

# *GhD14* Regulates Plant Architecture and Fiber Development in Cotton

**Liping Zhu**

Shaanxi Normal University

**Lingling Dou**

Xianyang Normal University

**Zailong Tian**

Shaanxi Normal University

**Huizhi Zhang**

Shaanxi Normal University

**Li Zhang**

Shaanxi Normal University

**Cuixia Liu**

Shangluo University

**Jianing Yu**

Shaanxi Normal University

**Guanghui Xiao** (✉ [guanghuix@snnu.edu.cn](mailto:guanghuix@snnu.edu.cn))

Shaanxi Normal University <https://orcid.org/0000-0003-2717-8012>

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## Research

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# Abstract

**Background:** Strigolactone (SL) signaling is essential in regulating plant development. DWARF14 (D14), the SL receptor, interacts with the F-box in MORE AXILLARY GROWTH (MAX2) to modulate SL signaling. However, the biological function of D14 protein is still unknown in cotton.

**Results:** Here, we identified GhD14s in *Gossypium hirsutum* and resolved its function in cotton plant architecture and fiber development. Subcellular location results revealed that the GhD14D protein was localized to both the cytoplasm and nucleus. GUS staining assay showed that GhD14D was mainly expressed in leaf primordium, inflorescence, axillary bud and stem and expression analysis revealed that *GhD14A/D* was highly expressed in stem, flower and fiber cells at 20 days post-anthesis (DPA). Silencing *GhD14A/D* gene expression in upland cotton significantly increased branch angle. Meanwhile, the fiber length and the transcripts of secondary cell wall biosynthesis related genes were also reduced after *GhD14A/D* gene silencing. In addition, overexpression of *GhD14D* in *Atd14* mutant successfully rescued the phenotype of the *d14* mutant with much shoot-branching and short plant height.

**Conclusions:** Our findings suggest that the *GhD14* gene contributes to shoot branch development and fiber cell development in cotton. This study deepens our understanding of the biological role of SL signaling in cotton and providing guidance for modifying cotton plant architecture and improving fiber development using genetic engineering to help us breed better cotton varieties in the future.

## Background

Cotton is an important cash crop in the world, the upland cotton (*Gossypium hirsutum* L.) is the largest source of textile fiber (Chen et al. 2007). Cotton plant architecture is a significant agronomic trait due to the correlation with planting density, the suitability of mechanical harvest and yield (Su et al. 2018). Compared with increased yield potential per plant, increasing the planting density significantly enhance yield per unit area in cotton production (Ji et al. 2021). Plant architecture is of great important in cotton production due to the modern cultivation in China requiring a compact structure for mechanized harvesting processes (Mao et al. 2015). The plant height and branch length are major limiting factors for planting density in cotton fields (Su et al. 1993). The investigation of cotton plant architecture is of great significant on increasing cotton yield.

Plant architecture refers to the organization of the plant body, includes shoot branching, branch length, branch angle, position of organs and so on in plants (Wang et al. 2008). Higher plants display various plant architectures for their different requirements. For crops, the crop yield and quality could be affected by plant architecture. After the Green Revolution, crop yields dramatically increased due to higher planting density, which was made possible by changes in plant architecture leading to shorter and more compact cultivars (Peng et al. 1999). The plant architecture is influenced by many environmental factors such as moisture and light. The drought stress reduces plant height and grain yield (Todaka et al. 2015). The dark and cold condition cause the plant to be dwarfed. Phytohormones play important roles in plant

development process (Blazquez et al. 2021; Xia et al. 2019). The plant architecture also be regulated by phytohormones (Domagalska et al. 2011). The short and compact characteristics are mainly controlled by plant architecture, which is regulated by plant hormones, such as auxins and strigolactones (SLs) (Domagalska et al. 2011; Ongaro et al. 2008).

SLs have been recognized as a new class of hormones that participate in regulating plant architectures in terms of shoot branching, lateral roots, shoot gravitropism and stem secondary thickening (Wang et al. 2020). Branching is a highly plastic determinant of plant shape to allow plants to respond to environmental stresses (Evers et al. 2011). Shoot branching is a ubiquitous phenomenon in higher plants and a basic characteristic of plant growth, that is essential in determining plant architecture. The SL biosynthesis gene (HIGH TILLERING AND DWARF 1/DWARF17) increases tiller number of rice and improves rice grain yield (Wang et al. 2020). In Arabidopsis, three key repressors of the strigolactone (SL) signaling pathway (SMXL6/SMXL7/SMXL8) directly interact with two transcriptional factors and suppress their transcriptional activation of a key repressor of branching (BRC1), thus promoting branching (Xie et al. 2020). The SL could trigger polyubiquitination and degradation of SMXL2, then regulating the gene expression and hypocotyl elongation (Wang et al. 2020). The SL also control the degradation of cytokinin via activating the transcription of *CYTOKININ OXIDASE/DEHYDROGENASE 9* in rice (Duan et al. 2019).

The SL receptor, Dwarf14 (D14), is a key receptor that interacts with an F-box protein to form a Skp1-Cullin-F-BOX (SCF) complex and regulates SL signaling pathway (Nakamura et al. 2013). In soybean, the GmMAX2 interacted with the GmD14 and then forming active D14/KAI-SCF<sup>MAX2</sup> complexes for function (Ahmad et al. 2020). The SCF complex then recruits SL repressor proteins to be ubiquitinated and then the SL be degraded, hence the SL repressor proteins could regulate the expression level of SL-dependent genes (Marzec et al. 2016). A Yoshimulactone Green (YLG) based *in vitro* assay of a high-throughput chemical screening identified a novel small molecule DL1 as a potent inhibitor of D14 that competes with endogenous SLs and increases shoot branching in Arabidopsis and rice (Yoshimura et al. 2018). In *Dendranthema grandiflorum*, *DgMAX2*, a key regulatory gene in SL signal transduction, can restore *atmax2-1* mutant branching to wild-type (WT) in Arabidopsis (Dong et al. 2013). In Populus, there are two *PtD14* genes, but only *PtD14a* was able to recover the *Atd14* mutant (Zheng et al. 2016). Recent study revealed that *GhMAX2* contributes to plant growth, architecture and fiber elongation in cotton, knockdown expression of *GhMAX2* significantly reduced plant height, slow growth, short internodes, and reduced fiber length (He et al. 2021), however the D14 in *G. hirsutum* has not been investigated.

In this study, a putative SL receptor, *GhD14*, was identified in *G. hirsutum* and it was localized to both the cytoplasm and nucleus. The expression analysis showed that *GhD14* was primarily expressed in buds, stems, flowers and 20 DPA fibers. *GhD14*-silenced plants had a phenotype of increased branch angle, reduced fiber length and the transcripts level of secondary cell wall biosynthesis related genes. Meanwhile, the *GhD14* rescued the phenotype of the *d14-1* mutant with much shoot-branching and short plant height. These result shows that *GhD14* is involved in regulating cotton architecture and fiber development, which lays a foundation for manually controlling cotton plant architecture in the future.

# Materials And Methods

## Plant materials and growth conditions

*G. hirsutum* cultivar Xuzhou 142 was planted in a climate-controlled greenhouse with the same condition as previously reported (Lin et al. 2019). The fibers at 0, 3, 5, 10, 15, and 20 days after anthesis (DPA), stems, leaves, flowers from four months plants and roots from one month cotton plants were collected and frozen in liquid nitrogen immediately, then stored at -80°C until use. Seeds of wild-type *A. thaliana* (Columbia, Col-0) and *d14* mutant (Waters et al. 2012) were placed at 4°C vernalization for 48-72 hours, then surface-sterilized and germinated on 1/2 strength Murashige and Skoog (1/2 MS) agar plates. Then, *A. thaliana* seedlings were transferred from the medium to soil pots grown in chambers at 24°C 14 h /10 h, light/dark cycle. The seeds of *Nicotiana Benthamiana* are uniformly scattered in the soil and culture conditions are the same as the *G. hirsutum*.

## Bioinformatics analysis of D14 proteins

The open reading frame (ORF) of GhD14 was identified by ORF Finder (<https://www.ncbi.nlm.nih.gov/orffinder/>). The physicochemical properties were calculated by ExPASy online software (Gasteiger et al. 2003). The GhD14 protein secondary structure was predicted by SOPMA software with secondary structure prediction method (Geourjon et al. 1995). Three dimensional (3D) structure of GhD14D protein was constructed by SWISS-MODEL (Waterhouse et al. 2018). The DNAMAN software was used to further annotate the protein structure of GhD14s. The protein sequences of AtD14, OsD14 and GhD14s were submitted to DNAMAN software to perform multiple protein sequences alignment.

The D14 genes from two monocots (*Zea mays* and *Oryza sativa*) and 13 dicots (*Theobroma cacao*, *Corchorus olitorius*, *Cicer arietinum*, *Solanum pennellii*, *Capsicum annum*, *Juglans regia*, *Arachis duranensis*, *Glycine max*, *Abrus precatorius*, *Phtheirospermum japonicum*, *Striga hermonthica*, *Orobancha cernua* and *Arabidopsis thaliana*) were selected for phylogenetic analysis. The full-length protein sequences of these homologous were obtained from the National Center for Biotechnology Information (NCBI). Phylogenetic tree was constructed by MEGA 7.0 with the Neighboring-Joining (NJ) method and 1000 bootstrap replications (Kumar et al. 2016).

## DNA, RNA extraction and expression analysis of *Ghd14*

The total DNA of cotton leaves was extracted using CTAB method (Huang et al. 2000). The total RNA of fibers at 0, 3, 5, 10, 15 and 20 days post-anthesis (DPA) and cotton roots, stems, leaves, and flowers were extracted by the PureLink™ RNA mini kit (Invitrogen, Lot no.1687455, USA) according to the manual instructions. First-strand complementary DNA (cDNA) was reverse-transcribed from 2 µg of the total RNA by the description of PrimeScript™ RT reagent Kit with gDNA Eraser (TaKaRa, Code No. RR047A, Japan).

The primers were designed by OLIGO 7.0 (Rychlik. 2007). For expression analysis, the ubiquitin gene *GhUBQ7* was used as a housekeeping gene (Xiao et al. 2016). All primers and sequences in this work are listed in Additional file 7 and Additional file 8. The qRT-PCR experiments were performed using SYBR® Premix Ex Taq™ II (Takara, Japan) kit on a Bio-Rad Real Time PCR detection system (Bio-Rad CFX96Touch, USA) with the following reaction parameters: 94°C for 2 min, followed by 40 cycles of 94°C for 30 s and 60°C for 30 s. The  $2^{-\Delta\Delta CT}$  method was used to calculate the relative expression levels of the target genes. All data were statistically analyzed using by One-way ANOVA method by SigmaStat software with default parameters (Ji et al. 2003).

## Cloning, plasmid construction and transformation

The protein sequence of AtD14 was downloaded from TAIR database, and used as a query sequence to against the *G. hirsutum* protein genome database (ZJU\_v2.1). The most similarity genes to AtD14 in *G. hirsutum* were recognized as GhD14s. The gene was amplified from *G. hirsutum* cDNA with PrimeSTAR® Max DNA Polymerase (TaKaRa, Japan) and sequenced by TSINGKE company (Beijing, China).

The 1715 bp upstream genomic DNA sequence of the initiation codon (ATG) from *GhD14D* was cloned as the promoter sequence from Xuzhou 142 DNA and full-length CDS of *GhD14* were amplified from Xuzhou 142 cDNA. For overexpression of *GhD14D* experiment, the full-length of *GhD14D* CDS sequence was assembled into pCAMBIA1305 by In-Fusion® HD Cloning Kit (Vazyme Biotech, China) according to the instructions. For  $\beta$ -glucuronidase (GUS) staining analysis, the *GhD14D* promoter sequences was introduced into pCAMBIA2300 to drive the GUS gene expression. All the final constructs were transformed into *A. thaliana* and *N. benthamiana* by *Agrobacterium tumefaciens* mediated transformation method as previously described [49] (Cheng et al. 2018). In addition, the full-length CDS of *GhD14D* was cloned into pTF486-GFP and pCHF3-GFP to generate the 35S::*GhD14*-GFP construct for subcellular localization assays. Two primers in pCAMBIA1305 were used for identification of whether *A. thaliana* contain pCAMBIA1305: *GhD14D*. All the primers used in vector construction are listed in Additional Table 1. Meanwhile, one month old *A. thaliana* seedlings was used for observation of the rosette leaves of seedlings and leaves phenotype and Two and a half month *A. thaliana* was used for the whole plants phenotype observation.

## GUS staining analysis

For GUS staining analysis, the GhD14Dpro::GUS construct was introduced into *A. tumefaciens* strain GV3101 and subsequently transformed into *A. thaliana* using the floral dip method (Lloyd et al. 1986). Transgenic plants were selected on solid half-strength MS media plates containing 50  $\mu$ g/mL chloramphenicol. The selected transgenic seedlings were further validated by genomic PCR. Various tissue at different developmental stages were collected, and stained with GUS staining solution according to the instructions. The microscope (Nikon, Japan) was used to observe the GUS staining result.

## Transient expression of GhD14D-GFP

For subcellular localization of GhD14D, the fused GhD14D-GFP vectors were transiently expressed in *A. thaliana* protoplast and *N. benthamiana* according to previous studies (Abel et al. 1994; Cheng et al. 2018). The two fusion constructs pTF486-GFP and pCHF3-GFP were used for transient expression GhD14D-GFP in *A. thaliana* protoplast and in *N. benthamiana* epidermis cells, respectively. In addition, the nuclei staining, the chemical reagent 4', 6'-diamidino-2-phenylindole (DAPI) staining method was used to confirm the nuclei localization of GhD14D (Liang et al. 2018). The positive transgenic plants were used for *in vivo* subcellular localization analysis using a confocal microscope (Nikon, Japan).

### **Virus-induced *GhD14A/D* gene silence in *G. hirsutum***

The binary CLCr vectors pCLCrVA and pCLCrVB were used for virus-induced gene silence (VIGS) analysis (Gu et al. 2014). Three fragments of *GhD14A/D* gene were amplified using cDNA with the primers listed in Additional Table 1. The PCR fragments were inserted into pCLCrVA and these vectors were transformed into *A. tumefaciens* GV3101 by electroporation. The *A. tumefaciens* culture containing pCLCrVA or the constructed vectors were mixed with an equal volume of an *A. tumefaciens* culture containing pCLCrVB to generate the mixed *A. tumefaciens* solutions for infiltration. The mixed solutions were infiltrated into fully expanded cotyledons of 10-day-old cotton seedlings with 15 plants through vacuum infiltration [52] (Gao et al. 2011). The inoculated seedlings were then transferred to a dark climate-controlled greenhouse at 25°C. The one month old seedling leaves and 25 DPA fibers from negative control and *GhD14A/D*-VIGS cotton plants were chosen to assess the silence effects of *GhD14A/D* gene by qRT-PCR experiment. The sixth branches from bottom to top were used for detecting the angle of the monopodial branch.

## **Results**

### **Identification of the GhD14 in *G. hirsutum***

The AtD14 protein sequence was used as the query sequence to against the *G. hirsutum* genome (Hu et al. 2019) to obtain the D14 protein in *G. hirsutum*. The sequences GH\_A02G1790.1 and GH\_D03G0270.1 in the cotton genome were the most similar to AtD14 and were chosen as the candidate orthologs of AtD14 and named GhD14A and GhD14D, respectively. The GhD14A and GhD14D have one intron, contain 816 base pairs (bp) (Fig. S1A), 271 amino acids (Fig. 1A) and have a molecular weight of 19.9 kDa and 30.0 kDa, respectively.

### **Conserved domain and promoter analysis of GhD14s**

The AtD14 protein structure in *A. thaliana* has been reported as an  $\alpha/\beta$  hydrolase (Li et al. 2020). We compared the protein sequences of GhD14A, GhD14D and AtD14, and identified eight  $\alpha$ -helixes, five  $\eta$ -helixes and seven  $\beta$ -strands in D14 protein (Fig. 1B, Fig. S1B, C). There are three catalytic residues, S95, D217 and H246 in GhD14A and GhD14D protein sequences (Fig. 1B). The sequence similarities of AtD14 with GhD14A and GhD14D were 52.7% and 53.5%, respectively. As *GhD14A* and *GhD14D* have 97.4% similarity in coding sequences, we cloned *GhD14D* from *G. hirsutum* cultivar Xuzhou 142 and used it as

the representative *GhD14* gene for *cis*-elements distribution, subcellular location, GUS and overexpression analysis.

The *cis*-element distribution of 1,715 bp *GhD14D* promoter sequence and promoter driven GUS expression were investigated to study the potential function of *GhD14D*. The *GhD14D* promoter has high A and T content, and typical *cis*-elements TATA-box and CAAT-box (Fig. S2). The auxin responsiveness *cis*-element AUXRR-core and jasmonic acid methyl ester (MeJA) responsiveness regulatory element CGTCA-motif were also present in the *GhD14D* promoter region (Fig. S2), indicating that auxin and MeJA might function in regulating *GhD14D* gene expression via combine the corresponding *cis*-elements present in the promoter of *GhD14D*.

### Phylogenetic analysis of GhD14s

To explore the evolutionary relationships among GhD14D and other fifteen D14s from typical higher plant species, we performed phylogenetic analysis of D14 proteins from *G. hirsutum* and other fifteen higher plant species using MEGA 7.0 software with Neighboring-Joining (NJ) method and 1000 bootstrap replications. The phylogenetic tree showed that the D14s in sixteen higher plants have the same evolutionary origin. The D14s was resolved into two evolutionary branches in phylogenetic tree and both of which contain D14 proteins from monocotyledons and dicotyledons. Meanwhile, GhD14s was more closely related to eudicots, and was most closely related to *T. cacao*, with moderate bootstrap support of 79% (Fig. S3).

### GhD14D is localized to the cytoplasm and nucleus

The pTF486-GhD14D:GFP vector was constructed to investigate the subcellular location of GhD14D protein and DAPI staining was used to test whether the GhD14D protein was localized in the nucleus. The pTF486-GhD14D:GFP was transiently expressed in *A. thaliana* protoplasts and the cells were stained with DAPI, and then examined using confocal microscopy. The results showed that GhD14D-GFP fusion protein was accumulated throughout the nucleus and cytoplasm and was also co-localized with DAPI (Fig. 2A). The pCHF3-GhD14D:GFP was bombarded into tobacco epidermal cells by agroinfiltration to further confirm the subcellular localization of GhD14D protein. We found the green fluorescence of GhD14D-GFP distributed in the cytoplasm and nucleus and the fluorescence of GhD14D-GFP coincided with the DAPI fluorescence, confirming that GhD14D protein is co-localized in the cytoplasm and nucleus (Fig. 2B). Both GhD14D-GFP fusion protein and DAPI methods in *A. thaliana* protoplast and tobacco epidermal cells clearly showed strong nucleus and cytoplasm localization, indicating that the hydrolase GhD14D functions in the cytoplasm and nucleus.

### Expression patterns of *GhD14A/D*

Expression patterns of *GhD14A/D* in fibers at 0, 3, 5, 10, 15 and 20 days post-anthesis (DPA) and root, stem, leaf, and flower were investigated to explore the function of *GhD14* using quantitative real-time PCR (qRT-PCR). Since *GhD14A* and *GhD14D* have 97.43% coding sequence similarity and they can't separate

with specific primer, the expression level of *GhD14A/D* was investigated. As shown in Fig. 2, *GhD14A/D* gene transcripts were increased during the fiber development with peak value at 20 DPA (Fig. 3A), and abundantly expressed in flower and stem (Fig. 3B), indicating that *GhD14A/D* may function in fiber, flower and stem development.

In order to test the tissue specific expression of *GhD14D*, the *GhD14D* promoter driving *GUS* gene expression in *A. thaliana* was visualized by histochemical staining of transgenic *A. thaliana*. The color of transgenic *A. thaliana* seedlings represents the promoter driven *GUS* gene expression. As shown in Fig. 4, the *GhD14D* promoter-driven *GUS* gene was highly expressed in buds, stems and flowers (Fig. 4). These observations demonstrated that *GhD14D* may play important role in buds, stems and flowers.

### ***GhD14A/D* gene silencing in cotton increased branch angles and reduced fiber length**

Virus-induced gene silencing (VIGS) is an efficient and rapid method to reduce gene transcripts and investigate gene functions in plants (Gu et al. 2014). To further uncover potential functions of *GhD14* in cotton, the VIGS strategy was used to reduce *GhD14A/D* transcript levels in *G. hirsutum*. Positive control of *GhPDS* gene silencing shows in Fig. S5. Expression levels of the *GhD14A/D* gene were detected using qRT-PCR strategy and the result showed that *GhD14A/D* gene expression was significantly reduced in *GhD14A/D*-silenced plants compared with that in the control plants (CLCrVA) (Fig. 5A). The phenotype of branch angles of control and VIGS plants (*GhD14A/D*-V1, *GhD14A/D*-V2 and *GhD14A/D*-V3) showed that the monopodial branch angles were significantly increased in *GhD14A/D*-silenced plants compared with the negative control plants (CK). The branch angles of *GhD14A/D*-silenced plants were increased about two times compared with the CK (Fig. 5B, C, Fig. S4). In order to explore the potential functions of *GhD14A/D* in cotton fiber development. The fiber length was observed after *GhD14A/D* silencing, and the result showed that the cotton fiber length was significantly reduced in *GhD14A/D*-silenced plants (Fig. 5D, E), indicating the *GhD14A/D* gene may play a critical role in cotton fiber elongation. In general, reducing *GhD14A/D* gene expression led to wider branch angles and decreased fiber length, suggesting that *GhD14A/D* functions in cotton architecture and fiber development.

### ***GhD14A/D* gene silencing in cotton reduce the transcripts of secondary cell wall biosynthesis genes**

The cotton fiber is produced by the specific elongation of cell in ovule epidermal and cell wall biosynthesis is required during the fiber elongation process. To further investigate the mechanism of *GhD14A/D* in regulating fiber development, the relative expression level of secondary cell wall biosynthesis genes (Sun et al. 2017) was investigated in *GhD14A/D*-silenced plants. As shown in Fig. 6, The six genes (*GhLBD30*, *GhCesA7*, *GhMYB46*, *GhXCP1*, *GhIRX8* and *GhXCP2*) related secondary cell wall biosynthesis were down regulated in *GhD14A/D*-silenced plant, while other three genes (*GhIRX10*, *GhCesA8* and *GhCesA4*) related secondary cell wall biosynthesis have no significant difference in expression levels compare with the negative control (CLCrVA) (Fig. 6). This result demonstrates that downregulating of *GhD14* gene expression reduced the transcripts of genes involved in secondary cell wall biosynthesis.

## Overexpression of *GhD14d* rescued the *d14* mutant phenotype in Arabidopsis

Overexpression of *GhD14D* in the *d14* Arabidopsis mutant, a heterologous complementation approach, was performed to further explore the function of *GhD14D* in stem development and shoot branching, since *AtD14* is the receptor of SLs and has essential functions in the SL signal transduction process (Chevalier et al. 2014). The pCAMBIA1305-35S-*GhD14D* was constructed and transformed into *A. thaliana* mutant *d14-1*. The seven transgenic lines were obtained after PCR detection (Fig. S6). The third generation of six transgenic lines with stable and highly expressed *GhD14A/D* was used for further analysis. As shown in Fig. 6, mutant *d14-1* plants had smaller length-width ratio of leaf (Fig. 7A, B) and more shoot branches as well as shorter plant height (Fig. 6C, D, E) compared with wild-type *A. thaliana* plants. Overexpression of *GhD14D* in *A. thaliana d14-1* mutant restored the leaf phenotypes and the number and length of branches to the phenotype of wild-type plants (Fig. 7). The phenotype of the *d14-1* mutant with more shoot branches and short stature could be rescued by *GhD14D* and there was no significant phenotype difference between wild-type *A. thaliana* lines and *d14-1/35S:GhD14D* transgenic lines (Fig. 7). These results suggested that *GhD14D* and *AtD14* may have similar functions in regulating branching number, plant height, petiole length, and length-width ratio of leaf.

## 4 Discussion

Cotton is an important source of protein and oil and cotton fiber is commonly used as natural fiber in the textile industry, hence, regulating growth and controlling branching are essential for cotton cultivation. Several key genes involved in the SL biosynthesis signaling pathway and participating in plant architecture development, especially shoot-branching have been identified (Beveridge et al. 2010). In this study, a homolog of *AtD14* was cloned from upland cotton cultivar Xuzhou 142 and named *GhD14s*. *GhD14s* showed high identity with *AtD14* (53.5%). The D14 functional domains were conserved in cotton and the model plants rice (monocotyledon) and Arabidopsis (dicotyledon) (Fig. 1). Phylogenetic analysis showed that the D14s in higher plants have a close relationship with D14s from other higher plants (Fig. S3). The conservation of D14 among different species indicates it play critical role in higher plants.

Subcellular localization results showed that *GhD14D* is localized to the nucleus and cytoplasm, which is consistent with the localization results of the D14 genes in Arabidopsis and rice (Chevalier et al. 2014; Yao et al. 2018). The subcellular localization results provided evidence that *GhD14D* functions in both the nucleus and cytoplasm. GUS staining showed that D14 in rice was mainly expressed in parenchyma cells surrounding the xylem in leaves, stems and axillary buds (Arite et al. 2009). In this study, the *GhD14D* pro::GUS gene expression was mainly observed in the buds, stems and flowers (Fig. 4). The transcripts of *GhD14D* in cotton were also significantly accumulated in stem, leaf, flower and 20 DPA fibers (Fig. 3). These results demonstrate that *GhD14D* functions in stem, leaf, flower and fiber development. Previous studies in other plants found that *D14* was involved in the development of stem, leaf and flower: The *d14* mutant had a larger stomatal aperture in leaf, slower abscisic acid (ABA)-mediated stomatal closure, lower anthocyanin and reduced plant senescence under drought stress (Li et al. 2020). OsMADS57

interacts with TEOSINTE BRANCHED1 (OsTB1) and targets OsD14 to control the outgrowth of axillary buds in rice (Guo et al. 2013).

The OsD14 is the receptor of SL protein, which acts as a new component of the SL-dependent branching inhibition pathway and inhibits rice tillering (Airte et al. 2009). AtD14 hydrolyzes SLs into a D-ring-derived intermediate CLIM and irreversibly binds CLIM to trigger SL signal transduction, thus regulating shoot branching in Arabidopsis (Yao et al. 2016). Furthermore, overexpression of *GhD14D* in the *d14* mutant, was performed to further explore the function of *GhD14D* in stem development and shoot branching. As a result, overexpression of *GhD14D* in *AtD14* mutant (*d14-1*) reduced the mean number of branches and restored the plant height (Fig. 7). Meanwhile, the branching and leaf phenotypes were restored to wild-type plants phenotype and there was no significant difference between wild-type plants and *d14-1/35S::GhD14D* transgenic lines. The phenotype of the *d14-1* mutant with fewer shoot branches and higher stature could be rescued via *GhD14D* overexpression in *d14* mutant, suggesting that *GhD14D* and *AtD14* may have similar functions in regulating branching number, plant height, petiole length, and length-width ratio of leaf. Consistently, the *OsD14* gene also restored the phenotype of *d14* to the wild type both in plant height and tiller development in rice (Yao et al. 2018).

The *GhD14A/D* had high expression levels in flower, stem, leaf, and 20 DPA fibers, but was weakly expressed in cotton root (Fig. 3). In rice, transcripts of *D14* were highly accumulated in leaves and the first leaf buds, but not in root tip (Airte et al. 2009). In petunia, high expression levels of *DAD2* were observed in axillary bud and leaf, but not in root (Hamiaux et al. 2012). In chrysanthemum, *DgD14* had the highest expression level in stem, followed by node, and was only weakly expressed in root (Wen et al. 2015). These results reflect that the *D14* gene may not be involved in root structural development. The *GhD14A/D* showed high expression level at 20 DPA fibers (Fig. 3), and this stage is the secondary cell wall thickening stage (Zhang et al. 2015). Meanwhile, silencing the *GhD14A/D* gene expression reduced the fiber length (Fig. 5) and the expression levels of the secondary cell wall biosynthesis related genes (Fig. 6). These results indicating that *GhD14* might play a role in fiber development through impact secondary cell wall biosynthesis of cotton fiber. In the future, molecular mechanisms and regulatory relationships between *GhD14* and secondary cell wall biosynthesis needs to be further investigated to deep our understanding of the strigolactone and secondary cell wall biosynthesis.

The *GhD14A/D* gene silencing enhanced fruit branch angles and reduced fiber length in cotton (Fig. 5), indicating that the *GhD14A/D* transcripts accumulation level affects cotton plant architecture and fiber length. The plant architecture is also regulated by auxin, cytokinin, and gibberellic acid (GA). D14 is the receptor of SL, which involved in plant stature and inhibition of plant shoot branching (Arite et al. 2009). The strigolactone signal can regulate the auxin polar transport and the cytokinin content in the stem (Ferguson et al. 2009). In the future, molecular mechanisms and regulatory relationships between *GhD14* and auxin or cytokinin should be examined to deepen our understanding of strigolactone signaling in cotton architecture. Although we did not investigate this, we speculate that *GhD14* might regulate branch number and angles through auxin and GA in cotton. In the future, molecular mechanisms and regulatory

relationships between *GhD14* and auxin or GA should be examined to deepen our understanding of SL signaling in cotton architecture.

## Conclusion

Strigolactone (SL) signaling functions directly in regulating plant development. DWARF14 (D14), a non-canonical SL receptor, interacts with the F-box protein MAX2 to modulate SL signaling in rice and *Arabidopsis*. In this study, the *GhD14D* was cloned from *G. hirsutum*, which containing a hydrolase fold motif, and had similar protein domains as typical D14 proteins. Subcellular localization analysis revealed that GhD14D was located in the cytoplasm and nucleus. qRT-PCR results showed that *GhD14A/D* was highly expressed in stem, flower and 20 DPA fiber cells; GUS staining assay indicated that *GhD14D* was mainly expressed in leaf primordium, inflorescence, axillary bud and stem. Reducing the transcripts of *GhD14A/D* in cotton significantly increased branch angle and reduced fiber length and the transcripts of secondary cell wall biosynthesis related genes. In addition, overexpression of *GhD14D* successfully rescued the phenotype of *Arabidopsis d14* mutant with reduced shoot-branching, length-width ratio of rosette leaves and plant height. These results indicate that *GhD14D* contributes to fiber development and plant architecture development in cotton, which will lay a foundation for manipulating cotton plant architecture by genetic engineering and provide a candidate gene for producing cultivars with ideal architectures in the future.

## Abbreviations

SL: Strigolactone; D14: DWARF14; MAX2: MORE AXILLARY GROWTH; DPA: days post-anthesis; WT: wild-type; bp: base pairs; MeJA: jasmonic acid methyl ester; NJ: Neighboring-Joining; VIGS: Virus-induced gene silencing; ABA: abscisic acid; MS: Murashige and Skoog; ORF: open reading frame

## Declarations

### Ethics approval and consent to participate

All the cotton lines used and analyzed for this study were collected from the Institute of Cotton Research of the Chinese Academy of Agricultural Sciences, which were public and available for non-commercial purpose. This article did not contain any studies with human participants or animals performed by any of the authors.

### Consent for publication

Not applicable.

### Availability of data and materials

All sequences data analyzed in this study are included in supplementary files. Other materials that support the findings of this study are available from the corresponding author on reasonable request.

### **Competing interests**

The authors have no conflict of interest to declare.

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### **Author contributions**

Conceptualization, J.Y. and G.X.; methodology, L.Z. and L.D.; software, LP.Z. and C.L.; validation, L.D., Z.T. and H.Z.; formal analysis, LP.Z. and Z.T.; investigation, C.L. and Z.T.; resources, H.Z.; data curation, L.Z. and L.D.; writing—original draft preparation, LP.Z. and G.X.; writing—review and editing, LP.Z.; visualization, L.Z. and Z.T.; supervision, G.X.; project administration, J.Y.; funding acquisition, G.X. All authors have read and agreed to the published version of the manuscript.

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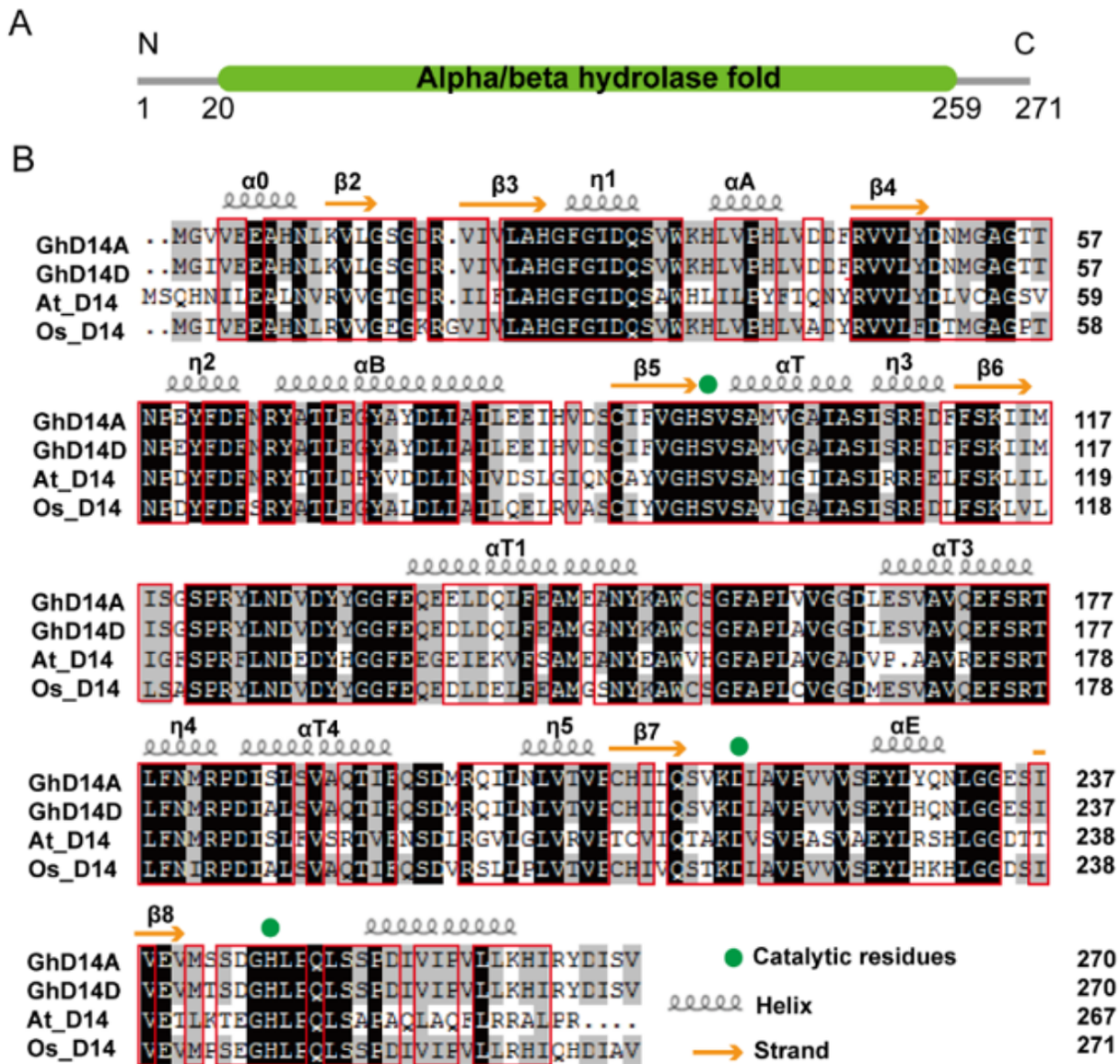
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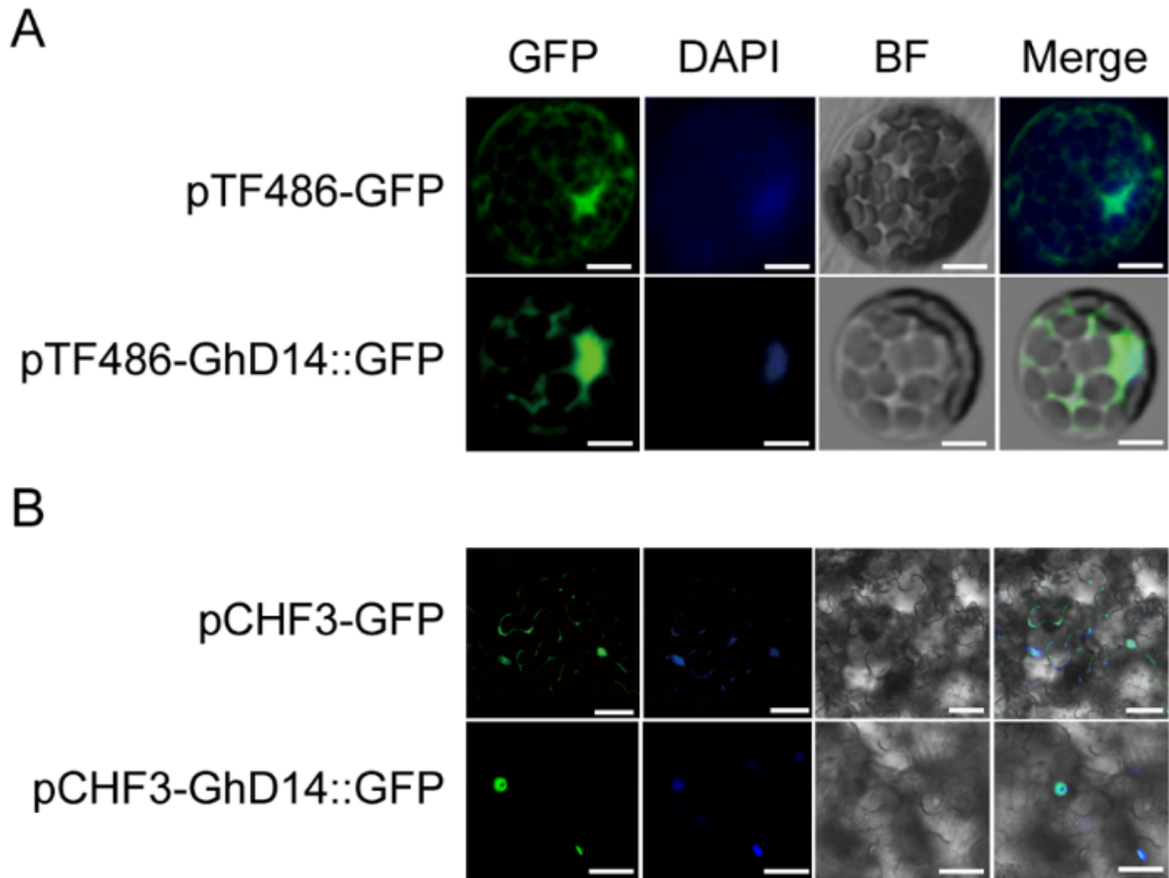
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## Figures



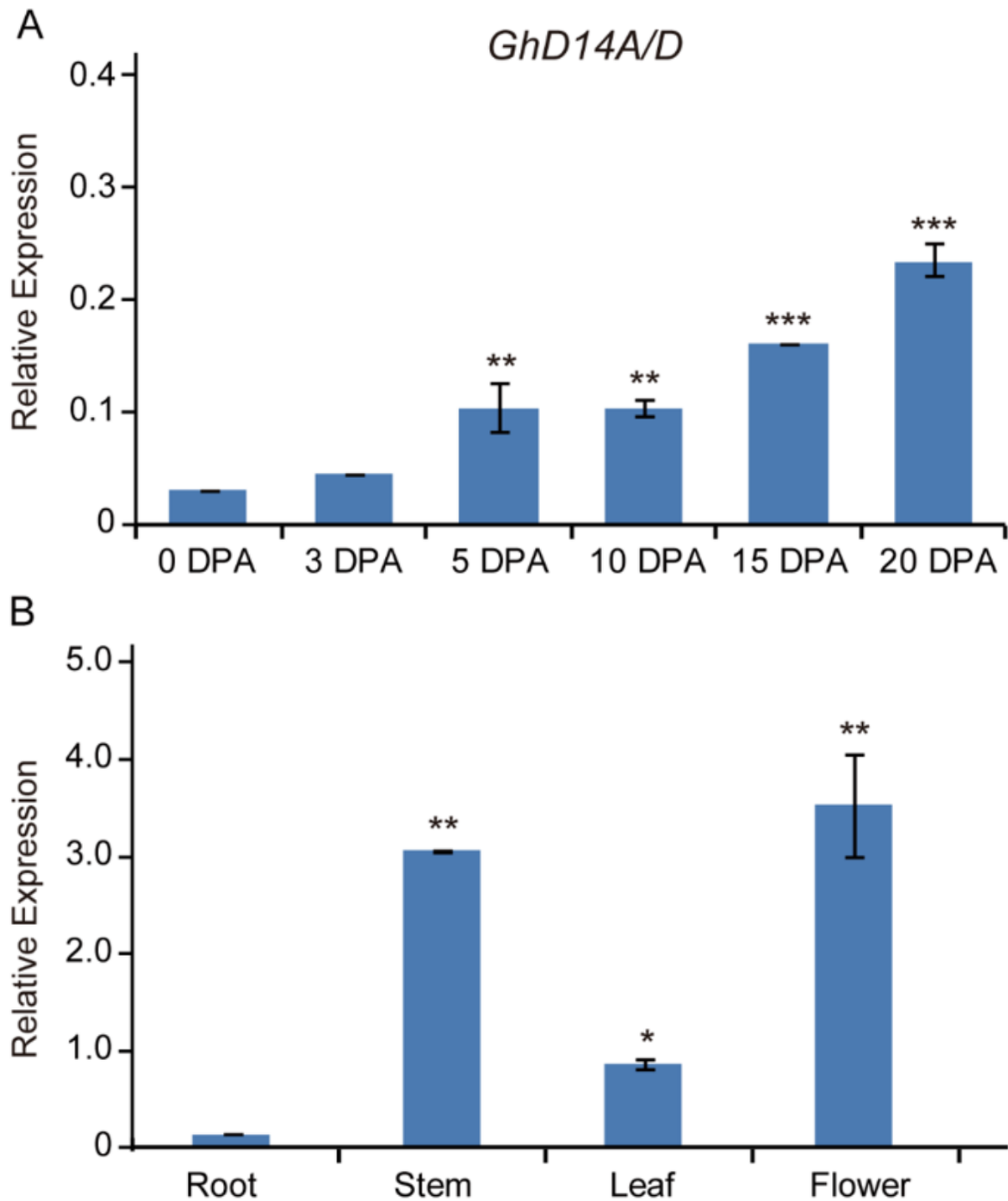
**Figure 1**

Multiple protein sequence alignment and structure diagram of D14 proteins. (a). D14 protein structure diagram. N indicates N-terminus; C indicates C-terminus. The green box indicates the alpha/beta hydrolase fold domain. The numbers below the protein represent the protein length and specific location of the domain. (b). Multiple protein sequence alignment of D14 proteins from cotton, Arabidopsis and rice. Secondary structure elements are displayed on top of the sequences. Identical and conserved residues are highlighted black and gray, respectively. The catalytic residues, helices and strands are indicated by green dots, curved lines and orange arrows, respectively. Numbers on the right indicate the sequence length.



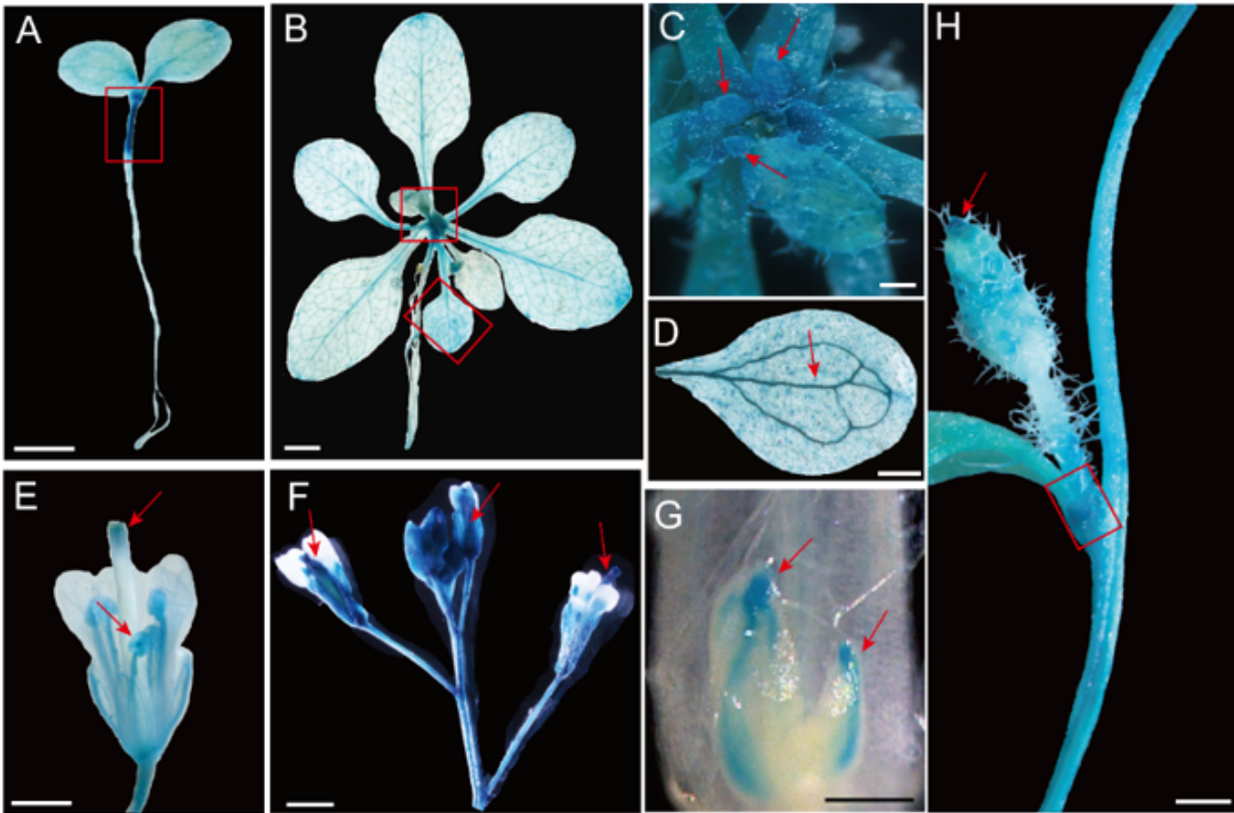
**Figure 2**

Subcellular localization of GhD14D protein. (a). Subcellular localization of GhD14D protein in Arabidopsis protoplasts. (b). Subcellular localization of GhD14D protein in tobacco epidermal cell. Pictures from left to right are GFP fluorescence, DAPI fluorescence, bright field (BF) and BF merged with DAPI and GFP fluorescence. Scale bars = 10  $\mu\text{m}$  (a) and 50  $\mu\text{m}$  (b).



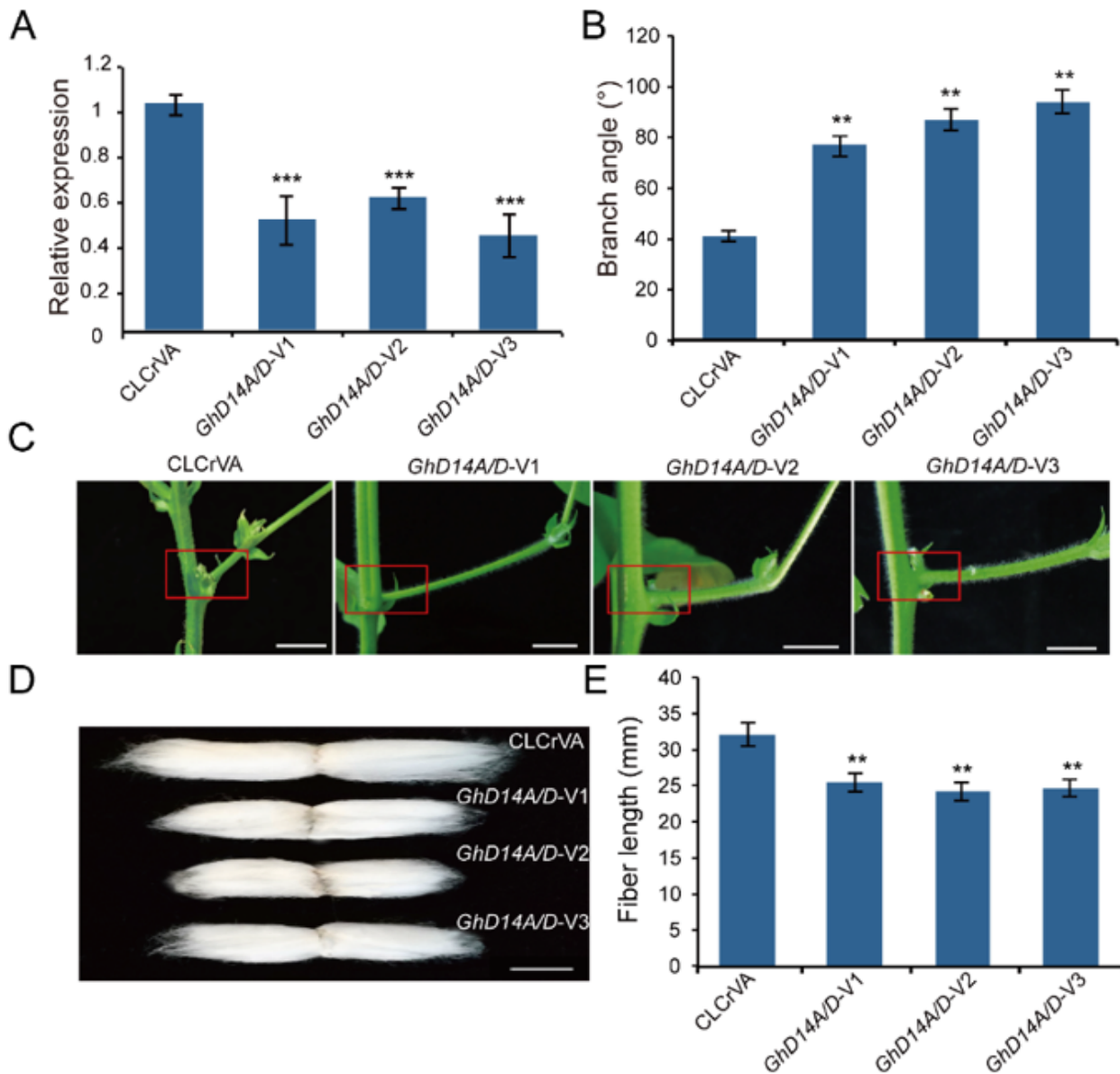
**Figure 3**

Expression analysis of GhD14A/D gene in different fiber development stages and tissues. The expression levels of GhD14A/D in fibers from 0 to 20 days post-anthesis (DPA) (a) and in different tissues (b). Statistical significance was determined using one-way ANOVA using 0 DPA samples (a) and root as the control (b), respectively. (\*  $P < 0.05$ ; \*\*  $P < 0.01$ ; \*\*\*  $P < 0.001$ ).



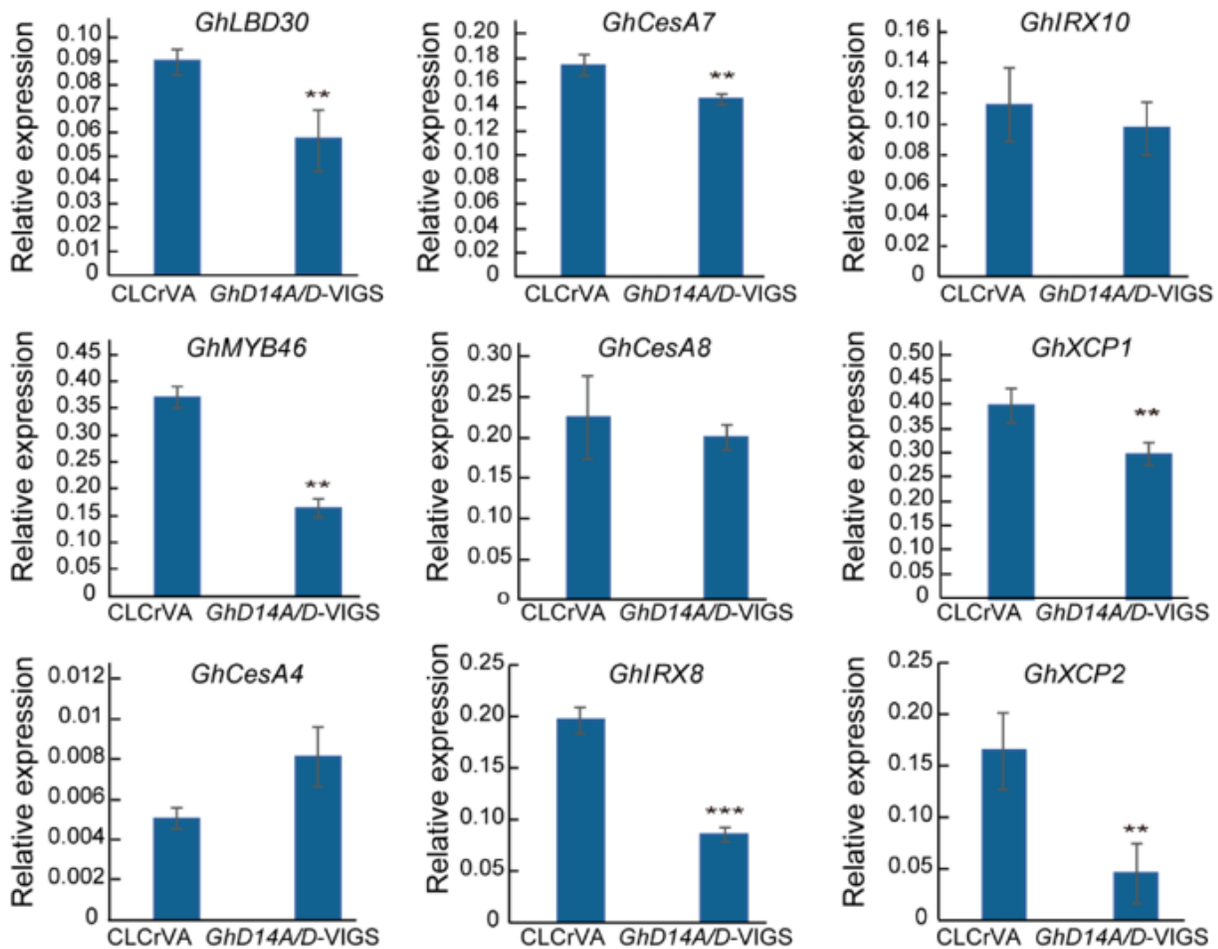
**Figure 4**

GUS staining assay of GhD14D gene promoter. The GUS staining result of 5-day-old transgenic seedlings (a), 15-day-old transgenic plant (b), leaf primordium (c), cotyledon (d), flower (e), inflorescence (f), axillary bud (g) and stem (h). Scale bars = 0.2 mm in (a), 0.5 mm in (b), 1 mm in (c), 1 mm in (d), 0.5 mm in (e), 2 mm in (f), 1 mm in (g), and 2 mm in (h). The red rectangles and arrows indicate GUS staining locations.



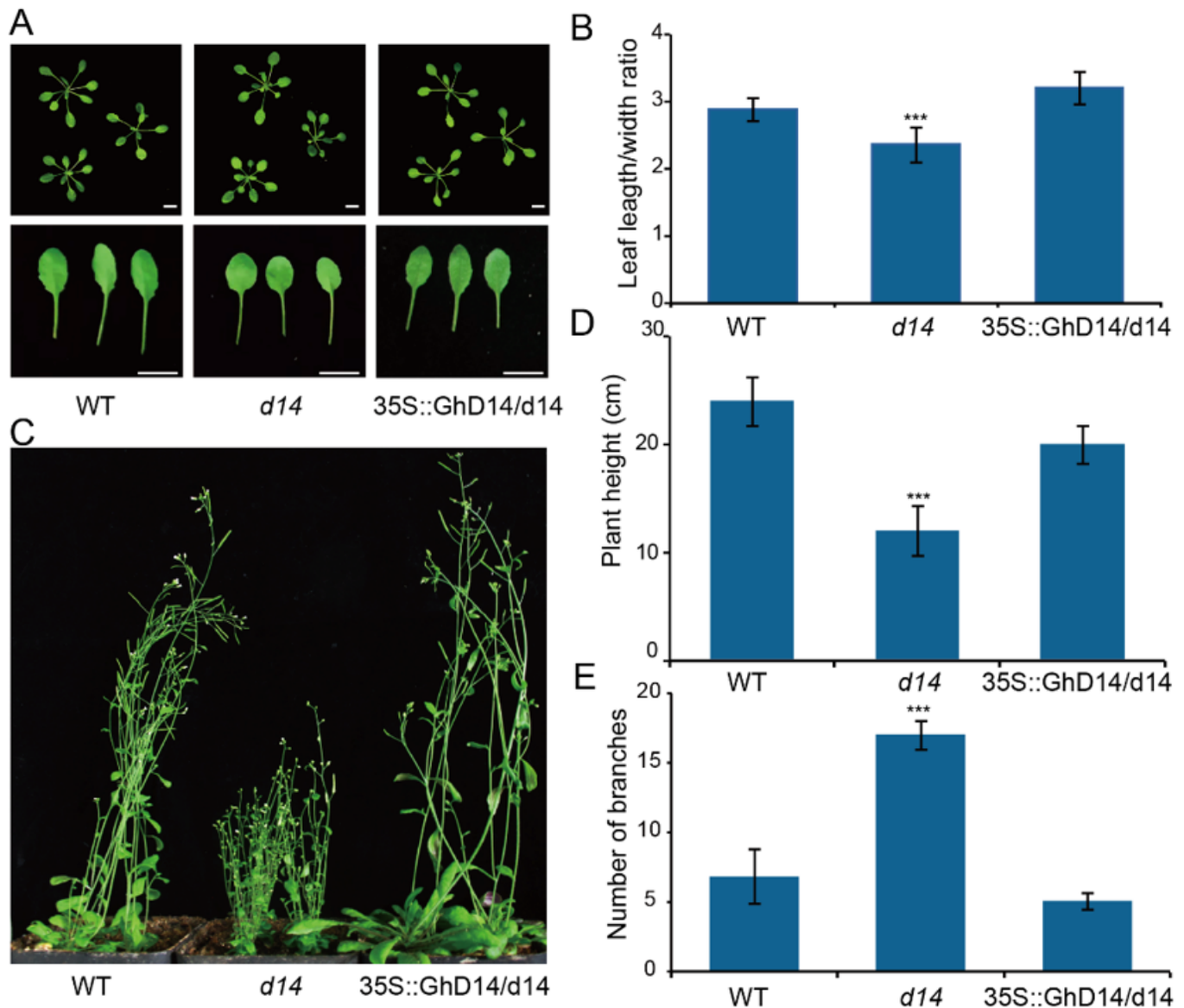
**Figure 5**

Silencing GhD14A/D modified plant architecture and reduced fiber length. (a). Relative expression levels of GhD14A/D in 25 DPA fibers from negative control (CLCrVA) and GhD14A/D-silenced cotton plants. (b). Average branch angles of negative control (CLCrVA) and GhD14A/D-silenced cotton plants. °, the degree of branch angle. (c). Branch phenotypes of negative control (CLCrVA) and GhD14A/D-silenced cotton plants. Red rectangles show the branch angles. Scale bars = 1 cm. (d). Representative seeds with attached fibers from negative control (CLCrVA) and GhD14A/D-silenced cotton plants. Scale bars = 1 cm. (e). Fiber length of negative control (CLCrVA) and GhD14A/D-silenced cotton plants. negative control (CLCrVA) and GhD14A/D-V1, GhD14A/D-V2, GhD14A/D-V3 represent cotton seedlings infiltrated with pCLCrVA empty vector and pCLCrVA-GhD14A/D vector, respectively. Error bars represent the mean  $\pm$  SE. Statistical significance was determined using one-way ANOVA. (\*\* P<0.01; \*\*\* P<0.001).



**Figure 6**

qRT-PCR analysis of secondary wall-associated formation genes. Relative gene expression of nine secondary cell wall biosynthesis genes in 20 DPA fibers of GhD14A/D-VIGS plants and negative control (CLCrVA) were calculated using GhUBQ7 as the internal reference gene. Error bars represent  $\pm$  SE of triplicate experiments. Asterisks indicate statistically significant differences, as determined using one-way ANOVA (\*P < 0.05; \*\*P < 0.01).



**Figure 7**

GhD14D recovered the d14 mutant phenotype in Arabidopsis. The rosette leaves (a) and plant height (c) phenotypes of WT, d14 mutant and GhD14D complementation line in d14 mutant background. Scale bars = 1 cm. Average length-width ratio (b), plant height (d) and average number of branches (e) of WT, d14 mutant and GhD14D complementation lines in the d14 mutant background. WT, wild type. Statistical analysis was performed with three biological replicates, and the error bars represent the mean  $\pm$  SE. Statistical significance was determined using one-way ANOVA. (\*  $P < 0.05$ ; \*\*\*  $P < 0.001$ ).

## Supplementary Files

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