Characterization and functional analysis of GhWRKY42, a group IId WRKY gene, in upland cotton (Gossypium hirsutum L.)

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Abstract

Background: WRKY transcription factors (TFs) participate in various physiological processes of plants. Although WRKY genes have been well studied in model plants, knowledge of the functional roles of these genes is still extremely limited in cotton.

Results: In this study, a group IId WRKY gene from cotton, GhWRKY42, was isolated and characterized. Our data showed that GhWRKY42 localized to the nucleus. A transactivation assay in yeast demonstrated that GhWRKY42 was not a transcriptional activator. A β-glucuronidase (GUS) activity assay revealed that the promoter of GhWRKY42 showed fragment deletion activity in Nicotiana tabacum and was mainly expressed in the roots, stems and leaves of ProGhWRKY42::GUS transgenic Arabidopsis plants. Quantitative real-time PCR (qRT-PCR) analysis indicated that GhWRKY42 was up-regulated during leaf senescence and was induced after exposure to abiotic stresses.

Conclusions: Constitutive expression of GhWRKY42 in Arabidopsis led to a premature aging phenotype, which was correlated with an increased number of senescent leaves, reduced chlorophyll content and elevated expression of senescence-associated genes (SAGs). In addition, virus-induced gene silencing (VIGS) was used to silence the endogenous GhWRKY42 gene in cotton, and this silencing reduced plant height.

Keywords: GhWRKY42, Arabidopsis, Leaf senescence, VIGS, Cotton, Plant height

Background

Plants are constantly challenged by various factors that affect plant growth and development throughout their life cycle. To combat these challenges, some responsive genes, including WRKY transcription factors (TFs), are induced to help plants adapt through physiological and morphological changes [1]. WRKY TFs are plant-specific proteins and constitute one of the largest TF families in plants [2]. WRKY TFs share the common feature of a highly conserved WRKY domain that consists of the peptide sequence motif WRKYGQK at the N-terminus and a zinc-finger-like motif at the C-terminus [3]. WRKY TFs have one or two conserved WRKY domains, and these domains contain a Cx4-5Cx22-23HxH or Cx7Cx23HxC zinc-finger-like motif. Based on the number of conserved WRKY domains and the structural characteristics of the zinc-finger-like motifs, WRKY TFs can be categorized into group I, group II or group III. Group II can be further divided into subgroups IIa, IIb, IIc, IId and IIe [3–7]. WRKY TFs can recognize and bind to the W-box sequences [TTGAC(C/T)] in the promoter region of target genes to participate in regulatory networks [8].

In plants, WRKY TFs are mainly involved in defense responses, trichome development, plant growth and development and leaf senescence [9]. Various TFs are involved in modulating leaf senescence, and 1533 TFs have been identified via leaf senescence transcriptome analyses in Arabidopsis [10, 11]. WRKY TFs are quantitatively important members of those TFs involved in leaf senescence [11]. In Arabidopsis, AtWRKY6 is associated
with the senescence process by targeting the promoter of the SIRK gene, which likely encodes a signaling component related to leaf senescence [12]. AtWRKY45 was recently reported to interact with the DELLA protein RGA-LIKE1 (RGL1) and to directly target the SAG12, SAG13, SAG113 and SEN4 genes, to positively modulate leaf senescence via the gibberellic acid-mediated signaling network [13]. In rice, OsWRKY24 promotes senescence in transgenic rice plants by binding to the promoter of OsMT1d to repress ROS scavenging [14]. OsWRKY23 is markedly increased during dark-induced leaf senescence, and OsWRKY23-overexpressing lines can accelerate leaf senescence under dark conditions [15]. Furthermore, TaWRKY7 from wheat can significantly promote senescence in transgenic Arabidopsis under dark conditions [16].

According to previous reports, WRKY TFs are thought to be involved in the regulation of plant tissue growth and development. For example, VvWRKY2 is specifically expressed in the lignified cells of young grapevine stems, and overexpression of VvWRKY2 in N. tabacum affects the lignin biosynthesis pathway, thus influencing xylem development [17]. Li et al. reported that Atwrky13 mutants exhibit weaker stems due to altered development of parenchyma cells [18]. Another WRKY TF, WRKY71/EXB1, positively regulates plant branching by controlling axillary meristem initiation and bud activities [19]. In addition, the pollen-specific WRKY TF AtWRKY34 is phosphorylated by two mitogen-activated protein kinases, MPK3 and MPK6, in the regulation of male gametogenesis [20]. Furthermore, emerging evidence has demonstrated that WRKY TFs are widely involved in stress responses. For example, GhWRKY40 is involved in pathogen responses [21], and GhWRKY68 is involved in salt and drought stress responses [22]. These reports further emphasize the importance of studying WRKY TFs.

Cotton (Gossypium hirsutum) is an important economic crop that is widely cultivated around the world. As a significant source of fiber, oil and biofuel products, cotton has become an important industrial raw material. In field production, the growth and yield of cotton are severely restricted by both external environmental factors and internal factors. A growing number of studies have shown that WRKY TFs play important roles in the responses to these factors. Therefore, it is particularly important to study the functional roles of WRKY genes in cotton. In the present study, a group IId WRKY gene, GhWRKY42, was isolated and characterized. We performed a preliminary analysis of the gene structure, evolutionary relationships and expression patterns of GhWRKY42. Overexpression of GhWRKY42 in Arabidopsis accelerated leaf senescence. In addition, silencing GhWRKY42 in VIGS plants significantly reduced plant height.

Results

Sequence and evolutionary analysis of GhWRKY42

We previously identified several WRKY genes in cotton that were up regulated by abiotic stresses, during leaf senescence and in vegetative organs using cDNA microarray and RNA-Seq data [23]. Among them, we selected GhWRKY42 for further study. The sequence analysis results showed that GhWRKY42 contained a 1038-bp ORF, encoding 345 amino acids. The predicted protein isoelectric point was 9.38, and the molecular weight was 37.88 kDa. The results of comparative analysis of the GhWRKY42 coding and genomic sequences indicated that GhWRKY42 harbored three exons and two introns (Fig. 1a). The multiple sequence alignment results revealed that the GhWRKY42 protein contained one WRKY domain, consisting of a conserved WRKYGQK core sequence and a C2H2 (C-X5-C-X23-H-X1-H) zinc-finger-like motif. Therefore, GhWRKY42 belongs to the group II WRKY subfamily according to Eulgem et al. [3]. Furthermore, a putative nuclear localization signal (NLS) sequence (KKRK) and a conserved HARF structural motif were found within the GhWRKY42 amino acid sequence, which are shared among group IId WRKY proteins (Fig. 1b). A phylogenetic tree was built to evaluate the evolutionary relationship between GhWRKY42 and other group II WRKY members from different species (Fig. 2). As shown in Fig. 2, GhWRKY42 was closely associated with group IId members, which was consistent with the results of the amino acid alignment analysis.

GhWRKY42 localizes to the nucleus

Consistent with the identified NLS sequence, the subcellular location prediction software Plant-mPloc (http://www.csbio.sjtu.edu.cn/bioinf/plant-multi/) predicted that the GhWRKY42 protein localizes to the nucleus. To confirm our prediction, the 35S-GhWRKY42::GFP vector was constructed and transferred into onion epidermal cells. The 35S::GFP construct served as a control. The onion epidermal cells harboring the 35S-GhWRKY42::GFP construct emitted green fluorescence predominantly in nuclei (Fig. 3a), whereas 35S::GFP fluorescence occurred widely throughout the cell [24].

Transcriptional activation assay of GhWRKY42

The transcriptional activation of GhWRKY42 was examined with a GAL4 yeast system. The plasmids pGADT7-largeT+pGBK7-GhWRKY42 (experimental group), pGADT7-largeT+pGBK7-p53 (positive control) and pGADT7-largeT+pGBK7-laminC (negative control) were transformed into Y2HGold yeast cells. All transformants grew well on SD−Trp−Leu medium. The transformants of the positive control grew well on SD−Trp−Leu−His−Ade medium, but similar to the
negative control, the experimental group did not grow on this medium (Fig. 3b).

Promoter analysis of GhWRKY42
A 1943-bp GhWRKY42 promoter fragment was obtained, and putative cis-elements were analyzed using the PlantCARE database. A group of putative cis-elements were identified in the promoter region, which were mainly involved in defense, stress, light and metabolic responses (Additional file 1: Table S1).

The results of GUS staining for the promoter deletion constructs showed that pBI121 (positive control) as well as ProGhWRKY42::GUS (−1943 bp to −1 bp), ProGhWRKY42−1::GUS (−1407 bp to −1 bp) and ProGhWRKY42−2::GUS (−778 bp to −1 bp) produced blue dots, whereas ProGhWRKY42−3::GUS (−391 bp to −1 bp) did not (Fig. 4a). The GUS staining results for different tissues showed that GUS was mainly active in the roots, stems and leaves of ProGhWRKY42::GUS transgenic Arabidopsis plants and was also detectable in the stamens but not in the pistils, petals or pods (Fig. 4b).

Expression analysis of GhWRKY42 under stress treatments
To evaluate the expression patterns of GhWRKY42 following various stresses, ten-day-old cotton seedlings were exposed to MeJA, ABA, drought and salt treatments. As shown in Fig. 5, GhWRKY42 was found to be differentially up-regulated under MeJA, ABA, drought and salt treatments. GhWRKY42 expression was rapidly induced at 2 h after MeJA treatment, reaching its maximum accumulation at 4 h (4.6-fold induction) and then gradually decreasing (Fig. 5a). Similarly, GhWRKY42 expression was induced at 2 h after ABA treatment but exhibited maximum transcript levels at 6 h with 2.7-fold induction (Fig. 5b). Under drought treatment, the GhWRKY42 transcript was differently elevated at different time points and peaked at 12 h (6.5-fold induction) (Fig. 5c). However, under salt treatment, the expression of GhWRKY42 was dramatically increased at 2 h, and a high expression level was maintained in the subsequent 4–12 h (Fig. 5d).

Expression analysis of GhWRKY42 in different tissues and during leaf senescence
qRT-PCR was performed to detect the transcript levels of GhWRKY42 in the roots, stems, leaves, petals, pistils, stamens, fiber and ovules. GhWRKY42 was found to be differentially expressed in different tissues. GhWRKY42 was strongly expressed in vegetative organs, including the stems, roots and leaves but was weakly expressed in the petals, pistils, stamens, fiber and ovules (Fig. 6a).

To evaluate the expression pattern of GhWRKY42 during leaf senescence, qRT-PCR was performed using cotton leaves at different senescence stages. The transcriptome data analysis [23, 25] showed that the expression level of GhWRKY42 gradually increased with the senescence of leaves (Fig. 6b). We further examined the expression level of GhWRKY42 in true leaves of CCR74 plants at different senescence stages [24]; the results revealed that the transcript levels of GhWRKY42 gradually increased as the leaves aged (Fig. 6c). In addition, the expression level of GhWRKY42 was detected in cotyledon samples from the

![Fig. 1 Gene structure and sequence analysis of GhWRKY42. a Gene structure of GhWRKY42. b Sequence alignment of the deduced GhWRKY42 protein with its homologous proteins AtWRKY15 (At2g23320), OsWRKY17 (XP_015633572), ZmWRKY21 (XP_008673611), GmWRKY15 (XP_006573647) and NtWRKY15 (XP_016498573). The approximately 60-amino acid WRKY domain is indicated by the two-headed arrow. The highly conserved WRKY domain core sequence WRKYGQK is boxed. The C and H residues in the zinc-finger motif are indicated by arrows. The KKRK NLS and conserved HARF structural motif are indicated by a horizontal line. The abbreviations before the gene names of different species are as follows: At, Arabidopsis thaliana; Gh, Gossypium hirsutum; Zm, Zea mays; Gm, Glycine max; Os, Oryza sativa; Nt, Nicotiana tabacum.](image)
early-aging cotton variety CCRI10 and the non-early-aging variety Liao4086. The qRT-PCR results showed that the transcript levels of *GhWRKY42* increased gradually during cotyledon senescence and were significantly higher in CCRI10 than in Liao4086 (Fig. 6d).

**GhWRKY42** promotes leaf senescence in transgenic *Arabidopsis*

The transcript of *GhWRKY42* was highly accumulated in the senescent leaves of cotton. To further clarify the functional role of *GhWRKY42* in response to leaf senescence, *GhWRKY42* was transformed into *Arabidopsis* plants. The transgenic lines were confirmed by qRT-PCR (Fig. 7a). As shown in Table 1, the *GhWRKY42* transgenic plants flowered earlier and had fewer rosette leaves than the WT plants. In addition, the senescence phenotypes of the transgenic and WT plants were observed at different developmental stages, and the ratio of senescent leaves was counted. Compared with the WT, the transgenic lines exhibited severe aging phenotypes at four, five and seven weeks (Fig. 7b), which were reflected by a significantly higher ratio of senescent cotyledons at four weeks (Fig. 7c), a higher ratio of senescent true leaves (rosette leaves) at five weeks (Fig. 7d) and a lower chlorophyll content at seven weeks (Fig. 7e).

**Increased expression levels of senescence-associated marker genes and ABA-responsive genes in *GhWRKY42*-overexpressing plants**

To elucidate the possible mechanisms of *GhWRKY42*-mediated precocious senescence, we examined the effects of *GhWRKY42* on the transcript levels of senescence-associated marker genes during natural leaf senescence. The genes included *AtNAP* (NAC domain TF) (*At1g69490*), *AtSAG12* (*At5g45890*), *AtSAG13* (*At2g29350*), *AtWRKY6* (WRKY DNA-binding protein 6) (*At1g62300*) and *AtORE1/AtNAC6* (NAC domain TF) (*At5g39610*), which are all factors that are up-regulated during aging in *Arabidopsis* [26–30]. As shown in Fig. 8a-e, the expression of all senescence-associated marker genes in the
Fig. 3  Subcellular localization and transcriptional activity assays of GhWRKY42.  

a Transient expression of the 35S-GhWRKY42::GFP construct in onion epidermal cells.  
b Transcriptional activity of GhWRKY42 in Y2HGold yeast cells. The ORF of GhWRKY42 was cloned into the pGBKT7 vector. The constructs were transformed into Y2HGold yeast cells and identified on SD/-Trp/-Leu and SD/-Trp/-Leu/-His/-Ade medium.  
PC, positive control (pGADT7-largeT+pGBKT7-p53); NC, negative control (pGADT7-largeT+pGBKT7-laminC); AD+BD-42, experimental group (pGADT7-largeT+pGBKT7-GhWRKY42).

Fig. 4  GUS activity analysis of the GhWRKY42 promoter.  

a Detection of the GUS activity of the GhWRKY42 promoter deletion vector transiently expressed in Nicotiana tabacum. From the left to right, the pBI121 vector (positive control) and the ProGhWRKY42::GUS (~1943 bp to ~1 bp), ProGhWRKY42-1::GUS (~1407 bp to ~1 bp), ProGhWRKY42-2::GUS (~778 bp to ~1 bp) and ProGhWRKY42-3::GUS (~391 bp to ~1 bp) constructs are shown.  
b GUS activity of ProGhWRKY42::GUS transgenic Arabidopsis in different tissues. (a-e) roots, stems, leaves, floral organs and pods.
transgenic plants was significantly up-regulated compared with that in the WT plants. In addition, we identified the expression levels of two ABA-responsive genes, *AtABF2* (ABA-responsive element binding factor 2) (AT1G45249) [31] and *AtHAB1* (hypersensitive to ABA1) (AT1G71770) [32], in *Arabidopsis*. The expression levels of both genes were significantly elevated in the transgenic plants compared with the WT plants (Fig. 8f, g).

**Decreased plant height in GhWRKY42-silenced cotton plants obtained via VIGS**

To further identify the functional role of *GhWRKY42*, VIGS of *GhWRKY42* was performed using the cotton variety CCR110. Two weeks later, the cotton plants harboring pCLCrVA-PDS showed an albino phenotype, suggesting that the VIGS assay was successful (Fig. 9a). qRT-PCR was performed to evaluate the effect of gene silencing. Expression level of *GhWRKY42* was significantly lower in the silenced plants (pCLCrVA-*GhWRKY42*) than in the control plants (pCLCrVA) (Fig. 9b). The expression of the senescence-associated marker gene *GhNAP* was also markedly reduced in the silenced plants (Fig. 9c). As shown in Fig. 9a, the silenced plants exhibited a relatively lower plant height phenotype than the control plants, and the lower plant height phenotype was statistically analyzed (Fig. 9d).

**Discussion**

The WRKY TF family is one of the largest superfamilies of regulatory proteins in plants [7]. In the past several years, growing evidence has shown that members of the WRKY gene family mainly participate in stress responses, plant growth and development and leaf senescence [33]. However, studies on WRKY TFs have mainly focused on model plant species, while only a few of these genes have been evaluated in cotton. In this study, we isolated a group IId *GhWRKY42* gene from upland cotton and characterized its functional roles. The results of multiple sequence alignment and phylogenetic tree analyses revealed that the *GhWRKY42* gene is a member of the group IId WRKY family. Subcellular localization analysis revealed that the *GhWRKY42* protein is located in the nucleus. These findings are consistent with the predicted nuclear-targeting signal sequence and with the results of studies on another group IId TF, *GhWRKY11*,...
in cotton [34]. The results of transcriptional activation analysis in yeast have shown that the *GhWRKY42* protein has no transcriptional activation activity; these findings are similar to those reported for *PtrWRKY40* from *Populus trichocarpa* [35]. These results suggest that *GhWRKY42* may be a nuclear protein that functions in the cell nucleus but may be not a transcriptional activator.

The expression patterns of genes are often used as an indicator of their functional roles [36]. For example, *GhWRKY17* has been shown to be induced by salt and drought treatments, and overexpression of *GhWRKY17* in *N. tabacum* results in a more sensitive phenotype to drought and salt stresses [37]. Previous studies have shown that a large number of genes can be induced by various abiotic stresses [38]. In our study, *GhWRKY42* was demonstrated to be differentially induced under MeJA, ABA, drought and salt treatments in cotton. In addition, many stress response *cis*-elements were found in the promoter region of *GhWRKY42*. These findings suggested that *GhWRKY42* might be involved in the regulation of abiotic stress networks.

The 5′ promoter deletion assay is often used to investigate promoter expression characteristics and the functional roles of regulatory elements in promoter regions. The structure and function of promoter deletion fragments can be suggested by evaluating promoter deletion construct-driven reporter genes in transgenic plants [39]. In our study, a promoter deletion assay showed that *ProGhWRKY42–2:GUS* (−778 bp to −1 bp) contained the shortest sequence exhibiting promoter activity. Therefore, it is speculated that critical *cis*-elements may exist within the −778 bp to −391 bp upstream region of the *GhWRKY42* promoter. The *cis*-elements in this region were predicted, it was found to contain not only TATA-box and light response elements but also stress response elements such as ABRE (ABA response element), CGTCA motif (MeJA response element), HSE

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**Fig. 6** Expression profiles of *GhWRKY42* in different tissues and during leaf senescence in cotton. **a** Expression levels of *GhWRKY42* in different tissues. **b** Expression profile of *GhWRKY42* during leaf senescence in 15-, 25-, 35-, 45-, 55-, and 65-day-old leaves. RPKM, reads per kilobase per million mapped reads. **c** Expression level of *GhWRKY42* in true leaves at different developmental stages. Stage 1, a fully expanded, young, non-senescent leaf; Stage 2, a fully expanded, mature, non-senescent leaf; Stage 3, a leaf in early senescence with < 25% leaf area yellowing; Stage 4, a leaf in mid-senescence with approximately 50% leaf area yellowing; Stage 5, a leaf in late senescence with > 75% leaf area yellowing. **d** Relative expression level of *GhWRKY42* in the cotyledons of the CCR10 and Liao 4086 varieties. Cotyledon samples were collected weekly at eight different development stages, ranging from the flattened cotyledon stage to the completely aged stage. *GhActin* served as the reference gene. The bars represent the means ± SEs from three independent experiments.
(heat response element) and the TGACG motif (MeJA response element). These elements may play an important role in ensuring that the promoter drives the expression of downstream genes.

WRKYs can directly bind to the W-box [TTGAC(C/T)] in the promoter of target genes to modulate stress responses, plant development and leaf senescence [40, 41]. Liu et al. reported that W-box and G-box cis-elements are important positive regulators during leaf senescence in rice. Both elements are significantly plentiful in the promoter regions of up-regulated TFs (including WRKYs) that regulate leaf senescence [41]. W-box and G-box cis-elements were identified in the promoter region of GhWRKY42, suggesting that GhWRKY42 may be involved in leaf senescence and be regulated by other GhWRKYs or by the gene itself during this process. These findings laid the foundation for further analysis of the upstream regulatory mechanism of GhWRKY42.

Senescence is a natural phenomenon and prevails among all living organisms, including plants. During leaf senescence, genetic and environmental factors affect mature leaves, leading to the initiation of leaf senescence; this senescence is accompanied by chlorophyll, membrane, protein and nucleic acid degradation as well as nutrient relocation from senescing leaves to growing organs or storage tissues [42–45]. Crop productivity is mainly determined by the yield per area, but leaf senescence severely affects crop yield [46]. Thus, studying the mechanism of leaf senescence is particularly important. In the present study, the expression level of GhWRKY42 was found to be up-regulated during natural senescence and exhibited significantly higher expression in the early-aging cotton variety CCRI10 than in the non-early-aging variety Liao4086. It has been reported that the GhNAC12 gene, which is more highly expressed in CCRI10 than in Liao4086 during leaf senescence,
**Fig. 8** Expression levels of SAGs and ABA-responsive genes in WT and transgenic plants during leaf senescence. Expression levels of AtNAP (a), AtSAG12 (b), AtSAG13 (c), AtWRKY6 (d), AtORE1/AtNAC6 (e), AtABF2 (f) and AtHAB1 (g) in WT and transgenic plants. The AtNAP, AtSAG12, AtSAG13, AtWRKY6 and AtORE1/AtNAC6 genes are SAGs. The AtABF2 and AtHAB1 genes are ABA-responsive genes. Total RNA was isolated from five-week-old Arabidopsis rosette leaves. GhActin served as the reference gene. The bars represent the means ± SEs from three independent experiments. Independent t-tests revealed highly significant (**P < 0.01) differences between the WT and the transgenic lines.

**Fig. 9** Silencing GhWRKY42 via VIGS decreases plant height in cotton. a Plant phenotypes of empty control and GhWRKY42-silenced plants. b Expression level of GhWRKY42 in empty control and GhWRKY42-silenced plants. c Expression level of GhNAP in empty control and GhWRKY42-silenced plants. d Plant height of empty control and GhWRKY42-silenced plants. GhActin served as the reference gene. The bars represent the means ± SEs from three independent experiments. Independent t-tests revealed highly significant (**P < 0.01) differences between the control and silenced plants.
causes an early-aging phenotype in Arabidopsis [47]. Therefore, \textit{GhWRKY42} may be involved in the aging process and may play a regulatory role during leaf senescence. Consistent with our prediction, overexpression of \textit{GhWRKY42} did lead to an advance of leaf senescence in transgenic \textit{Arabidopsis}. In a previous study, overexpression of \textit{AtWRKY45} in \textit{Arabidopsis} was observed to up-regulate expression of representative SAGs during age-triggered leaf senescence [13]. Pheno- typic observations of overexpressing \textit{Arabidopsis} lines and RNAi cotton lines show that \textit{GhNAP} positively reg- ulates leaf senescence through ABA-mediated pathways [48]. Phytohormones, such as ABA, ethylene, MeJA, and salicylic acid, have been demonstrated to promote leaf senescence [45]. The ABA content increases in aged leaves, and endogenously applied ABA promotes expres- sion of some SAGs [44]. ABA-responsive genes, which are involved in the ABA signaling pathway, are induced in senescing \textit{Arabidopsis} [49]. In our study, SAGs and ABA-responsive genes were significantly accumulated in the \textit{GhWRKY42} transgenic lines, suggesting that \textit{GhWRKY42} may be associated with leaf senescence via ABA-mediated pathways.

Previous studies have shown that some genes are closely associated with plant height. Wei et al. identified the QTL DTH8 in rice, which includes the HAP3 gene and regulates yield, plant height and flowering time [50]. WRKY TFs such as \textit{LP1} in foxtail millet and \textit{OsWRKY78} in rice have all been shown to play an important role in stem elongation and plant height [51, 52]. In our study, we detected high expression levels and strong GUS ac- tivity of \textit{GhWRKY42} in the stem and reduced height in VIGS plants. Therefore, we hypothesized that \textit{GhWRKY42} might be related to stem development. Plant height is an important plant architecture trait, and decreased height is beneficial for mechanical harvesting and lodging resistance [53]. Our findings provide a basis for breeding new cotton varieties with an ideal plant type. However, further studies are needed to elucidate the pathways involved in the \textit{GhWRKY42}-mediated mechanism.

Conclusions
\textit{GhWRKY42}, a group IId WRKY member, is closely asso- ciated with leaf senescence and plant development. \textit{GhWRKY42} is located in the nucleus and exhibits no transcriptional activity. \textit{GhWRKY42} is induced by leaf senescence and various stresses. Ectopic expression of \textit{GhWRKY42} in \textit{Arabidopsis} promotes leaf senescence, and VIGS cotton plants exhibit a decreased plant height phenotype. Our work could lead to a better understand- ing of the functional roles of WRKY genes in cotton. However, how the \textit{GhWRKY42} gene regulates leaf senescence and plant height development requires fur- ther study and clarification.

Methods

Plant materials, growth conditions and stress treatments
Two early-aging cotton varieties, CCR110 and CCR174, and a non-early-aging variety, Lia4086, were used in our experiments. The cotton varieties were cultivated in the field of the Cotton Research Institute of the Chinese Academy of Agricultural Sciences (Anyang, Henan, China). Different tissues were collected from CCR110 plants. Roots and stems were collected from two-week-old seedlings. Leaves were collected from newly flattened leaves. Petals, pistils, and stamens were sampled at anthesis, and fiber and ovules were harvested at 10 days post anthesis.

To evaluate the expression pattern of \textit{GhWRKY42} during leaf senescence, cotyledons were collected from two cotton varieties, CCR110 and Lia4086, which exhibit different aging characteristics. We collected cotyledon samples weekly at eight different developmental stages, ranging from the flattened cotyledon stage to the completely aged stage. The expression patterns of \textit{GhWRKY42} were further evaluated in true leaves of the early-aging cotton variety CCR174 at five aging stages, as described previously [24]. Each sample included material from eight different individual plants, and we performed three repetitions for each sample.

To evaluate the stress response of \textit{GhWRKY42} in cot- ton, 10-day-old CCR110 cotton seedlings were planted in pots for subsequent stress treatments. The CCR110 cot- ton seedlings were planted in a growth chamber at 25 ° C, with a 16 h light/8 h dark cycle. For the abiotic stress treatment, the seedlings were irrigated with 15% poly- ethylene glycol 6000 (PEG6000) and 200 mM sodium chloride (NaCl); for the signaling molecule treatment, the seedlings were sprayed with 100 μM methyl jasmo- nate (MeJA) and 200 μM abscisic acid (ABA). Each coty- ledon sample included material collected from eight uniform plants, and each treatment was repeated three times. The samples were harvested at 0 h, 2 h, 4 h, 6 h, 8 h and 12 h. All samples were quickly frozen in liquid nitrogen for subsequent RNA extraction.

Gene cloning and sequence analysis
To amplify the full-length cDNA and genomic se- quences of \textit{GhWRKY42}, primers were designed based on the coding sequence of \textit{GhWRKY42} (accession KF669797) submitted to NCBI by Dou et al. [23]. The primers used for this purpose are listed in Additional file 2: Table S2. The full-length cDNA and gen- omic fragments of \textit{GhWRKY42} were amplified from cDNA and DNA, respectively, obtained from CCR110 leaves at the five-leaf stage. The fragments were subse- quently inserted into the pMD18-T vector (TaKaRa,
brane was washed with 70% and 100% alcohol, plasmids with 75% alcohol. After the particulate carrier membrane and some accessories were cleaned and sterilized and was placed on a clean bench, and the bombardment chamber and some accessories were cleaned and sterilized and was placed on a clean bench, and the bombardment chamber and some accessories were cleaned and sterilized and was placed on a clean bench.

- A solid Murashige and Skoog (MS) medium and cultivated at 28 °C for 3 days in darkness.

- Approximately 1.5 × 1.5 cm pieces with a scalpel on a clean bench. The epidermal pieces were then transferred to fresh onion. The inner epidermis of a fresh onion was cut into approximately 1.5 × 1.5 cm pieces with a scalpel on a clean bench.

- The epidermal pieces were then transferred to fresh onion. The inner epidermis of a fresh onion was cut into approximately 1.5 × 1.5 cm pieces with a scalpel on a clean bench.

- The epidermal pieces were then transferred to fresh MS agar medium at 25 °C for 12 h in darkness. The resulting green fluorescence was detected using a confocal laser scanning microscope (Zeiss LSM 700) at a wavelength of 488 nm.

RNA isolation, cDNA synthesis and qRT-PCR

Total RNA was isolated using RNAprep PurePlant Kit (Polysaccharides & Polyphenolics-rich) (Tiangen, China). One microgram of total RNA was prepared for cDNA synthesis in a 20 μl reaction system using a PrimeScript™ RT reagent kit with gDNA Eraser. The cDNA was diluted 5 times for qRT-PCR. Transcript levels were determined using a 7500 Real-Time PCR system (Applied Biosystems) and SYBR® Premix Ex Taq™ II (TaKaRa). The 20 μl reaction volume contained the following components: 10 μl of SYBR Premix Ex Taq™ II (Tli RNaseH Plus) (TaKaRa). The 20 μl reaction volume contained the following components: 10 μl of SYBR Premix Ex Taq™ II (Tli RNaseH Plus) (2x), 0.8 μl of the PCR forward primer (10 μM), 0.8 μl of the PCR reverse primer (10 μM), 0.4 μl of ROX Reference Dye II (50x), 2 μl of cDNA and 6 μl of ddH₂O. The optimal PCR amplification procedure used was as follows: a pre-denaturation step at 95 °C for 30 s; 40 cycles of 95 °C for 5 s and 60 °C for 34 s; and a melting curve step at 95 °C for 15 s, 60 °C for 1 min and 95 °C for 15 s. *GhActin* and *AtActin2* were used as reference genes. The 2−ΔΔCT method was applied to calculate relative expression levels [54]. Three independent experiments were performed, and all reactions were performed with three technical replicates.

Subcellular localization

The open reading frame (ORF) of *GhWRKY42* without the termination codon was cloned into the pBI121-GFP vector driven by the cauliflower mosaic virus (CaMV) 35S promoter. The 35S::GhWRKY42::GFP plasmid was extracted to obtain a plasmid concentration of at least 1 μg/μl. The inner epidermis of a fresh onion was cut into approximately 1.5 × 1.5 cm pieces with a scalpel on a clean bench. The epidermal pieces were then transferred to solid Murashige and Skoog (MS) medium and cultivated at 28 °C for 3–6 h in darkness. The gene gun device was sterilized and was placed on a clean bench, and the bombarding chamber and some accessories were cleaned with 75% alcohol. After the particulate carrier membrane was washed with 70 and 100% alcohol, plasmids encased in gold powder were added to the middle of the particulate carrier membrane. After the membrane dried slightly, the onion epidermis was bombarded using the gene gun with the following parameters: particle bombardment running distance, 9 cm; rupture disk pressure, 1300 psi; and vacuum degree, 28 mmHg. The epidermis after bombardment was transferred to fresh MS agar medium at 25 °C for 12 h in darkness. The resulting green fluorescence was detected using a confocal laser scanning microscope (Zeiss LSM 700) at a wavelength of 488 nm.

Transcriptional activation assays

The ORF of *GhWRKY42* was cloned into the pGBKTT7 vector to construct pGBKTT7-*GhWRKY42*. The pGADT7-largeT+pGBKT7-35S::GhWRKY42 constitutive group, pGADT7-largeT+pGBKT7-p53 (positive control) and pGADT7-largeT+pGBK7-laminC (negative control) plasmids were transformed into yeast competent cells. The transformed yeast products were spread on corresponding dropout selective medium plates that did not contain tryptophan or leucine (SD/−Trp/−Leu) and incubated for 3–5 days at 30 °C. Positive clones were identified and streaked on SD/−Trp/−Leu medium plates and plates containing medium without tryptophan, leucine, histidine or adenine (SD/−Trp/−Leu/−His/−Ade). The plates were inverted and incubated at 30 °C for 3–5 days to identify transcriptional activity.

Genetic transformation of *Arabidopsis thaliana*

The ORF of *GhWRKY42* was inserted into the binary expression vector pBI121 driven by the 35S promoter to generate the 35S-35S::GhWRKY42 construct. The 35S::GhWRKY42 promoter fragment was also inserted into the pBI121 vector by replacing the 35S promoter to generate the Pro35S::GhWRKY42 construct. The 35S:GhWRKY42 and Pro35S::GhWRKY42 constructs were individually introduced into *Agrobacterium tumefaciens* strain LBA4404 and transformed into *Arabidopsis thaliana* using the floral-dip method [55]. For the screening of positive plants, seeds of the T0 generation (harvested from the wild-type (WT)) were sterilized and selected on 1/2 MS solid medium plates (0.22% MS modified basal salt mixture, 3% sucrose and 0.8% agar powder) containing kanamycin (50 mg/L). The plates containing the seeds were chilled at 4 °C for 3 days in darkness, after which they were transferred to an incubator at 22 °C under a 16 h light/8 h dark cycle with a light intensity of 100 μmol m−2 s−1. Two weeks later, the green seedlings on the plates were selected and transplanted into the nutrient soil in a growth chamber. The positive plants were further verified using PCR, and selfed seeds harvested from the positive plants were employed as the T1 generation. Using the same method, the seeds were screened until the T3 homozygous.
generation. The phenotypic characteristics of the transgenic and WT plants were observed at different developmental stages.

**Transient transformation of N. tabacum and GUS histochemical staining assays**

Based on the position of stress response cis-elements in the GhWRKY42 promoter, four promoter deletion fragments were delimited. The four fragments were amplified from the pMD18-T vector containing the GhWRKY42 promoter and inserted into the pBI121 vector by replacing the 35S promoter. As a result, four promoter deletion plasmids, ProGhWRKY42::GUS (−1943 bp to −1 bp), ProGhWRKY42−1::GUS (−1407 bp to −1 bp), ProGhWRKY42−2::GUS (−778 bp to −1 bp) and ProGhWRKY42−3::GUS (−391 bp to −1 bp), were constructed and transformed into LBA4404. Transient expression in N. tabacum was performed in accordance with previously described methods [56]. Transgenic Arabidopsis plants harboring the ProGhWRKY42::GUS construct were used to analyze organizational expression characteristics. GUS staining was performed as follows: the prepared materials were soaked in the GUS dye solution, after which the materials were placed in darkness at 25–37 °C overnight; the materials were then decolorized approximately 2–3 times using 70% alcohol until the negative control materials turned white, and the blue dots in the white background observed under microscopy were identified as GUS expression sites.

**VIGS assay**

For the VIGS assay, approximately 300-bp fragments amplified from the pMD18-T vector containing the GhWRKY42 gene were integrated into the pCLCrVA vector to construct pCLCrVA-GhWRKY42, which was then transformed into LBA4404. The LBA4404 strains carrying pCLCrVA-GhWRKY42, pCLCrVA (negative control) or pCLCrVA-PDS (positive control) were mixed with the strain harboring pCLCrVB (helper vector) (1:1 ratio, OD600 = 1.5) and co-injected into two fully expanded cotyledons of CCR110 plants. In the VIGS assay, at least 20 seedlings were used per group. For qRT-PCR detection, samples from at least 6 uniform injected plants were used. The cotton plants were then cultivated at 22 °C with a 16 h light/8 h dark cycle in a greenhouse. The experiment was repeated three times. The detailed VIGS procedure was performed as previously described [57, 58].

**Determination of chlorophyll content**

Determination of the chlorophyll content was performed as described by Shah et al. [59].

**Additional files**

- **Additional file 1:** Table S1. Predicted cis-acting elements in the promoter region of GhWRKY42. (DOCX 36 kb)
- **Additional file 2:** Table S2. Primers used in this study. (DOCX 32 kb)

**Abbreviations**

- ABA: Abscisic acid
- DAS: Days after sowing
- GUS: β-glucuronidase
- MeJA: Methyl jasmonate
- NaCl: Sodium chloride
- NLS: Nuclear localization signal
- ORF: Open reading frame
- PEG6000: Polyethylene glycol 6000
- qRT-PCR: Quantitative real-time PCR
- SAGs: Senescence-associated genes
- SEs: Standard errors
- TFs: Transcription factors
- VIGS: Virus-induced gene silencing
- WT: Wild-type

**Competing interests**

The authors declare that they have no competing interests.
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