the cells (Additional file 1: Figure S1). Result also revealed that blue color fluorescence of DAPI-stained nuclei is overlapped with yellow color fluorescence of YFP, thereby, confirming nuclear localization of CaWRKY70.

CaWRKY70 overexpression triggers susceptibility in chickpea to Foc1
To examine the possible role of CaWRKY70 in defense regulation against Foc1, full-length CaWRKY70 gene was isolated from Foc1 treated shoot tissues of susceptible chickpea cDNA sample. Chimeric CaWRKY70 gene construct was prepared using pCAMBIA2301 vector and positive clones were selected by PCR analyses (Additional file 1: Figure S2). CaWRKY70 gene containing plasmids were delivered into resistant chickpea genome via Agrobacterium mediated transformation and transgenic chickpea plants were established (Fig. 2a, b). Two independent CaWRKY70 overexpressing chickpea lines i.e., OEX1, and OEX3 of T2 generation were obtained from primary transformants. Real-time PCR analysis detected that overexpressing plants exhibit significantly higher expression of CaWRKY70 transcript than control vector transformed plants, where lowest expression of CaWRKY70 mRNA was noted (Fig. 2b). To compare disease symptoms on control transgenic and CaWRKY70 overexpressing chickpea, we treated plants with Foc1 for 0, 3, 7 and 12 days, respectively. Result shows that CaWRKY70 overexpressing chickpea plants exhibit enhanced susceptibility to Foc1 infection in comparison to the control transgenic plants. Disease symptoms were much pronounced in overexpressing chickpea plants at 12 dpi with Foc1 (Fig. 2c). Foliar symptom that developed on control transgenic and CaWRKY70 overexpressing chickpea upon interaction with Foc1 was used to measure the disease intensity index. Based on incidence of infected plants and foliar symptoms, CaWRKY70 overexpressing chickpea was found to be highly susceptible to Foc1 than control transgenics. Disease symptom started developing at 3 dpi, progressed at
Fig. 2 CaWRKY70 overexpression induces susceptibility in chickpea under Foc1 infection. a Diagram of gene construct used for CaWRKY70 overexpression in resistant chickpea. b qRT-PCR determination of CaWRKY70 mRNA level in control transgenic (VC) and overexpressing chickpea. Error bars indicate ±SD of three independent biological samples. Data was normalized to CaGAPDH. Fold change values relative to control vector transformed plants. c Development of disease symptom in control transgenic and CaWRKY70 overexpressing chickpea. Disease phenotype of transgenic plants were photographed under control treatment (0 dpi) and Foc1 infection at 3, 7 and 12 dpi, respectively. d Assessment of disease intensity index in control transgenic and CaWRKY70 overexpressing chickpea under control treatment (0 dpi) and Foc1 infection. Chickpea plants grown in the soil-rite mixture were inoculated with Foc1. e Effect of Foc1 infection on the control transgenic and CaWRKY70 overexpressing chickpea based on incidence of dead or wilted plants. In d and e, each data point represents mean values of three pots with five plants per pot. Error bars indicate ±SD of three independent biological samples. f Electrolyte leakage in control transgenic (VC) and CaWRKY70 overexpressing chickpea roots under control treatment (0 dpi) and Foc1 infection. Each time point represents ±SD of three biological replicates. *P ≤ 0.05 and **P ≤ 0.01 indicate values show significant differences between control transgenic (VC) and CaWRKY70 overexpressing chickpea as determined by Student’s t test. g Quantitation of Foc1 5.8S rDNA at root tissues of control transgenic (VC) and CaWRKY70 overexpressing chickpea plants by real-time PCR. Chickpea DNA amount was normalized by CaGAPDH expression. Each bar represents mean ± SD of three independent biological replicates and fold change is relative to control treatment (0 dpi). **P ≤ 0.01 indicate mean values significantly different from control vector transformed plant determined by Student’s t test. h Relative water content percent (RWC%) in control transgenic and CaWRKY70 overexpressing chickpea leaves at different infection time-points. Asterisks (*) indicate significant difference at ***P ≤ 0.001 by one-way ANOVA followed by multiple comparison of means using tukey’s post-hoc test. i Chlorophyll A and j Chlorophyll B content in control transgenic and CaWRKY70 overexpressing chickpea leaves at different infection time-points. In i and j, error bars represent ±SD (n = 3). *P ≤ 0.05 indicate mean values are significantly different from control vector transformed plants as determined by Student’s t test.
higher rates in *CaWRKY70* overexpressing chickpea resulting in 100% plants with vascular wilt developed at 12 dpi (Fig. 2d), *CaWRKY70* transgenic plants developed highly susceptible reaction to Foc1 reaching 100% incidence of dead plants at 12 dpi. In contrast, control transgenic plants demonstrated less severe reaction at 12 dpi with only 20% of the plant’s death (Fig. 2e). Foc1 infection progressively enhances cell-death induced ion-leakage at root tissues of control transgenic and *CaWRKY70* overexpressing chickpea. Time-dependent analyses show significant difference in Foc1 induced electrolyte leakage in control transgenic and *CaWRKY70* overexpressing chickpea with higher leakage of ions in overexpressing chickpea roots and least in control transgenic plants. The conductivity was found to be significantly increased in *CaWRKY70* overexpressing root at 12 dpi of Foc1 infection than control transgenics (Fig. 2f). Amount of Foc1 biomass in root tissues of *CaWRKY70* overexpressing chickpea appears to be significantly higher than control transgenic. The relative accumulation of Foc1 5.8S rDNA was ~2.0-fold and ~7.0-fold at root tissues of control transgenic and *CaWRKY70* overexpressing chickpea, respectively, under Foc1 infection (Fig. 2g). Relative water content (RWC) was determined to compare percentage amount of water restored within the plant body of control transgenic and *CaWRKY70* overexpressing chickpea under control condition and Foc1 infection. Result shows that control transgenic plants retain ~77% RWC upon Foc1 infection whereas, the value was markedly reduced in *CaWRKY70* overexpressing chickpea plants under Foc1 stress i.e., ~46% and ~34% RWC (Fig. 2h). Foc1 inoculated chickpea plants show chlorosis of leaves accompanied by yellowing to browning or cell death (Fig. 2i). Decrease in chlorophyll A and chlorophyll B content was measured using leaves of control transgenic and *CaWRKY70* overexpressing chickpea in time dependent manner. Control transgenic plants retain significantly higher chlorophyll A and chlorophyll B content than overexpressing chickpea on Foc1 stress (Fig. 2j, k). Nevertheless, loss of total chlorophyll content in chickpea leaves might be a secondary effect of Foc1 infection. Taken together, our results confirm that *CaWRKY70* promotes susceptibility in chickpea to Foc1.

**CaWRKY70 reduces ROS accumulation and cell death in chickpea**

Histochemical DAB and trypan blue staining was performed to compare H₂O₂ accumulation and cell death between control transgenic and *CaWRKY70* overexpressing chickpea under Foc1 stress. Result demonstrates intense DAB and trypan blue colouration in Foc1 infected control transgenic roots (Fig. 3a, b). On the contrary, DAB and trypan blue staining was not observed at *CaWRKY70* overexpressing chickpea roots under Foc1 stress. H₂O₂ treatment was carried out as primary ROS inducer in transgenic chickpea roots which show intense DAB and trypan blue staining at roots of control transgenic plants. DAB colouration was not demonstrated by *CaWRKY70* overexpressing chickpea roots. However, faint trypan blue colour was retained by overexpressing plant roots only after H₂O₂ treatment (Fig. 3a, b). DAB and trypan blue staining was also performed using leaves and stem parts of the control transgenic and *CaWRKY70* overexpressing chickpea plants. Here, no such DAB or trypan blue staining was noted upon control treatment, however, Foc1 infection resulted in higher deposition of brownish DAB precipitates in leaves and stem tissues of control transgenic plants in comparison to *CaWRKY70* overexpressing chickpea (Fig. 3c). Similarly, strong trypan blue colouration was noted on leaves and stem portions of control transgenic plants than *CaWRKY70* overexpressing chickpea upon Foc1 infection (Fig. 3c). It is noteworthy that *CaWRKY70* overexpressing chickpea leaves and stem show mild histochemical DAB and trypan blue staining. We further measured DAB and trypan blue colour intensities of control transgenic and *CaWRKY70* overexpressing chickpea root, leaf and stem tissues. Quantitative measurements revealed significant reduction in the DAB and trypan blue intensities at roots, leaves and stem parts of *CaWRKY70* overexpressing chickpea compared to control transgenic plants under Foc1 stress (Fig. 3d, e). Although, control treatment does not exhibit such drastic changes in DAB and trypan blue intensities of empty vector transformed and *CaWRKY70* overexpressing chickpea plants. Therefore, it can be concluded that *CaWRKY70* negatively affects ROS accumulation and cell death induction and thus promotes susceptibility in *CaWRKY70* overexpressing chickpea plants upon Foc1 infection.

**CaWRKY70 inhibits SA signaling in transgenic chickpea**

ICS1 and PAL contribute to the pathogen induced SA production through isochorismate and phenylpropanoid pathways, respectively [18, 40]. In *CaWRKY70* overexpressing chickpea shoots, *CaICS1* and *CaPAL* transcripts were found to be downregulated after Foc1 infection at 2, 3, 4 and 7 days. Interestingly, pCAMBIA2301 vector transformed plant demonstrates mRNA induction upto 4 dpi and then downregulation at 7 dpi under Foc1 stress (Fig. 4a, b). SA concentrations were also significantly decreased at shoot tissues of *CaWRKY70* overexpressing chickpea in comparison to the control transgenic plants at 7 dpi with Foc1 (Fig. 4c). Result shows higher accumulation of SA in non-inoculated systemic shoot tissues of control vector transformed chickpea after Foc1 infection. It has been
observed that control transgenic and CaWRKY70 overexpressing chickpea exhibit basal accumulated level of SA production at shoot tissues under control treatment. Thus, CaWRKY70 reduces both SA biosynthesis genes expression and SA accumulation at shoot tissues of transgenic chickpea.

**Differential expression pattern of defense related transcripts in vector transgenic and CaWRKY70 overexpressing chickpea under Foc1 infection**

We compared the expression of defense responsive transcripts between vector inoculated and CaWRKY70 overexpressing transgenic chickpea on Foc1 challenge. CaWRKY33 transcript was remarkably lower at the shoot tissues of overexpressing chickpea than vehicle transgenics (Fig. 5a). WRKY54 and WRKY70 are the defense related transcription factors that negatively regulate osmotic stress tolerance in Arabidopsis [41]. At shoot tissues of CaWRKY70 overexpressing chickpea, CaWRKY54 mRNA was upregulated after Foc1 treatment (Fig. 5b). CaWRKY40 expression in Arabidopsis enhanced resistance to virulent Pseudomonas syringae pv. tomato DC3000 infection [42]. Furthermore, CaMPK9-CaWRKY40 signaling promotes primary defense responses in chickpea against Foc1 [43]. CaWRKY70 inhibits CaWRKY40 expression in transgenic chickpea (Fig. 5c). CaWRKY70 overexpressing chickpea also demonstrates a sharp decrease in CaMPK9 transcript level (Fig. 5d). EDS1 and PAD4 complex formation was found to be required for pathogen infection induced SA accumulation [44]. However, both CaEDS1 and CaPAD4 mRNAs were downregulated in
CaWRKY70 overexpressing chickpea plant type (Fig. 5e, f). Recent finding showed that phosphorylation dependent changes in AtNPR1 promotes its interaction with AtWRKY70 which suppresses Pathogenesis Related (PR) gene transcription in Arabidopsis [45]. Present result suggests that CaNPR1, CaPR1 and CaPR5 transcript levels were reduced in CaWRKY70 overexpressing chickpea in comparison to the vehicle treated plants (Fig. 5g, h and k). Among other SA signaling genes, expression of CaTGA1 and CaTGA6 mRNAs were downcast in CaWRKY70 overexpressing chickpea (Fig. 5i, j). Induction of JA-signaling gene CaDefensin mRNA is also inhibited at shoot tissues of CaWRKY70 over-accumulating plants than control vector transformed chickpea (Fig. 5l). Overall, CaWRKY70 negatively regulates induction of defense related genes expression at shoot tissues of transgenic chickpea upon Foc1 infection.

CaWRKY70 represses CaWRKY40 promoter activity

As described above, CaWRKY70 overexpressing chickpea significantly lowered the expression of CaWRKY40 mRNA under Foc1 infection; we got interested to check whether CaWRKY70 plays any direct modulatory role in the suppression of CaWRKY40 promoter activity. ChiP-PCR assay revealed that CaWRKY70 physically associates with W-box 2 at CaWRKY40 promoter. CaWRKY70 binding was observed at shoot tissues of susceptible (JG62) chickpea plants under Foc1 stress, whereas resistant chickpea shoot does not exhibit in vivo CaWRKY70 binding (Fig. 6a). Next, we tested this binding through DNA-protein docking experiment. Phyre2 server used WRKY transcription factor 1 (PDB ID: c2aydA) as template structure with 100% confidence and 34% coverage (Additional file 1: Figure S3a-c). Furthermore, qualitative assessment of the predicted model was evaluated by Ramachandran plot analysis using RAMPAGE server. Ramachandran plot analyses revealed that in CaWRKY70 model 94.7% residues are in favoured region, 5.3% residues are in the allowed region and none of the residues in outlier location (Additional file 1: Figure S3d-e). For understanding the molecular mechanism of its interaction with W-Box DNA, an in silico DNA-protein docking was carried out with HADDOCK (Fig. 6b). HADDOCK web server clustered 174 structures in 12 clusters, representing 87.0% of the HADDOCK generated water-refined models. Clusters were ranked according to HADDOCK score and Z-score for plotting DNA-protein interaction. Top five best scoring clusters are provided in Additional file 2: Table S1. HADDOCK score is calculated as the weighted sum of van der Waals, electrostatic, desolvation and restraint violation energies whereas, the Z-score indicates how many standard deviations from the average of these clusters is in terms of score. For predicted protein model, HADDOCK score v/s i-RMSD (interface-RMSD) plot was created. i-RMSD was calculated based on the backbone (CA, C, N, O, P) atoms of all residues involved in intermolecular contact using 10 Å cut-off. i-RMSD (ligand-RMSD) was also calculated on the backbone atoms of all (N > 1) molecules (Additional file 2: Table S1, Fig. 6b). DNA-protein interaction was further confirmed by in vitro EMSA.
Result indicates in vitro CaWRKY70 binding to wild-type (TGAC), and mutated W-boxes (TAGC and TAGA) (Fig. 6c). Approximately, 150 ng of purified histidine tagged CaWRKY70 exhibits strong DNA binding to W-box 2 at CaWRKY40 promoter. However, such binding was completely inhibited upon mutating G nucleotide of TGAC i.e., TAAC. In vitro DNA binding of CaWRKY70 was outcompeted using 5 and 10 M excess cold probes, respectively (Fig. 6d). Effect of CaWRKY70 binding at W-box 2 of CaWRKY40 promoter was tested by transient co-infiltration experiments using Nicotiana xanthi protoplasts and Nicotiana tabacum leaf discs, respectively. Protoplast co-transfection experiment showed that CaWRKY70 effectively inhibits CaWRKY40 promoter mediated expression of YFP in N. xanthi protoplast, which suggests CaWRKY70 mediated negative regulation of CaWRKY40 promoter activity in vivo (Fig. 6e). Agrobacterium-mediated transient co-infiltration of p35S:CaWRKY70 (effector construct) and pCaWRKY40:GUS (reporter construct) in tobacco leaf discs demonstrates reduction in CaWRKY40 promoter driven GUS expression (Fig. 6f). Quantitative data also revealed ~2.5-fold reduction in the histochemical GUS staining upon constitutive induction of CaWRKY70 (Additional file 1: Figure S4). Therefore, CaWRKY70 binds to and represses CaWRKY40 promoter activity.
Fig. 6 CaWRKY70 suppresses CaWRKY40 promoter activity in vivo and in planta. 

(a) In vivo Chromatin immunoprecipitation (ChIP) PCR assay shows CaWRKY70 binding to W-box 2 of CaWRKY40 promoter at shoot tissues of susceptible (JG62) chickpea under Foc1 stress. Diagram shows presence of W-boxes at CaWRKY40 promoter. (In) denotes input amplified from pre-cleared chromatin samples. Arrow indicates position of the primers. Plus (+) and minus (−) signs indicate anti-CaWRKY70 antibody and pre-immune sera immunoprecipitated chromatin. (+ 1) indicates transcription start site (TSS). 

(b) In silico molecular docking of CaWRKY70 and W-box 2 containing CaWRKY40 promoter DNA of cluster 3. 

c, d Electrophoretic mobility shift assay (EMSA) shows in vitro histidine tagged WRKY70 binding at pWRKY40 W-box 2. Approximately, 200 ng of WRKY70-His protein specifically binds at W-box 2 (−217 to −245 bp upstream of TSS). Plus (+) and minus (−) signs indicate presence or absence of specific components. BP indicates bound probe and FP represents free probe. Box indicates W-box 2. Asterisk (*) indicates mutated W-box. The experiment was repeated twice with similar results. 

e CaWRKY70 mediated trans-inhibition of CaWRKY40 promoter activity. p35S::CaWRKY70 and pWRKY40:YFP constructs were co-transfected in protoplasts obtained from Nicotiana tabacum cv. Xanthii (Brad) cell suspension culture. mCherry was used as transformation marker. Scale bar = 10 μm. 

(f) CaWRKY70 reduces GUS expression driven by CaWRKY40 promoter in tobacco leaf discs. Diagram shows constructs used for Agrobacterium mediated transient co-infiltration experiment. Plus (+) and minus (−) signs indicate presence or absence of specific vehicles.
CaWRKY40 positively regulates CaMPK9 promoter activity
Recent report shows that higher activation of CaMPK9 in resistant chickpea phosphorylates CaWRKY40 under Fusarium oxysporum (Foc1) stress [43]. We further anticipated that phosphorylated CaWRKY40 positively regulates CaMPK9 expression via feed-back mechanism in resistant chickpea on Foc1 challenge. Here, increased expression of CaMPK9 transcript in resistant chickpea also suggests its positive regulatory role in the defense activation against Foc1. Transcript level was found to be ~ 3.0-fold higher in resistant chickpea plants over control treatment. By contrast, susceptible chickpea plants show sharp downregulation of CaMPK9 transcript upon Foc1 challenge (Additional file 1: Figure S5). ChIP-PCR data support in vivo association of CaWRKY40 with W-boxes at CaMPK9 promoter in resistant genotype plants upon exposure to Foc1 (Fig. 7a). EMSA was used to establish in vitro binding of recombinant WRKY40 protein at CaMPK9 promoter DNA. Binding reactions were performed using 200 ng purified 6× histidine-tagged WRKY40 protein. Result shows the formation of sharp bound complexes after incubation of recombinant WRKY40 protein and labelled W-boxes. The complexes were competed out using 20 (for W-box 1) or 50 (for W-box 2) molar excess cold competitors (Fig. 7b, c). To ascertain the specific role of CaWRKY40 in CaMPK9 promoter activation, W-box-specific deletion constructs were generated and stably transformed into tobacco genome by Agrobacterium mediated gene transfer method (Fig. 7d). Integration of the CaMPK9 promoter fragments within tobacco genome was further confirmed by genomic PCR. Result shows sharp amplification of CaMPK9 promoter deletion derivatives (Fig. 7e). CaMPK9 promoter activity was further monitored in the presence or absence of specific effector constructs that constitutively express CaWRKY40. Reduction in the GUS activity was highest when both W-boxes were deleted (Fig. 7f). However, deletion of a single W-box results in mild reduction of the GUS activity in effector construct-infiltrated setup. These results suggest that CaWRKY40 binds at CaMPK9 promoter via both W-box 1 and W-box 2, which in turn positively modulates CaMPK9 expression.

Physical interaction between CaWRKY70 and CC-NB-ARC-LRR protein suppresses cell death in chickpea
The present group has recently established that physical interaction between RPP2-like CC-NB-ARC-LRR (CC-NLR) protein and CaWRKY64 triggers in planta ectopic cell death [30]. To further investigate whether CaWRKY70 similarly influences cell death signaling by CC-NLR protein, we tested their in planta interaction through bimolecular fluorescence complementation (BiFC) assay using Nicotiana benthamiana leaves. Results demonstrate that CaWRKY64 and CaWRKY70 tagged to C-terminus of YFP (cYFP) interact with full-length CC-NLR protein fused to N-terminus of YFP (nYFP). Result shows that reconstitution of YFP signal was observed in the nucleus (Fig. 8a). However, no such interaction was detected with control vector i.e., WRKY70-cYFP+nYFP. After testing their potential interaction between CaWRKY70 and CC-NLR protein, we were curious to check the effect of CaWRKY70 interaction on CaWRKY64 and CC-NLR protein mediated cell death phenomenon. Thus, we co-expressed HA tagged CaWRKY64, CaWRKY70 and myc tagged CC-NLR protein in chickpea leaves by Agrobacterium. The infiltrated leaves were further subjected to histochemical DAB and trypan blue staining. DAB staining shows that co-expression of epitope tagged CaWRKY64 and CC-NLR protein results in high levels of H2O2 accumulation in infiltrated chickpea leaves, which is suppressed upon CaWRKY70 expression (Fig. 8b). Similarly, trypan blue staining also depicted that CaWRKY70 effectively inhibits the retention of blue colouration and cell death in chickpea leaves when myc tagged CC-NLR protein and HA tagged CaWRKY64 were co-expressed (Fig. 8b). Next, we performed co-immunoprecipitation (Co-IP) to show physical interaction between myc epitope tagged NB-ARC domain of CC-NLR protein and CaWRKY70. Proteins were transiently co-expressed in N. benthamiana leaves by Agrobacterium. Reciprocal Co-IP analyses detected that CaWRKY70 and myc-NB-ARC proteins were co-precipitated by anti-myc and anti-WRKY70 antibodies, respectively. Immunoblotting was performed with anti-WRKY70 and anti-myc antibodies. Similarly, input samples show the presence of both CaWRKY70 and myc-CC-NLR proteins after probed with anti-WRKY70 and anti-myc antibodies (Fig. 8c). Effect of CaWRKY70 interaction on CC-NLR and CaWRKY64 mediated DNA binding has been further tested by in vitro EMSA experiment. Our group has previously shown that CC-NLR protein stimulates in vitro DNA binding of epitope tagged CaWRKY64 protein at CaEDS1 promoter [30]. This binding was found to be reduced by the addition of an increasing amount of recombinant his-tagged WRKY70 protein (Fig. 8d). Together, we establish that physical interaction between CaWRKY70 and CC-NLR protein negatively regulates cell death signaling in chickpea.

Discussion
WRKY70 transcription factor is well characterized as an important transcriptional modulator of SA mediated signal transduction pathways in Arabidopsis and wheat [39, 46]. AtWRKY70 overexpressing Arabidopsis plants demonstrated enhanced resistance to biotrophic pathogens Pseudomonas syringae and Erysiphe chichoracearum
However, the plants showed hyper-susceptibility to necrotrophic fungus *Alternaria brassicicola* [9]. Contrastingly, Atwrky70 mutants displayed susceptibility towards *B. cinerea* infection [47]. In wheat, *TaWRKY70* positively regulates defense against stripe rust pathogen *Puccinia striiformis* f. sp. *tritici* [46]. Both *Arabidopsis*
Fig. 8 CaWRKY70 attenuates R-protein signaling in chickpea. 

**a** BifC assay for physical interaction between CC-NLR-nYFP and CaWRKY70-cYFP. (nYFP + CaWRKY70-cYFP) and (CC-NLR-nYFP + CaWRKY64-cYFP) interactions were used as negative and positive controls, respectively. N denotes nucleus. 

**b** H2O2 accumulation and cell death in chickpea leaves. Myc-CC-NLR, HA-WRKY64 and CaWRKY70 were transiently co-expressed in chickpea leaves by *Agrobacterium*. Infiltrated chickpea leaves were subjected to DAB and trypan blue staining. 

**c** Co-immunoprecipitation of c-myc-NB-ARC and CaWRKY70 after *Agrobacterium* mediated transient expression in *N. benthamiana* leaves. Arrow indicates the immunoprecipitated protein band of CaWRKY70. Asterisks show non-specific bindings. 

**d** His-WRKY70 binding at W-box of chickpea *EDS1* promoter by EMSA. Dotted lines represent W-box. BP and FP indicate bound and free probes, respectively. In **b**, **c** and **d**, plus (+) and minus (−) signs indicate presence or absence of specific components.
Arabidopsis mutants inhibit subset of JA-responsive genes [9]. Enhanced WRKY70 promotes SA-responsive genes expression and CaWRKY70 [55]. However, JA treatment was found to be ineffective AtWRKY70 exhibited SA hyperaccumulation and subsequent induced transcript, whereas mutant plants AtWRKY70 in SA signaling demonstrate reduced level of SA treatment [9]. SA signaling was also found to be induced almost 30-fold at 2 h post-treatment in susceptible plant compared to the resistant one (Fig. 1). Likewise, AtWRKY70 expression was also found to be induced almost 30-fold at 2 h post-SA treatment [9]. SA signaling invokes positive inducer of SAR development and CaWRKY70 in chickpea shoots upon Foc1 infection. SA response is common for induction of several WRKY genes involved in distinct stages of the SAR activation in plants [50, 51]. SA treatment appears to be a positive inducer of SA development and CaWRKY70 expression in chickpea. Its mRNA level increases approximately 5-fold after 6 h of SA application over control treatment in susceptible plant compared to the resistant one (Fig. 1). Likewise, AtWRKY70 expression was also found to be induced almost 30-fold at 2 h post-SA treatment [9]. SA signaling invokes AtWRKY70 expression in young and senescing leaves. Our result shows that CaWRKY70 expression was highest at shoot tissues of chickpea (Fig. 1). Constitutive expression of bacterial salicylate hydroxylase NahG removes free SA and the subtle induction of AtWRKY70 transcript [52]. Arabidopsis mutants eds1, pad4 and npr1 compromised in SA signaling demonstrate reduced level of AtWRKY70 transcript, whereas mutant plants edr1, cpr5 and acd11 exhibited SA hyperaccumulation and subsequent induction of AtWRKY70 [53, 54]. SA and JA are two antagonistic signaling molecules that influence plant defense [55]. However, JA treatment was found to be ineffective for CaWRKY70 transcript induction. In most cases, WRKY70 promotes SA-responsive genes expression and inhibits subset of JA-responsive genes [9]. Enhanced expression of AtWRKY70 in coi1 mutant suggests that JA-responsive factor represses AtWRKY70 expression depending on endogenous JA levels [9]. Although, CaWRKY70 suppresses the expression of SA biosynthesis and signaling genes at shoot tissues of transgenic chickpea upon Foc1 infection (Figs. 4 and 5). ICS1 and PAL expression positively influences activation of SA signaling pathways and phenyl propanoid biosynthesis pathways leading to the production of anti-microbial secondary metabolites that protects chickpea and tomato plants from nematode penetration and Fusarium oxysporum infection, respectively [56, 57]. Our previous transcriptomic and metabolite analyses also revealed induction of CaICS1 and CapAL transcripts and associated SA accumulation in resistant (WR315) chickpea after Foc1 inoculation [38]. By contrast, CaICS1, CaPAL expression and SA concentrations were significantly depleted at shoot tissues of CaWRKY70 overexpressing chickpea under Foc1 stress (Fig. 4). EDS1 and PAD4 are two such important regulatory components of SA biosynthesis in plants upon pathogen stress [44, 58]. In chickpea, we previously observed constant induction of CaEDS1 and CaPAD4 transcripts at both shoot and root tissues of resistant genotypic plant under Foc1 infected condition [38]. However, CaEDS1 and CaPAD4 transcripts were downregulated at CaWRKY70 overexpressing chickpea shoot tissues in response to Foc1 infection (Fig. 5). This may strengthen negative regulatory role of CaWRKY70 in systemic defense reactions. SA signaling in plants depends on the activation of TGA transcription factors and NPR1. These two transcriptional modulators synergistically control expression of the two critical SA marker genes i.e., PRI and PR5. The effective induction of CaTGA1 and CaTGA6 mRNAs were observed at shoot tissues of Foc1 infected resistant chickpea, whereas susceptible plants were unable to stimulate such mRNA accumulation [38]. CaWRKY70 expresses at shoot tissues of susceptible chickpea after Foc1 inoculation and its overexpression in resistant chickpea plants markedly reduces CaTGA1 and CaTGA6 transcripts accumulation (Figs. 1 and 5), which indicates to the impairment of conserved SA signaling in susceptible chickpea. CaNPR1, CaPRI and CaPR5 transcripts follow the same pattern of Foc1 induced downregulation at shoot tissues of CaWRKY70 overexpressing chickpea than control transgenics (Fig. 5). Importantly, CaWRKY70 represses the expression of SA and JA-marker genes i.e., CaPR1, CaPR5 and CaDefensin that promotes susceptibility in transgenic chickpea (Fig. 5). SA-mediated repression of JA-responsive gene expression is governed by cytosolic NPR1 [59]. WRKY70 controls JA-repressors based on cytosolic modification of NPR1 protein [59]. CaWRKY70-mediated suppression of CaNPR1 might play negative role in SA-responsive CaPRI, CaPR5 and JA-induced CaDefensin gene expression in chickpea (Fig. 5). PDF1.2 transcript in Atwrky70 mutant was found to be low which enhanced upon B. cinerea infection. However, in CaWRKY70 overexpressing chickpea, CaDefensin expression has been downregulated under Foc1 infection [47]. Low levels of SA promote PR genes expression in chickpea, whereas high concentrations inhibit both SA and JA signaling pathways. Thus, CaWRKY70 acts as an integrator of SA and JA responses in the regulation of chickpea defense responses to Foc1. Mutual antagonism and interaction between SA and JA pathways are common regulatory steps for CaWRKY70 expression and its activation that governs wilt-disease resistance phenomenon in chickpea. Our previous study revealed that differential SAR
induction in resistant and susceptible chickpea plants are analogous to the SA dependent gene expression and here, we convey mechanism of its attenuation. Hence, our present study explains the complexity of SA biosynthesis, signaling and its feed-back inhibition by CaWRKY70 that control Foc1 resistance/susceptibility in two contrasting chickpea accessions, respectively.

Regulatory sequences and DNA-binding activity of WRKY family members remarkably govern various cellular and stress responsive phenotypes in plants [60]. DNA binding role of CaWRKY70 is not an unusual phenomenon since, it is a nuclear localized protein (Additional file 1: Figure S1). In Foc1 infected chickpea, CaWRKY70 inhibits the expression of CaWRKY40 signaling genes i.e., CaWRKY33 and CaMPK9 (Figs. 5 and 7). Recent finding suggests CaMPK9 interaction and phosphorylation provide stability to CaWRKY40 protein in chickpea upon Foc1 infection [43]. CaWRKY40 mediated upregulation of CaMPK9 expression was suppressed in CaWRKY70 overexpressing chickpea. Shared transcriptional regulation of AtWRKY18, AtWRKY40 and AtWRKY60 adjusts abscisic acid (ABA) signaling mediated abiotic stress responses in plants [61]. We found that CaWRKY70 binds at CaWRKY40 cis-elements and represses its activity (Fig. 6). It is interesting to note that CaWRKY40 positively regulates the CaMPK9 promoter activation which has been established by both in vivo and in planta experiments (Fig. 7). CaMPK9 upstream elements deletion study also revealed its role in the modulation of promoter activity. CaWRKY70 activated transcription of CaWRKY54 gene in transgenic chickpea (Fig. 5). These two transcription factors co-ordinately function as negative regulators of leaf senescence, stomatal closure, and osmotic stress tolerances in Arabidopsis [41, 62]. Although, our study demonstrated that CaWRKY70 and CaWRKY54 co-operatively contribute to Foc1 susceptibility in chickpea. CaWRKY70 mediated promoter modulation suggests bidirectional transcriptional regulation. Therefore, CaWRKY70 mediated inhibition of appropriate immune signaling in chickpea accomplishes through its direct and indirect negative regulatory influence on defense genes expression under Foc1 stress condition.

Transcriptional responses behind SAR activation by WRKY proteins were previously established in Arabidopsis [63, 64]. However, the mechanism of its repression is not known. Present study shows that deactivation of SAR in pathogen-free systemic tissues of chickpea is mediated by CaWRKY70 upon Foc1 infection. Repressor activity of AtWRKY70 on SARD1 gene expression regulates the balance between growth and defense in Arabidopsis [19, 20]. Such regulatory functions are yet to be demonstrated in chickpea. CaWRKY70 is transcribed and translated at susceptible chickpea shoots upon Foc1 infection (Fig. 1). Since, SAR activation was prominent at shoot tissues of resistant chickpea [38], we monitored CaWRKY70 over-expression effect in this background. The overall suppression of SA signal transduction network confers Foc1 susceptibility in CaWRKY70 overexpressing chickpea (Figs. 4 and 5). Although, it is tempting to monitor CaWRKY70 knock-down effect in susceptible chickpea background as future attempt.

SA has immense roles in plant immunity, including resistance gene signaling. EDS1 is a well-known genetic regulator of SA production, resistance gene functioning and cell-death [65]. RPP2-like CC-NB-ARC-LRR protein and CaWRKY64 mount EDS1 dependent ectopic defense activation and cell-death in chickpea [30]. EDS1 mostly confers resistance as part of the TIR-NB-LRR signaling [66]. Based on our protein-protein interaction studies, we establish that physical interaction between CaWRKY70 and chickpea RPP2-like CC-NLRR protein effectively suppresses ROS accumulation and cell-death induction in planta (Fig. 8). RPP4-mediated resistance response against Hyaloperonospora parasitica Emoy2 was partially reduced in Atwrky70 mutants [32]. On the other hand, RPP7-triggered defense reaction was not affected in this mutant background. Thus, R-protein mediated defense response pathway is positively correlated with AtWRKY70 functioning in Arabidopsis, whereas it is inhibited by CaWRKY70 in chickpea might contradict WRKY70 dependent R-protein signaling in general. CaWRKY70 mediated inhibition of ROS accumulation in transgenic chickpea root, stem and leaves also support our interpretation (Fig. 3). Although, reduction in oxidative bursts signaling and SA production does not fully correlate with the increased ion-leakage and fungal biomass accumulation in CaWRKY70 overexpressing chickpea root than control transgenics (Figs. 2 and 3). We reasoned that electrolyte leakage is associated with membrane damage due to higher colonization of Foc1 biomass, whereas ROS accumulation induced cell death is a defense phenomenon that inhibits in planta Foc1 growth. Importantly, less cell death promotes higher Foc1 colonization in CaWRKY70 overexpressing chickpea roots. On the other hand, complete inhibition of EDS1 signaling not only affects SA induction, but also cell-death promotion (Figs. 5 and 8). In summary, CaWRKY70 functions as nodal suppressor that includes fundamental defense regulators like, SA response and EDS1 into R-protein mediated signal transduction pathways highlighted in chickpea upon Foc1 stress (Fig. 9).

Conclusions

Finally, this study provides information which may fill the gaps between already available knowledge about CaWRKY70 mediated transcriptional control of downstream defense signaling pathways. Promoter occupancy
and protein-protein interaction play crucial roles for suppression of chickpea defense to Foc1. Interpretation of our findings may be translated and recapitulated for serial examination of multiple layers of defense signaling in chickpea. Repressor role of CaWRKY70 in modulating ROS homeostasis, SA biosynthesis and signaling is an interesting finding. Interconnection between several signaling cues in turn confer resistance against Foc1 in definite ways depending on time point of infection, duration, and severity. Notably, spatiotemporal expression patterns of CaWRKY70 mediated immune signaling elements renovates our apprehension. Thus, present study is useful to develop strategies for protecting chickpea from *Fusarium* wilt disease.

**Methods**

**Plant materials and growth conditions**

Experiments were performed using two different genotypes of chickpea (*Cicer arietinum* L.) i.e., JG62 (wilt susceptible) and WR315 (wilt resistant) obtained from Dr. Suresh C. Pande, International Crops Research Institute for the Semi-Arid Tropics (ICRISAT), Patancheru, Andhra Pradesh, India. Surface sterilized seeds of both genotypes were sown in the pots containing autoclaved mixture of soil-rite and soil under natural greenhouse conditions at 22 to 25 °C temperature, 70% relative humidity, 100 μmol m⁻² s⁻¹ light intensity and 16 h photoperiod. Pots were watered from bottom at every 2 days and supplemented with half strength Hoagland’s medium (TS1094, Hi-media Laboratories, Mumbai, India). *Nicotiana tabacum* L. cv. Samsun NN and *N. benthamiana* seeds were gifted by Dr. Nrisingha Dey, Institute of Life science, Bhubaneshwar, India. Seeds were surface sterilized and grown aseptically on MS medium at 24 °C temperature and 60% relative humidity with light intensity of 100 μmol m⁻² s⁻¹ under 16 h photoperiod.

**Fungal inoculation**

*F. oxysporum* f. sp. *ciceri* Race1 (Foc1) fungal strain was obtained from Dr. Suresh C. Pande, International Crops Research Institute for the Semi-Arid Tropics (ICRISAT), Patancheru, Andhra Pradesh, India. Sixteen-days-old chickpea plants were inoculated with Foc1 using sick-soil method according to the previously described method [34, 43].

**Hormone treatment**

For inducer treatments, 2 mM salicylic acid (SA, Hi-media), 100 μM ABA (Abscisic acid, Hi-media) and 50 μM JA (Jasmonic acid, Hi-media) was sprayed on greenhouse gown sixteen-days-old chickpea plants of both susceptible and resistant accessions. Leaf tissues were collected at 6 h of treatment for RNA isolation.

**RNA isolation and quantitative real-time PCR (qRT-PCR) analyses**

Shoot tissues of sixteen-days-old susceptible and resistant chickpea plants were collected and frozen in liquid nitrogen. Total cellular RNA was extracted from frozen sample using TRIZOL reagent (HiMedia). First strand cDNA synthesis was carried out from 5 μg total RNA using First Strand cDNA synthesis Kit (Thermo Scientific, USA) following manufacturer’s guideline. qRT-PCR was performed using Bio-Rad iCycler (Bio Rad iQ5) with SyBr green (Bio Rad). The reaction mix containing SyBr green qPCR Super Mix (2×) (Bio Rad), 25 ng cDNA, and 0.3 μM of forward and reverse primers (Additional file 3: Table S2). Chickpea Glyceraldehyde-3-phosphate dehydrogenase (*CaGAPDH*) expression was used as internal control. Normalized fold change levels for all the genes were calculated using the 2⁻ΔΔCt method [67].

**Subcellular localization**

For subcellular localization study, YFP gene was cloned in *BamHI/ SacI* site of pBI121binary plant transformation vector. Full-length *CaWRKY70* gene was PCR amplified and fused in frame to N terminal part of the yellow fluorescent protein (YFP) gene in pBI121 vector for preparation of the 35S:WRKY70-YFP construct. The vectors were mobilized into the competent
Agrobacterium tumefaciens strain GV3101. 35S: CaWRKY70-YFP and control 35S:YFP vectors were transiently transformed into onion epidermal cells by Agrobacterium. At 48 h post-agroinfiltration, epidermal cells were observed under a confocal microscope (Leica TCS SP2 AOBS system) to monitor localization patterns of fusion proteins under excitation and emission at 514 nm and 527 nm, respectively.

Molecular cloning
Full-length coding sequence of CaWRKY70 (GenBank Accession No. XM_004502763.3) was amplified from chickpea cDNA pool by reverse-transcriptase polymerase chain reaction (RT-PCR) using gene specific primers (Additional file 3: Table S2). Purified PCR amplicons were restriction digested with BamHI/ XhoI (Roche, Mannheim, Germany) and cloned in modified pBI221 vector containing short multi-cloning sites (MCS) region by replacing GUS gene. Cassette was gel excised after treatment with HindIII/ EcoRI (Roche, Mannheim, Germany) and cloned in MCS of binary plant expression vector pCAMBIA2301. Cloning was facilitated by restriction digestion, which is followed by sequencing of the full-length gene. Binary plant transformation vector containing CaWRKY70 gene was mobilized to Agrobacterium tumefaciens strain AGL-1. Empty pCAMBIA2301 vector was also transformed into Agrobacterium strain AGL-1 was used as vector control.

Chickpea transformation
Agrobacterium-mediated chickpea transformation was carried out as described by [68] with a modified rooting protocol [69]. Briefly, chickpea transformation was carried out with single cotyledon and half-embryo explant followed by infection with Agrobacterium strain AGL-1 harbouring empty pCAMBIA2301 vector and modified vector carrying CaWRKY70 gene. Multiple shoots were regenerated from the explants and elongated with 0.25 mg/ l IAA (Indole-3-acetic acid) for 10 days. The elongated shoots were transferred to rooting medium (1/2 MS salts, B5 vitamins, 1 mg/ l IBA and 20 g/ l sucrose) [70]. Finally, rooted plantlets were properly hardened, transferred to glasshouse and established in the pots. Empty pCAMBIA2301 vector generated plants were used as control transgenics whereas, CaWRKY70 gene carrying modified vector transformed plants were considered as overexpressing chickpea.

Disease intensity index
Disease intensity index was determined on control transgenic and CaWRKY70 overexpressing chickpea based on the development of foliar symptoms at 0, 3, 7 and 12 dpi, respectively. Incidence of foliar symptoms (I) was set at 0 to 1 scale and disease severity (S) rated on a 0 to 4 scale (0 - no wilting; 1 - less wilting; 2 - partial wilting; 3 - wilting; 4 - severe wilting). Disease intensity index (DII) was calculated as DII = (I × S)/4 [71].

Incidence of dead plants
Development of Foc1 infection on control transgenic and CaWRKY70 overexpressing chickpea in the greenhouse experiment was recorded as incidence of dead plants at 0, 3, 7 and 12 dpi. The percentage of incidence of dead plants from transgenic chickpea was measured using the formula i.e., percentage of incidence of dead plants = (total number of infected plants/ total number of plants assessed) × 100 [72].

Chlorophyll estimation
For chlorophyll extraction, one gram of control transgenic and CaWRKY70 overexpressing chickpea leaf samples were crushed with 2 ml dimethyl sulfoxide (DMSO): acetone (1:1 vol/ vol) mix. The samples were then kept in a refrigerator at 4 °C for 4 h. The samples were then centrifuged at 500 rpm for 5 min. Following this, supernatant was transferred to fresh 2 ml eppendorf tubes. The colour absorbance (A) of extracts was determined using Shimadzu UV 1800 spectrophotometer (Shimadzu Corporation, Kyoto, Japan) at 645 and 663 nm wavelength against the blank solvent containing 80% acetone. Chlorophyll A and B content was estimated based on [73].

DAB staining
H2O2 accumulation in treated CaWRKY70 overexpressing chickpea root, stem and leaves were visualized by 3, 3'-diaminobenzidine (DAB) staining, according to the method of [74]. The plant tissues were immersed in 1 mg/ ml DAB (3, 3'-diaminobenzidine) solution (pH 3.8) and vacuum infiltrated for 2 h followed by incubation of 8 h at room temperature. Chlorophyll was removed by incubating in 96% ethanol for overnight and photographed with a digital camera. DAB stained samples were oven dried for 24 h and crushed with sterile double distilled water to measure their intensities by spectrophotometer at 500 nm wavelength and water as blank.

Trypan blue staining
For trypan blue staining, lactophenol-trypan blue solution was prepared by mixing 10 ml lactic acid, 10 ml glycerol, 10 ml distilled water, 10 g of phenol and 10 mg of trypan blue. H2O2 treated and Foc1 infected control and transgenic chickpea root, stem and leaves were subjected to trypan blue staining. Plant tissues soaked in trypan blue solution were warmed in a boiling water bath for 1 min and cleared with saturated chloral hydrate solution (2.5 g chloral hydrate dissolved in 1 ml distilled water) for 10 min. Decolorized plant samples were
photographed. Trypan blue stained samples were dried and extracted with sterile double distilled water. The crude extracts were quantified using spectrophotometer at 500 nm wavelength against water as blank.

**Bacterial expression and protein purification**

PCR amplified full-length CaWRKY40 and CaWRKY70 genes were inserted into EcoRl/ XhoI site of pET28a(+) (Novagen, Germany) and transformed into *Escherichia coli* BL21 (DE3) cells. Recombinant WRKY40 protein purification was carried out as previously described by [42]. Histidine tagged WRKY70 protein purification was performed according to previously described method by [49]. Protein induction was carried out with 1 mM isopropyl-β-D-galactoside (IPTG) at 37 °C for 1 h with vigorous shaking (160 rpm) and cells were harvested by centrifugation at 8000 rpm 4 °C for 5 min. Hexa-histidine tagged WRKY70 protein was purified from cell lysate by Ni-NTA affinity chromatography (Qiagen).

**Antibody production**

Anti-WRKY70 polyclonal antibodies were raised in rabbits. Two rabbits were immunized for antibody production. Rabbits were injected on 5 occasions with recombinant hexa-histidine tagged WRKY70 protein of 1.5 mg concentrations at every 3-week intervals. Serum obtained from each immunized rabbit was tested 2-weeks after each injection. Pre-immune serum was collected from each animal. Approximately, 20–30 ml serum/ rabbit was obtained. The serum was affinity purified. Antibodies were used at a final dilution of 1:10,000 for immunoblotting experiments.

**Protein extraction and immunoblotting**

Total soluble protein extraction from sixteen-days-old control and Foc1 inoculated susceptible and resistant chickpea shoots were performed using an ice-cold protein extraction buffer (50 mM Tris–HCl pH 7.5, 100 mM NaCl, 1 mM DTT, 0.5% Triton X-100, 0.1% SDS and 10% glycerol) followed by the addition of protease inhibitor cocktail (ETD/A-free, Roche). Protein concentration was measured by Bradford assay [75]. Approximately, 20 μg of total soluble protein was separated by 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) gel and detected by western blotting with anti-WRKY70 polyclonal primary antibody and an anti-rabbit IgG conjugated to horseradish peroxidase secondary antibody (Sigma, A-6667).

**Conductivity measurement assay**

For conductivity measurement experiment, control transgenic and CaWRKY70 overexpressing chickpea roots were subjected to Foc1 infection at various times and washed thoroughly with water. 200 milligram control and Foc1 infected chickpea roots were incubated overnight in sterile tubes filled with 20.0 ml distilled water. Following this, electrical conductivity of water was measured using an electrolyte meter at indicated time points.

**Estimation of Foc1 biomass**

Amount of Foc1 biomass was measured according to the previously described method [76]. Genomic DNA isolated from Foc1 inoculated root tissues of control transgenic and CaWRKY70 over-accumulating chickpea was used as template for real-time PCR with 5.8S rDNA primers listed in (Additional file 3: Table S2).

**Measurement of relative water content (RWC)**

RWC of vehicle transgenic and CaWRKY70 overexpressing chickpea plants were determined by weighing method upon control treatment and Foc1 infection [38].

**SA estimation by high performance liquid chromatography (HPLC)**

SA concentrations were determined by HPLC (Shimadzu, Japan) provided with two LC-10 pumps and a UV detector system SPD-10A [77]. Total SA was extracted from 200 mg control and Foc1 treated shoot tissues of transgenic chickpea. The samples were dissolved separately in 200 μl of running buffer (0.2 M NaOAc, pH 5.2, and 10% methanol) and injected in a C-18 HPLC column (4 μm, 250 × 4.6 mm, Phenomenex, USA). A two-pump linear gradient system was used for separation of methanolic plant extracts i.e., pump A contains 1% acetic acid and pump B was filled with acetonitrile. SA detection was carried out at 254 nm wavelength, 30 °C temperature with a flow rate of 0.8 ml/min. The data obtained were combined using Shimadzu Class VP series software. Samples were identified according to their respective retention time (Rt) of peaks and quantity was calculated in mg g⁻¹ FW based on area of the peak and the values obtained for standard used.

**In silico DNA-protein interaction study**

The sequence of putative CaWRKY70 protein was retrieved from NCBI (National Centre for Biotechnology Information). Template search and three-dimensional structure prediction of CaWRKY70 (PDB ID: c2aydA) was performed by homology modelling using Phyre2 tool (http://www.sbg.bio.ic.ac.uk/phyre2) [78]. Validation of the generated model was further carried out by Ramachandran plot analysis using RAMPAGE server (http://mordred.bioc.cam.ac.uk/~rapper/rampage.php) [79]. Molecular docking analysis of CaWRKY70 with W-Box DNA was done by HADDOCK (High Ambiguity Driven protein-protein Docking, www.haddocking.org) web server [80]. B DNA model with W-Box element (TGAC) was constructed using 3D-DART webserver, which
corrects the nucleotide according HADDOCK specification [81]. Protein and DNA models were subjected to molecular docking by uploading their respective PDB files using easy interface at the HADDOCK server. The W-Box element of modelled DNA and the “WRKYGQK” amino acid sequences present in CaWRKY70 transcription factor were selected as the active residues for docking. Passive residues were automatically selected surrounding the active residues. Finally, docked structure was illustrated and visualized using UCSF Chimera (https://www.cgl.ucsf.edu/chimera/) [82].

Chromatin immunoprecipitation (ChIP) assay
ChIP assay was performed according to the method previously described by [83] showing CaWRKY70 and CaWRKY40 binding at CaWRKY40 and CaMPK9 promoters via W-boxes, respectively. ChIP primers are listed in the (Additional file 3: Table S2).

Electrophoretic mobility shift assay (EMSA)
In vitro DNA binding activity of 6X histidine tagged WRKY70 protein was performed using EMSA experiments. To prepare the DNA probes for EMSA, equimolar concentration of each sense and antisense oligonucleotide of respective DNA duplexes were mixed in a reaction buffer containing 40 mM Tris–HCl; pH 7.5, 20 mM MgCl$_2$, 50 mM NaCl. The reaction mix was heated at 95°C for 10 min and slowly cooled down to room temperature for annealing. DNA duplexes were run on 7% PAGE (polyacrylamide gel electrophoresis) and gel-purified, followed by end labelling with $\gamma^{32}$P and T$_{4}$-polynucleotide kinase (NEB). The labeled probes were then purified using QIAquick Nucleotide Removal Kit (Qiagen). The binding assay was performed using His-tagged WRKY70 protein. Typically, the binding reaction contained 100–200 ng of purified protein, 10 ng of double-stranded synthetic oligonucleotides end-labeled with $\gamma^{32}$P in a binding buffer containing 20 mM HEPES, pH 7.5, 100 mM KCl, 0.2 mM EDTA, 1 mM DTT, 2 mM MgCl$_2$, and 1 μg of poly dl-dC. The protein was pre-incubated in binding buffer for 5 min prior to the addition of probe for eliminating the risk of non-specific binding and the reaction mix was incubated for another 30 min at room temperature. The complexes were resolved in 0.5X TBE-5% PAGE at (4°C, 100 V) for about 2 h. After electrophoresis, gels were dried and exposed to phosphorscreen for imaging in a phosphorimager (Typhoon, GE Healthcare). For competition assays, unlabelled duplexes were added during the incubation stage.

Protoplast transfection and trans-inhibition assay
Protoplasts from *Nicotiana tabacum* cv. Xanthi (Brad) cell suspension culture was isolated and electroporated according to previously described method by [84]. For trans-inhibition assay, *CaWRKY40* promoter region was cloned between HindIII/BamHI site by replacing CaMV35S promoter in YFP containing pBI121 vector to obtain pWRKY40:YFP. Next, p3SS:WRKY70 and pWRKY40:YFP vectors were co-transfected into protoplasts and incubated in dark for 48 h. Confocal microscopy was performed to monitor the promoter activity.

GUS assay
Histochemical GUS staining of transgenic tobacco seedlings were performed according to the protocol [85]. Briefly, the tissues were incubated in GUS staining solution containing 1 mM X-Gluc (Duchefa Biochemie, Netherlands), 100 mM sodium phosphate (pH 7.0), 2 mM potassium ferricyanide, 2 mM potassium ferrocyanide, 10 mM EDTA, and 0.1% Triton X-100 under dark conditions at 37°C for overnight (16 h). After staining, tissues were de-stained in 75% ethanol and photographed with a digital camera. CaWRKY40 (LOC101512877) and CaMPK9 (LOC101496681) promoter activity was monitored through the GUS expression analysis. CaWRKY40 and CaMPK9 promoter was inserted between HindIII/BamHI site of pBI121 vector by replacing CaMV35S promoter upstream to the GUS gene. Plasmids were transformed into *A. tumefaciens* strain LBA4404. p3SS:WRKY70 or p3SS:WRKY40 (effector constructs) and pWRKY40:GUS or pCaMPK9:GUS (reporter constructs) were co-infiltrated into the ventral surface of *N. tabacum* leaves and subjected to GUS activity assay at 2 days post-infiltration. 100 milligram of agro-infiltrated tobacco leaf discs were collected in a 1.5 ml micro-centrifuge tube and crushed with liquid nitrogen into fine powder. Five hundred microliter of GUS extraction buffer containing 50 mM NaHPO4, pH- 7.0, 10 mM 2-mercapto ethanol, 10 mM Na2EDTA, 0.1% SDS and 0.1% triton X-100 was added to grinded sample. The mixture was then centrifuged at 13,000 rpm for 15 min at 4°C and the cleared supernatant was collected. Data was normalized to protein concentration as measured by Bradford method [75]. Ten microgram crude protein extracts were mixed with 100μl of GUS assay solution 2 mM 4-methyl umbelliferyl-d-glucuronide (4-MU) in extraction buffer. The reaction was carried out at 37°C for 60 min and stopped by addition of 0.2 M Na2CO3. One hundred microliter re-action mixture was used to measure the fluorescence of GUS enzymatic activity using spectro-fluorometer (Hitachi, F-7000) under excitation and emission at 365 nm 455 nm.

*Agrobacterium* mediated transient infiltration
*Agrobacterium tumefaciens* GV3101 cells harbouring plasmids were grown overnight at 28°C in 25 ml Luria-
Bertani-broth with antibiotics. Next day, bacterial cells were pelleted by centrifugation at 8000 rpm and dissolved in 20 ml of induction medium (50 mM MES, pH 5.6, 0.5% (W/V) Glucose, 1.7 mM NaH2PO4, 20 mM NH4Cl, 1.2 mM MgSO4, 2 mM KCl, 17 μM FeSO4 and 70 μM CaCl2) in presence of 200 μM acetosyringone. The solution was incubated at 28 °C for 4 h in shaker incubator with rotation speed 160 rpm. After incubation, bacterial cells were pelleted, resuspended in 20 ml of agroinfiltration medium (10 mM MES and 10 mM MgCl2, pH 5.6) for 3 h with 100 μM acetosyringone at OD600 (0.4) and then infiltrated into the ventral surface of fully expanded 6-weeks-old N. benthamiana leaves. In case of chickpea leaves, transient infiltration was carried out using bath-sonication method as previously described by [30].

BiFC assay
For BiFC studies, full length CDS of the CaWRKY70 and CaRPP2-like CC-NC-ARC-LRR gene (GenBank accession XM_012712097.1) was cloned between BamHI/ Sal I site of pSPYCE and pSPYNE vectors to generate CaWRKY70-YFP C-ter. and CC-ARC-ARC-YFP N-ter. Full-length CaWRKY64 gene (GenBank accession XM_004489016.3) was cloned within Spel/ Xhol site of pSPYCE vector. The control and fusion plasmids were transformed into Agrobacterium strain GV3101. Agrobacterium cells carrying control and fusion plasmids were co-infiltrated into the abaxial-side of bacterium cells carrying control and fusion plasmids Agrobacterium transformed into pSPYCE vector. The control and fusion plasmids were the samples and incubated for additional 3 h at 4 °C. The bound proteins were separated by centrifugation at 14, 000 rpm for 20 min at 4 °C. Next, the beads were washed three times with ice cold wash buffer (50 mM HEPES pH 7.5, 150 mM NaCl, 10 mM EDTA and 0.1% Triton X-100 and protease inhibitor cocktail). Bound proteins were eluted from the beads by 100 µl 1X laemmlisi sample buffer containing 250 mM Tris-HCl (pH 6.8), 10% SDS (W/V), 0.5% bromophenol blue (W/V), 50% glycerol and 50 mM DTT in a boiling water bath for 10 min. Western blotting was performed with anti-c-Myc and anti-WRKY70 antibodies, which is followed by addition of the horse serum peroxidase conjugated secondary antibody.

Statistical analyses
For statistical differences, Student’s t-test was performed at a significance level of p < 0.05 come next to multiple comparison of means by Tukey’s post-hoc test.

Supplementary information
Supplementary information accompanies this paper at https://doi.org/10.1186/s12870-020-02527-9.

Additional file 1 Figure S1. Subcellular localization of control YFP and CaWRKY70-YFP after transient expression in onion epidermal cells by Agrobacterium. Bars represent 250 μm. Red and white arrows indicate nucleus and cytoplasm, respectively. Figure S2. PCR amplification of CaWRKY70 gene cloned in pCAMBA2301 vector after resolved on 1% agarose gel. Lane-M denotes DNA molecular weight marker in kilobases (kb). Lane-1, 2, and 3 show positive clones. Red arrows indicate CaWRKY70 PCR amplicons. Figure S3. Homology modelling and Ramachandran plot calculation of CaWRKY70 protein. a Predicted structure of CaWRKY70 showing five anti-parallel β-strands. d Predicted structure of CaWRKY70 showing five anti-parallel β-strands. e Qualitative assessment of stereo chemical and spatial arrangement of amino acids present on CaWRKY70 protein using RAMPAGE server. Figure S4. MUG assay quantitation of CaWRKY70 mediated reduction in pWRKY40-GUS activity. Plus (+) and minus (−) signs show presence or absence of the specific components. Error bars indicate ±SD (n = 5). Asterisks (*) indicate values are different from one another in statistically significant manner as determined by Student’s t test (**P < 0.001). Figure S5. CaMPK9 transcript accumulation in susceptible JG62 and resistant WR315 chickpea shoots under control treatment (0 dpi) and Foc1 infection (7 dpi) by real-time PCR. CaGA20DH mRNA level was used as internal control. Fold change was calculated relative to the control treatment. Error bars indicate ±SD of three biological replicates. Student’s t test was performed to determine its significance level as compared to the control treatment, **P < 0.01 and ***P < 0.001.

Additional file 2 Table S1. Statistics of the top five clusters of CaWRKY70-DNA complex generated by HADDOCK server.

Additional file 3 Table S2. List of primers used in this study.

Abbreviations
ABA: Abscisic acid; ARC: Apaf-1; Resistance and CED4 (Caenorhabditis elegans) cell death 4 protein; BIFC: Bimolecular Fluorescence Complementation; Ca: Cicer arietinum L.; CBBS: Coomassie Brilliant Blue; CC: Colleol; ChIP: Chromatin Immunoprecipitation; DNA: Deoxyribonucleic acid; EDS1: Enhanced Disease Susceptibility 1; EMSA: Electrophoretic Mobility Shift Assay; ET: Ethylene; HA: Haemagglutinin; HR: Hypersensitive response; ICS1: Isochorismate Synthase 1; IA: Jasmonic acid; LRR: Leucine rich repeat; MPK: Mitogen Activated Protein Kinase; mRNA: Messenger RNA; NB: Nucleotide binding; NLR: Nucleotide binding oligomerization domain leucine rich repeat; NPR1: Non-expressor of PR1; PAP4: Phytoalexin Deficient 4; PAGE: Polyacrylamide Gel Electrophoresis; PAL: Phenylalanine ammonia-
lyase; PCR: Polymerase chain reaction; PDB: Protein Data Bank; PR1: Pathogenesis Related 1; qRT-PCR: Quantitative real-time PCR; RNA: Ribonucleic acid; RPP2: Recognition of *Peronospora Parasitica* 2; SA: Salicylic acid; SDS: Sodium Dodecyl Sulfate; VFP: Yellow Fluorescent Protein

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### Authors’ contributions

JC and SD conceived and designed research. JC, SS, PG and AJ performed the experiments. JC, SS and SD analyzed the data and JC, PG, AJ and SD wrote the manuscript. The authors read and approved the final manuscript.

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### Availability of data and materials

No large-scale global data or any database was created for the study. All data generated during this study are included in this article and additional files.

### Ethics approval and consent to participate

Not applicable.

### Consent for publication

Not applicable.

### Competing interests

The authors declare that they have no competing interests.

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