

Genome-wide identification and characterization of the fibrillin gene family in *Triticum aestivum* (#41538)

1

Second revision

Guidance from your Editor

Please submit by **8 Mar 2020** for the benefit of the authors .



Structure and Criteria

Please read the 'Structure and Criteria' page for general guidance.



Author notes

Have you read the author notes on the [guidance page](#)?



Raw data check

Review the raw data. Download from the [materials page](#).



Image check

Check that figures and images have not been inappropriately manipulated.

Privacy reminder: If uploading an annotated PDF, remove identifiable information to remain anonymous.

Files

Download and review all files from the [materials page](#).

- 1 Tracked changes manuscript(s)
- 1 Rebuttal letter(s)
- 10 Figure file(s)
- 2 Table file(s)
- 2 Other file(s)



Structure and Criteria

Structure your review

The review form is divided into 5 sections. Please consider these when composing your review:

1. BASIC REPORTING
2. EXPERIMENTAL DESIGN
3. VALIDITY OF THE FINDINGS
4. General comments
5. Confidential notes to the editor

 You can also annotate this PDF and upload it as part of your review

When ready [submit online](#).

Editorial Criteria

Use these criteria points to structure your review. The full detailed editorial criteria is on your [guidance page](#).

BASIC REPORTING

-  Clear, unambiguous, professional English language used throughout.
-  Intro & background to show context. Literature well referenced & relevant.
-  Structure conforms to [PeerJ standards](#), discipline norm, or improved for clarity.
-  Figures are relevant, high quality, well labelled & described.
-  Raw data supplied (see [PeerJ policy](#)).

EXPERIMENTAL DESIGN

-  Original primary research within [Scope of the journal](#).
-  Research question well defined, relevant & meaningful. It is stated how the research fills an identified knowledge gap.
-  Rigorous investigation performed to a high technical & ethical standard.
-  Methods described with sufficient detail & information to replicate.

VALIDITY OF THE FINDINGS

-  Impact and novelty not assessed. Negative/inconclusive results accepted. *Meaningful* replication encouraged where rationale & benefit to literature is clearly stated.
-  All underlying data have been provided; they are robust, statistically sound, & controlled.
-  Speculation is welcome, but should be identified as such.
-  Conclusions are well stated, linked to original research question & limited to supporting results.

Standout reviewing tips

3



The best reviewers use these techniques

Tip

Support criticisms with evidence from the text or from other sources

Example

Smith et al (J of Methodology, 2005, V3, pp 123) have shown that the analysis you use in Lines 241-250 is not the most appropriate for this situation. Please explain why you used this method.

Give specific suggestions on how to improve the manuscript

Your introduction needs more detail. I suggest that you improve the description at lines 57- 86 to provide more justification for your study (specifically, you should expand upon the knowledge gap being filled).

Comment on language and grammar issues

The English language should be improved to ensure that an international audience can clearly understand your text. Some examples where the language could be improved include lines 23, 77, 121, 128 – the current phrasing makes comprehension difficult.

Organize by importance of the issues, and number your points

1. Your most important issue
2. The next most important item
3. ...
4. The least important points

Please provide constructive criticism, and avoid personal opinions

I thank you for providing the raw data, however your supplemental files need more descriptive metadata identifiers to be useful to future readers. Although your results are compelling, the data analysis should be improved in the following ways: AA, BB, CC

Comment on strengths (as well as weaknesses) of the manuscript

I commend the authors for their extensive data set, compiled over many years of detailed fieldwork. In addition, the manuscript is clearly written in professional, unambiguous language. If there is a weakness, it is in the statistical analysis (as I have noted above) which should be improved upon before Acceptance.

Genome-wide identification and characterization of the fibrillin gene family in *Triticum aestivum*

Yaoyao Jiang^{Equal first author, 1}, Haichao Hu^{Equal first author, 2}, Yuhua Ma³, Junliang Zhou^{Corresp. 3}

¹ School of Forestry and Biotechnology, Zhejiang Agriculture and Forestry University, Hangzhou, China

² College of Agriculture and Food Science, Zhejiang Agriculture and Forestry University, Hangzhou, China

³ Guizhou Institute of Pomological Sciences, Guizhou Academy of Agricultural Sciences, Guiyan, China

Corresponding Author: Junliang Zhou

Email address: gsszjl2008@163.com

Background. Fibrillin (*FBN*) is a highly conserved family of genes that is widely distributed in the photosynthetic organs of plants. Members of this gene family are widely involved in the growth and development of plants and their response to biotic and abiotic stresses. Wheat (*Triticum aestivum*), which is an important food crop, has a complex genetic background and little progress in the understanding of its molecular mechanism.

Methods. In this study, we identified 26 *FBN* genes in the whole genome of *T. aestivum* through bioinformatics. These genes were divided into 11 subgroups and distributed on 11 chromosomes of *T. aestivum*. Interestingly, most of TaFBN genes are located on the chromosomes 2A, 2B and 2D. The gene structure of each subgroup of gene family members and the position and number of motifs were highly similar. **Results.** The evolutionary analysis results indicated that the affinities of *FBNs* in monocots were closer together. Tissue-specific analysis revealed that *TaFBN* genes were expressed in different tissues and developmental stages. In addition, some *TaFBNs* were involved in one or more biotic and abiotic stresses. These results provide a basis for further study of the biological function of the *FBNs*.

Genome-wide identification and characterization of the fibrillin gene family in *Triticum aestivum*

Yaoyao Jiang^{#1}, Haichao Hu^{#1}, Yuhua Ma², Junliang Zhou^{2*}

¹. Zhejiang Agriculture and Forestry University, School of Forestry and Biotechnology, Hangzhou, Zhejiang, China

². Zhejiang Agriculture and Forestry University, College of Agriculture and Food Science, Hangzhou, Zhejiang, China

³. Guizhou Academy of Agricultural Sciences, Guizhou Institute of Pomological Sciences, Guiyan, Guizhou, China

*Corresponding: Zhou Junliang, <mailto:gsszjl2008@163.com> (JL.Z)

[#] YY.J. and HC.H. contributed equally to this work.

Abstract

Background. Fibrillin (*FBN*) is a highly conserved family of genes that is widely distributed in the photosynthetic organs of plants. Members of this gene family are widely involved in the growth and development of plants and their response to biotic and abiotic stresses. Wheat (*Triticum aestivum*), which is an important food crop, has a complex genetic background and little progress in the understanding of its molecular mechanism.

Methods. In this study, we identified 26 *FBN* genes in the whole genome of *T. aestivum* through bioinformatics methods. These genes were divided into 11 subgroups and distributed on 11 chromosomes of *T. aestivum*. Interestingly, most of *TaFBN* genes are located on the chromosomes 2A, 2B and 2D. The gene structure of each subgroup of gene family members and the position and number of motifs were highly similar.

Results. The evolutionary analysis results indicated that the affinities of *FBNs* in monocots were closer together. Tissue-specific analysis revealed that *TaFBN* genes were expressed in different

tissues and developmental stages. In addition, some *TaFBNs* were involved in one or more biotic and abiotic stresses. These results provide a basis for further study of the biological function of the *FBNs*.

Subjects: Bioinformatics, Genomics, Plant Science

Keywords: Fibrillin, *Triticum aestivum*, Abiotic stress, Gene duplication, Phylogenetic tree, *Cis*-regulatory elements

Introduction

Fibrillins (FBNs) are named after fibrils because these proteins were first detected in fibrils in the chromoplasts of dog rose (*Rosa rugosa*) and bell pepper (*Capsicum annuum*) fruit (Newman et al., 1989; Deruère et al., 1994; Kim et al., 2015). Since then, FBN proteins have been increasingly found in different organelles, including plastoglobules (PGs) in the chloroplasts and algal eyespots. Therefore, members of the FBN protein family have been given many different names, including plastid-lipid associated protein (PAP), plastoglobule (PGL), chloroplastic drought-induced stress protein of 34 kDa (CDSP 34), and chromoplast-specific carotenoid-associated protein (ChrC) (Pozueta-Romero et al., 1997; Ting et al., 1998; Kim et al., 2015). FBN proteins are located in the photosynthetic organs of cyanobacteria and some higher plants (Kim et al., 2015; Kim et al., 2017). Lundquist et al. (2012) identified 14 *FBN* genes in *Arabidopsis* by proteomic analysis, 50% of which were in PGs and others are mainly distributed in stroma and thylakoid membranes (Lundquist et al., 2012; Kim et al., 2015).

So far, the FBN protein family is mainly composed of 12 subfamilies; 11 of these have been found in higher plants and one has been identified in algae (Lohscheider and Río Bártulos, 2016; Kim et al., 2017). The members of these subfamilies were found to have similar hydrophobic structures; however, the biophysical properties of these proteins are quite diverse, including proteins with molecular weights of 20–42 kDa and isoelectric point (pI) values of 4–9 (Vidi et al., 2006; Lundquist et al., 2012). These findings suggest that each FBN protein may have specific

biological functions. In *Arabidopsis thaliana*, FBN proteins contain a conserved hydrophobic domain (lipocalin motif 1) in the N-terminus and amino acid residues near the C-terminus, including aspartic acid (Singh *et al.*, 2010). Furthermore, Lohscheider and Río Bártulos (2016) predicted that the three-dimensional structure of FBNs is similar to that of lipocalin, with the ability to bind and transport small hydrophobic molecules (Lohscheider and Río Bártulos, 2016), which suggests that the FBN family may have similar biological functions (Singh *et al.*, 2010; Francesc *et al.*, 2015; Kim *et al.*, 2015).

FBN proteins have a variety of important biological functions, such as participating in photosynthesis, the formation of lipoprotein structures, and responses to abiotic and biotic stresses (Kim *et al.*, 2015). Initially, researchers found that fibrillins located on the outer surface of red pepper chromoplast fibrils by Immunogold electron microscopy (Deruère *et al.*, 1994). Furthermore, fibril-like structures can be reconstituted *in vitro* from a mixture of FBN protein, lipids, and bicyclic carotenoids (Deruère *et al.*, 1994; Kim *et al.*, 2015). Compared to wild-type plants, RNAi-transgenic tomato plants with suppressed *LeChrC* (FBN1) accumulate 30% less carotenoids (Leitner-Dagan *et al.*, 2006; Singh *et al.*, 2010). In addition, when *FBN5* gene was deleted in *Arabidopsis thaliana* and rice, mutant plants were more sensitive to light stress and the levels of PQ-9 and PC-8 in the leaves were induced (Kim *et al.*, 2017). These results suggest that FBNs regulated the formation of chromoplast fibrils and the accumulation of carotenoids. In addition to structural roles, Fibrillin gene expression is also regulated by numerous abiotic and biotic stresses, especially oxidative stress (Youssef *et al.*, 2010). For example, the expression of the *Chrc* (FBN1) is induced in cucumber leaves infected with *Sphaerotheca fuliginea* (Leitner-Dagan *et al.*, 2006). Similar results have been reported when tomato plants infected with the fungus *Botrytis cinerea*. However, the expression patterns of fibrillins were varied and complexed during abiotic stress such as heat, cold, drought, high light and wounding treatment (Pruvot *et al.*, 1996; Kuntz *et al.*, 1998; Georg *et al.*, 2001; Leitner-Dagan *et al.*, 2006; Simkin *et al.*, 2008). *AtFBN1a* expression was induced and *AtFBN2* expression was repressed when subjected to drought or cold treatment (Laizet *et al.*, 2004). High levels expression of fibrillin

(*FBN1*) were observed in potato plants during water stress (Lee *et al.*, 2007). Similarly, the mutants of *pgl1* and *pgl2* were more sensitive to high light stress than was the wild-type in *Synechocystis* (Cunningham *et al.*, 2010). Moreover, when *LeChrC* (*FBN1*), *FBI4* and *AtFBN4* were knocked down in tomato, apple, and *Arabidopsis*, the mutant plants were more susceptible to the phytopathogenic fungus *Botrytis cinerea* and pathogenic bacteria *Erwinia amylovora* and *Pseudomonas syringae* pv. *tomato*, respectively (Cooper *et al.*, 2003; Leitner-Dagan *et al.*, 2006; Singh *et al.*, 2010). Meanwhile, *FBN* gene expression is regulated by hormones, including gibberellic acid, jasmonate, and abscisic acid, during plant growth and developmental stages, as well as when plants are subjected to stresses (Yang *et al.*, 2006; Youssef *et al.*, 2010; Kim *et al.*, 2017). The accumulation of *FBN* proteins were decreased in tomato *flacca* mutant plant, which is defective in ABA biosynthesis, subjected to drought stress. And the level of *FBN* protein can be induced when exogenous supply of ABA (Gillet *et al.*, 2001). Moreover, The *FBN1* and *FBN2* proteins are involved in the jasmonate biosynthesis pathway in response to light and cold stress (Youssef *et al.*, 2010). By contrast, *FBN1* mRNA and protein levels declined in red pepper fruit when treated with gibberellic acid (Deruère *et al.*, 1994).

Wheat (*Triticum aestivum* L.) is an important food crop that is widely grown around the world. Approximately 40% of the global population depends on *T. aestivum* as their staple food (Paux *et al.*, 2008; Han *et al.*, 2019). Common *T. aestivum* is a heterogenous hexaploid containing A, B, and D genomes; therefore, the genome information is large and complex (Ling *et al.*, 2013; Glover *et al.*, 2015; Han *et al.*, 2019). Moreover, owing to the complex genetic background of *T. aestivum*, only part of genes, which regulated important agronomic traits and molecular breeding, were reported in wheat. Therefore, the study of *T. aestivum* functional genomics is lagging far behind rice and corn. In recent years, high-quality wheat genome sequencing has been completed (International Wheat Genome Sequencing Consortium *et al.*, 2018); this will play an important role in elucidating the molecular mechanisms involved in growth and development, resistance, and high yield (Pradhan *et al.*, 2019; Rahimi *et al.*, 2019).

Although there is increasing evidence that *FBNs* play major roles in photosynthetic

organisms, to date, they have been identified and characterized in few plant species. In addition, the biological functional study of *T. aestivum* FBNs (*TaFBNs*) is limited in wheat. The identification and functional characterization of the *FBN* family in *T. aestivum* will contribute to elucidating the stress response mechanisms. In this study, we performed a genome-wide survey using the reported FBN protein sequences in the *T. aestivum* database. We identified 26 *FBN* genes in *T. aestivum* and used bioinformatic methods to analyze their biophysical properties, including gene structures and conserved motifs, as well as the chromosome distribution and gene duplication of *FBN* genes. In addition, we analyzed the expression profiles of *TaFBN* genes in different tissues, at different developmental stages, and in response to abiotic and biotic stresses using the *T. aestivum* expression database. These results may provide a basis for studying the biological function of the *FBN* gene in different growth and development stages of *T. aestivum*.

Materials & Methods

Plants material cultivation and treatments

The common *T. aestivum* cultivar “Chinese spring” was used in this study. *Triticum aestivum* seeds were sterilized with 1% NaOCl for 15 min, rinsed thoroughly with distilled water five times, and soaked in distilled water overnight at room temperature (18 °C). The seeds were transferred to filter paper and germinated for three days. The seedlings were cultured in nutrient solution and grown in a growth chamber with 16 h light (22 °C), 8 h dark (18 °C), and 50% humidity. The nutrient solution was replaced every three days at the growth stage. At 21 days old, seedlings were treated with 20% (m/V) PEG 6000 (Sigma-Aldrich, St. Louis, MO, USA) for 6 h. Untreated seedlings were used as a control, and each treatment contained three independent biological replicates. The roots, shoots, and leaves were collected separately for further analysis at 1 and 6 h after treatment.

Identification of *TaFBN* genes

We used protein sequences of *Arabidopsis thaliana* FBN (*AtFBN*) and *Oryza sativa* FBN (*OsFBN*) genes as queries to perform a BLAST (E-value $1e^{-10}$) search against the *T. aestivum*

genome database (genome assembly from IWGSC; <http://ensembl.gramene.org/>). We obtained a dataset of *TaFBN* sequences and filtered out the redundant sequences. The protein sequences of *AtFBN* and *OsFBN* genes were downloaded from the *Arabidopsis* Information Resource database (<https://www.arabidopsis.org/>) and the Rice Annotation Project database (<https://rapdb.dna.affrc.go.jp/>). Since a typical FBN protein is reported to contain a conserved PAP_fibrillin domain (PF04755), the online tools SMART (<http://smart.embl-heidelberg.de/>) and InterProScan (<http://www.ebi.ac.uk/interpro/>) were used to predict the functional domains of potential TaFBN proteins. To verify our results, all of the proteins were compared to the PAP_fibrillin domain using the HMMER 3.0 program with the default E-value ($E\text{-value} < 10^{-3}$). Proteins without the PAP_fibrillin domain were removed. The biophysical properties of the final TaFBN proteins were calculated using the ExPASy ProtParam tool (<https://web.expasy.org/>), including the theoretical values of pI, relative molecular mass, and the grand average of hydrophobicity (GRAVY). The subcellular localization of *TaFBNs* was analyzed using ProComp (<http://linux1.softberry.com>) and WoLF PSORT II (<https://www.genscript.com/wolf-psort.html>). In addition, the signal peptide and chloroplast transit peptides of *TaFBN* genes were predicted using the SignalP 4.1 server (<http://www.cbs.dtu.dk/services/SignalP-4.1/>) and ChloroP 1.1 server (<http://www.cbs.dtu.dk/services/ChloroP/>).

Multiple sequence alignments and phylogenetic analysis

Full-length protein sequences of *FBN* gene family members identified in 13 plant species, including eight monocotyledon species and five dicotyledon species, were downloaded from the NCBI database (<https://www.ncbi.nlm.nih.gov/>), the Ensembl Plants database (genome assembly from IWGSC; <http://ensembl.gramene.org/>), and the Phytozome v12.1 database (<https://phytozome.jgi.doe.gov/pz/portal.html>). Full-length protein sequences of these *FBN* genes were aligned using MAFFT software (<https://mafft.cbrc.jp/alignment/server/>). Based on FASTA files, a neighbor-joining phylogenetic tree was constructed using Molecular Evolutionary Genetics Analysis (MEGA) version 7.0 software with 1000 bootstrap replicates.

Analysis of gene structures and conserved motifs

To investigate the structure of *TaFBN* genes, we used the Gene Structure Display Server 2.0 database (<http://gsds.cbi.pku.edu.cn/>) to analyze the distribution of exons and introns in *TaFBN* genes. Conserved motifs were predicted using the Multiple EM for Motif Elicitation (MEME) database (<http://alternate.meme-suite.org/>; the number of motifs was set to 10 and the motif width was set to 6–50).

Chromosome distribution of FBN genes in *T. aestivum*

The *TaFBN* gene distribution map was analyzed based on the sequencing genome information of the *TaFBN* gene family. The information on the position of the *TaFBN* gene family on chromosomes was obtained from the Ensembl Plants database (<http://plants.ensembl.org/>). The distribution of *FBN* genes on chromosomes were plotted using TBtools software (<https://github.com/CJ-Chen/TBtools>) (*Chen et al., 2018*).

Analysis of the *cis*-regulatory element of *FBN* gene promoters

In this study, 2000-bp sequences upstream of translational start sites of *TaFBN* genes were set as promoter sequences. PlantCARE software (http://bioinformatics.psb.ugent.be/webtools/plant_care/html/) was used to predict the *cis*-regulatory elements based on these promoter sequences. The distribution of *cis*-regulatory elements in the promoter of the *TaFBN* gene was displayed using TBtools software (<https://github.com/CJ-Chen/TBtools>) (*Chen et al., 2018*).

Analysis of *TaFBN* gene expression patterns

The expression profile data used in this study were obtained via the Wheat Expression Browser database (<http://www.wheat-expression.com/>) (*Philippa et al., 2016; Ricardo et al., 2018*). We searched for *FBN* genes on the website using the gene ID as query terms. The expression of *TaFBNs* in different tissues, at different developmental stages, and under different abiotic and biotic stress conditions (including drought, cold, heat, and stripe rust) were analyzed. The results were presented as heatmaps, with different colors representing the absolute signal values. The color scale of the heatmap was given in log₂ ratio values. The cultivar used in the gene expression profiles analysis was “Chinese spring”.

Total RNA isolation and real-time PCR analysis

Total RNA from different tissues was extracted using TRIzol Reagent (Invitrogen). The total RNA was treated with RNase-free DNase I for 15 min to remove the remaining genomic DNA. First-strand cDNA was synthesized according to the manufacturer's instructions (TOYOBO, Kita-ku, Osaka, Japan), diluted 20 times, and used as a template for quantitative real-time PCR (qRT-PCR), which was performed using AceQ qPCR SYBR Green Master Mix (Vazyme, Nanjing, China). For an endogenous control, we used the *T. aestivum actin* gene (AB181991). At least three biological replicates, with three technical replicates each, were used for each treatment. Relative expression levels were calculated using the comparative $2^{-\Delta\Delta Ct}$ method (Willemse *et al.*, 2008). The *TaFBN* primers used for qRT-PCR are listed in Table S1.

Results

Identification and characterization of *FBN* genes in *Triticum aestivum*

In this study, a total of 26 *FBN* genes were identified in *T. aestivum*, which we named *TaFBN-A1–TaFBN-D10* according to their genome location (Table 1). The *TaFBN* characteristics, including the chromosomal position, intron number, gene length, number of amino acids, molecular mass, CDS, subcellular localization, signal peptide, and instability index, are listed in Table 1. As shown in Table 1, the *TaFBN* protein sequences ranged from 219 to 402 amino acids and the molecular weights ranged from 23.75 to 43.59 kDa. The prediction of subcellular location indicated that 18 *TaFBNs* were located in the chloroplasts and eight were located extracellularly. At present, the GRAVY values is an important indicator to measure the hydrophobicity of protein; the GRAVY values of almost all *TaFBN* proteins are less than 0, except for *TaFBN-A1*, *TaFBN-B1*, and *TaFBN-B6*. Meanwhile, the prediction results showed that no signal peptides were found in any *TaFBN* proteins, but all *TaFBN* proteins contain chloroplast transit peptides.

Gene structure analysis of *TaFBN* genes

To gain insight into the evolution of the *TaFBN* gene family, a diagram of the *TaFBN* exon–intron gene structure was constructed based on cDNA and genomic DNA sequence

information (Supplementary text 1) using the Gene Structure Display Server (Figure. 1b). A neighbor-joining phylogenetic tree was also constructed to explore the evolutionary relationship and the phylogenetic classification of *FBN* gene in wheat. Gene structure analyses indicated that homologous genes had similar exon–intron distribution patterns (Fig. 1b). However, the number of introns in different *TaFBN* gene family members varied greatly (ranging from 2 to 10 introns), while there was almost no difference between members of the same subfamily. We found that all *TaFBN* genes contain a conserved PAP_FBN domain (PF04755), and the distribution of domains was consistent with the genetic homology (Fig. 1c). These results suggested that members of the same subfamily may have similar biological functions. In addition, we used the MEME online tool to analyze the conserved motifs of *TaFBN* genes; the results showed that all *TaFBN* members contained five to nine conserved motifs (Fig. 2). The logo representation of the 10 conserved motifs identified for proteins encoded by *TaFBN* genes is described in Supplementary Fig. 1. Figure 2 showed that motif 1, motif 2, motif 3, motif 4, and motif 5 were highly conserved and widely distributed in all *TaFBN* proteins. Motif/domain analysis revealed that motif 1 contained conserved amino acid residues in the C-terminal and motif 3 contained a conserved lipocalin motif (Supplementary Fig. 2). The types and distribution of conserved motifs may be the reason for the functional diversity of *TaFBNs*.

Phylogenetic and evolutionary analysis of *TaFBN*

An unrooted phylogenetic tree was constructed for 179 *FBN* genes from eight monocotyledon species (with 26 *FBNs* from *T. aestivum*, 9 from *Oryza sativa*, 11 from *Zea mays*, 10 from *Sorghum bicolor*, 9 from *Panicum hallii*, 20 from *Panicum virgatum*, 10 from *Setaria italica*, and 4 from *Hordeum vulgare*) and five dicotyledon species (with 14 *FBNs* from *A. thaliana*, 12 from *Brassica oleracea* var. *capitata*, 11 from *Nicotiana tabacum*, 21 from *Glycine max*, and 22 from *Coffea arabica*) to study the evolutionary relationships of *TaFBN* members (Fig. 3). Based on the *FBN* gene characteristics of *A. thaliana*, these *FBN* genes can be classified into 11 subfamilies (Group 1 to Group 11). Interestingly, the members of *TaFBNs* were identified into nine subfamilies, each subfamily containing two or three *FBN* genes. The analysis

also revealed that the *FBN* genes in monocots (i.e., *T. aestivum*, *O. sativa*, *Z. mays*, *P. hallii*, and *S. bicolor*) were more closely related than those of the dicots (i.e., *A. thaliana*, *B. oleracea* var. *capitata*, and *N. tabacum*).

Analysis of *TaFBN* cis-regulatory elements

To further identify the *cis*-regulatory elements located upstream of the *TaFBN* genes, 2000-bp sequences upstream from translational start sites of putative *TaFBN* gene families were analyzed using the PlantCARE tool. As shown in Fig. 4, many *cis*-regulatory elements were identified in the promoters of *TaFBN* genes. These *cis*-regulatory elements can be divided into three types: hormone response elements, stress response-related elements, and light response-related elements. The hormone response elements, including the methyl jasmonate (MeJA)-responsive, abscisic acid-responsive, gibberellin-responsive, salicylic acid-responsive, and auxin-responsive elements, were widely distributed in promoters of the *TaFBN*s. The responses to abiotic stress were the light response-related, low temperature response-related, and drought stress-related response elements, respectively. These results suggested that *TaFBN* genes may be involved in photosynthesis, stress responses, and in maintaining the hormone balance in plants, thereby improving the chances for organisms to escape or better cope with the damaging effects of adverse environmental conditions.

Chromosomal location of *TaFBN* in the *T. aestivum* genome

To clarify the distribution of *TaFBN* genes on *T. aestivum* chromosomes, the 26 *TaFBN* genes were mapped onto *T. aestivum* chromosomes; these *TaFBN* genes were distributed on 11 chromosomes (Fig. 5). As shown in Fig. 5, we found that *TaFBN* genes were not randomly distributed on the chromosome; chromosomes 2A, 2B, and 2D contained six *TaFBN* genes; however, the other chromosomes (i.e., 1A, 1B, 1D, 4A, 4B, 4D, 5A, and 5B) had only one *TaFBN* gene. The unequal distribution of *TaFBN* genes on the chromosomes may be caused through ancient gene duplications.

Tissue specific expression patterns of *TaFBN*s at different developmental stages

To explore the tissue-specific expression patterns of *TaFBN* genes at different growth and

developmental stages in *T. aestivum*, publicly available expression data sets for 26 *TaFBN*s were analyzed to examine the transcription levels in various *T. aestivum* tissues, including the root, shoot, anther, spikelet, and leaf. Most of the *TaFBN* genes can be detected in at least two or more different tissues. The results suggested that *TaFBN* genes may be widely expressed in wheat tissues (Fig. 6a). However, the expression levels of *TaFBN* genes varied among the different tissues. The expression levels of *TaFBN*s in the tissues with high chlorophyll contents (leaf, shoot, and internode) were significantly higher than those in other tissues. As shown in Fig. 6b, the expression levels of *TaFBN* were notably different at different developmental stages. *TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, *TaFBN-B2*, *TaFBN-D2*, *TaFBN-A3*, *TaFBN-A6*, *TaFBN-B6*, and *TaFBN-D6* were highly expressed at all developmental stages. However, the expression levels of *TaFBN-B4*, *TaFBN-D5*, *TaFBN-A9*, *TaFBN-B9*, and *TaFBN-D9* were inhibited at all developmental stages. The expression levels of other *TaFBN* genes did not change significantly during any of the developmental stages. These data indicated that *TaFBN* genes have tissue-specific expression patterns, and some *TaFBN* genes play a vital role in the growth and developmental stages of *T. aestivum*.

Expression profiles of *TaFBN* genes in response to abiotic stresses

To further clarify the potential functions of *TaFBN* genes under abiotic stress responses, the expression levels of *TaFBN* genes were analyzed under drought, stripe rust, cold and heat conditions. Most of the *TaFBN* genes could be shown to be involved in the response to one or more abiotic stresses (Fig. 7). The transcripts of *TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, *TaFBN-B2*, *TaFBN-D2*, and *TaFBN-B6* were significantly upregulated by drought, stripe rust, cold, and heat treatments. However, the expression levels of *TaFBN-A5*, *TaFBN-B5*, *TaFBN-D5*, *TaFBN-A9*, *TaFBN-B9*, *TaFBN-D9*, *TaFBN-A10*, *TaFBN-B10*, and *TaFBN-D10* were slightly downregulated under drought, stripe rust and heat stresses. In addition, most of *TaFBN* genes were up-regulated after 12h of drought treatment and 11 days of stripe rust infection, respectively. Interestingly, almost all *TaFBN* genes had a high level of expression under cold stress. In addition, other *TaFBN*s can be induced to express under some of the stress conditions. The transcription levels

of the tested *TaFBN* genes were significantly downregulated under drought stress conditions. These results indicated that *TaFBN* genes might participate in response to abiotic stresses, especially drought, stripe rust, cold and heat stress in *T. aestivum*.

Validation of *TaFBNs* by qRT-PCR

To further detect the expression level of *TaFBN* genes in different tissues, we selected nine representative genes from the *TaFBN* gene family (*TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, *TaFBN-B2*, *TaFBN-D2*, *TaFBN-B5*, *TaFBN-B6*, *TaFBN-A9*, *TaFBN-B9*, and *TaFBN-D9*) based on their expression profile, and analyzed the expression level using qRT-PCR (Fig. 8a). The results showed that the expression of nine *TaFBNs* in the leaves and shoots was significantly higher than that in the roots. We also analyzed the *TaFBN* gene expression in the leaf under drought stress in *T. aestivum* seedlings (Fig. 8b). The results suggest that the expressions of some *TaFBN* genes, such as *TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, *TaFBN-B2*, *TaFBN-D2* and *TaFBN-B5*, were induced at different time points under drought stress. However, *TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, and *TaFBN-B2* displayed downregulation after drought treatment. In addition, as the treatment time increased, the expression level was significantly upregulated or downregulated. These results are consistent with the data of the above expression profiles.

Discussion

In this study, we identified 26 *FBN* genes in the *T. aestivum* genome. These genes were distributed on 11 chromosomes and had molecular masses ranging from 23.75 to 43.59 kDa and pI values ranging from 4.59 to 9.61. This diversity suggests that *TaFBN* genes may have specific biological functions in different metabolic processes. Furthermore, the study results indicate that most of the *TaFBN* genes are located on the chloroplast and contained chloroplast transit peptides. This provides strong evidence that various *FBNs* might participate in photosynthesis. GRAVY was used to calculate the overall hydrophobicity of the protein sequence, with higher positive GRAVY values indicating a greater level of hydrophobicity (Faya *et al.*, 2015). Almost all of the *TaFBN* genes' GRAVY values were less than 0, which means that most of the proteins are hydrophilic. In contrast, previous studies have reported that the *FBN* family can bind to and

transport small hydrophobic molecules in *A. thaliana* (Singh and McNellis., 2011; Kim et al., 2015). However, the specific spatial structure and the percentage of hydrophobic residues may affect the hydrophobicity of proteins (Dyson et al., 2004). Therefore, these different results may reflect the biological function diversity of *TaFBN* genes.

The phylogenetic tree analysis showed that the number of *FBN* gene family members in *T. aestivum* was higher than those in the other monocots and dicots in this study. In the other hands, gene family members of *FBN* in wheat are always clustered together with rice, maize and sorghum. However, the members of *FBN* gene in *P. hallii*, *P. virgatum*, *S. italic* and *H. vulgare* have a distant evolutionary relationship with *TaFBN* genes. And surprisingly, only four members of *FBN* gene family have been found in *H. vulgare* genome. Based on these phenomena, we believe that wheat, rice, maize and others monocots have a common ancestor. The evolutionary separation time of wheat and rice may be closer than others species such as *P. hallii*, *P. virgatum*, *S. italic* and *H. vulgare*. However, because the genome sequences of some species are incomplete, more evidences is needed to support these conclusions. To analyze the evolutionary relationships of *FBN* genes, we constructed a phylogenetic tree with 179 *FBNs* from *T. aestivum*, *O. sativa*, *S. bicolor*, *Z. mays*, *P. hallii*, *P. virgatum*, *S. italica*, *H. vulgare*, *A. thaliana*, *B. oleracea* var. *capitate*, *N. tabacum*, *G. max*, and *C. arabica*. These *FBN* genes were divided into 11 subfamilies using the classification method described for *FBN* in *A. thaliana* (Singh and McNellis., 2011). Interestingly, the exon–intron structures and numbers of conserved motifs were similar in the same subgroups. This phenomenon suggests that *TaFBN* genes in the same subgroup may have similar functions.

Gene expression levels in different tissues and at different developmental stages may be determined by gene function. Previous studies have shown that *FBNs* are regulated by a variety of biological and environmental factors at different growth and developmental stages (Singh and McNellis, 2011). We analyzed the expression patterns of the *TaFBN* gene family during different growth and development stages and under biotic and abiotic stresses in *T. aestivum* through publicly available expression data. In these studies, we obtained 26 *TaFBN* gene expression

349 profiles, which showed that most of the genes were highly expressed in the leaf, shoot, and
 350 internode. Similar results have been reported in potato, *Arabidopsis*, and *Brassica rapa* (Monte et
 351 al., 1999; Kim et al., 2001; Yang et al., 2006). Furthermore, the expression profile data
 352 suggested that *TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, *TaFBN-B2*, *TaFBN-D2*, and *TaFBN-B6*
 353 expressions were strongly induced under drought, stripe rust, cold, and heat stresses, but *TaFBN-*
 354 *A5*, *TaFBN-B5*, *TaFBN-D5*, *TaFBN-A9*, *TaFBN-B9*, *TaFBN-D9*, *TaFBN-A10*, *TaFBN-B10*, and
 355 *TaFBN-D10* expressions were slightly inhibited under these stresses. In addition, other *TaFBNs*
 356 responded to one or more stresses. These is some evidences that fibrillin protein, such *FBN1a*,
 357 *FBN1b* and *FBN2*, level increases in leaves of rice, *Arabidopsis*, *Brassica* and potato plants
 358 subjected to drought and cold stress (Gillet et al., 2001; Kim et al., 2001; Laizet et al., 2004; Lee
 359 et al., 2007). Furthermore, the accumulation of FBN1 protein in tomato *flacca* mutant, which is
 360 defective in ABA biosynthesis, was significantly declined than wild type during drought stress
 361 (Gillet et al., 2001). We get similar results in wheat plants subjected to drought stress. It is
 362 possible that fibrillin gene expression was regulated through endogenous ABA concentration in
 363 response to numerous stresses (Singh and McNellis, 2011). As we know, transcription factors
 364 participate in various biological processes by regulating the expression of downstream gene *cis*-
 365 regulatory elements (Ning et al., 2017). In this study, many *cis*-regulatory elements were
 366 detected in the promoter sequences of *TaFBN* genes. These elements contained light response-
 367 related elements, drought response-related elements, and hormone response elements, such as
 368 MeJA, abscisic acid, gibberellic acid, salicylic acid, and auxin. Interestingly, all of the *TaFBN*
 369 genes include many light response-related elements. For example, Rey et al. (2000) found that
 370 overexpressing *FBN1* can promote plant height and flowering under high light levels in tobacco
 371 (Rey et al., 2000; Singh and McNellis., 2011). Leitner-Dagan et al (2006) studies show that
 372 fibrillin gene expression and carotenoid accumulation in flower tissue of cucumber were
 373 increased during GA treatment, and nice GA-responsive elements were found on FBN promoter
 374 sequences (Leitner-Dagan et al., 2006). By contrast, auxin (IAA) can delay the accumulation of
 375 FBN protein in bell pepper fruit, but abscisic acid (ABA) can promote this process (Deruère et

al., 1994; Singh and McNellis, 2011). Although the expression patterns of *TaFBN* genes were varied and **complexed**, overall, these genes had similar functions in plant stress resistance and chromoplast development (Singh and McNellis, 2011).

Conclusion

In this study, we identified 26 *FBN* genes in *T. aestivum* using a genome-wide screening approach. Based on their phylogenetic relationships, these *FBN* genes were classified into 11 subfamilies. The *TaFBN* gene structures and conserved motifs were highly conserved in the same subgroup. Many *cis*-regulatory elements were found in the *TaFBN* gene promoter sequences, which showed that the expression of *TaFBN* genes was regulated by various hormones and environmental factors. Moreover, almost all *TaFBN* genes were highly expressed in the leaf, shoot, and internode. The expression profiling data suggest that *TaFBN-A1*, *TaFBN-B1*, *TaFBN-A2*, *TaFBN-B2*, *TaFBN-D2*, and *TaFBN-B6* were responsive to many biotic and abiotic stresses. These results can help us to clarify the structural and functional relationships among *TaFBN* gene family members.

References

- Chen CJ, Xia R, Chen H, He YH. 2018.** TBtools, a Toolkit for Biologists integrating various HTS-data handling tools with a user-friendly interface. *bioRxiv*. DOI: 10.1101/289660.
- Cunningham FX, Tice AB, Pham C, Gantt E. 2010.** Inactivation of genes encoding plastoglobulin-like proteins in *Synechocystis* sp. pcc 6803 leads to a light-sensitive phenotype. *Journal of Bacteriology* **192**(6):1700-1709 DOI:10.1128/JB.01434-09.
- Data Availability.Cooper B, Clarke JD, Budworth P, Kreps J, Hutchison D, Park S. 2003.** A network of rice genes associated with stress response and seed development. *Proceedings of the National Academy of Sciences of the United States of America* **100**(8):4945-4950 DOI:10.1073/pnas.0737574100.

- Deruère J, Römer S, d'Harlingue A, Backhaus RA, Kuntz M, Camara B. 1994.** Fibril assembly and carotenoid overaccumulation in chromoplasts: a model for supramolecular lipoprotein structures. *Plant Cell* **6**: 119–133 DOI:10.2307/3869680.
- Dyson MR, Shadbolt SP, Vincent KJ, Perera RL, McCafferty J. 2004.** Production of soluble mammalian proteins in *Escherichia coli*: identification of protein features that correlate with successful expression. *BMC Biotechnology* **4**:32 DOI:10.1186/1472-6750-4-32.
- Francesc HG, Albert B. 2015.** A hydrophobic proline-rich motif is involved in the intracellular targeting of temperature-induced lipocalin. *Plant Mol Biol* **88**:301-311 DOI 10.1007/s11103-015-0326-x.
- Faya N, Penkler DL, Tastan Bishop Ö. 2015.** Human, vector and parasite hsp90 proteins: a comparative bioinformatics analysis. *FEBS Open Bio* **5**(1):916-927 DOI:10.1016/j.fob.2015.11.003.
- Georg L, Nathalie Mh, Mélanie B, Stephan C, Noëlle B, Marcel K, Pascal, Rey. 2001.**
 1. Accumulation of plastid lipid - associated proteins (fibrillin/CDSP34) upon oxidative stress, ageing and biotic stress in Solanaceae and in response to drought in other species. *Journal of Experimental Botany* (360):360 DOI:10.1093/jexbot/52.360.1545.
- Glover NM, Daron J, Pingault L, Vandepoele K, Paux E, Feuillet C, Choulet F. 2015.**
 5. Small-scale gene duplications played a major role in the recent evolution of wheat chromosome 3B. *Genome Biology* **16**(1):188 DOI:10.1186/s13059-015-0754-6.
- Gillet B, Beyly A, Peltier G, Rey P. 2001.** Molecular characterization of *CDSP 34*, a chloroplastic protein induced by water deficit in *Solanum tuberosum* L. plants, and regulation of *CDSP 34* expression by ABA and high illumination. *Plant Journal* **16**(2):257-62 DOI:10.1046/j.1365-3113x.1998.00292.x.
- Han Z, Liu Y, Deng X, Liu D, Liu Y, Hu Y. 2019.** Genome-wide identification and expression analysis of expansin gene family in common wheat (*Triticum aestivum* L.). *BMC Genomics* **20**(1):101 DOI:10.1186/s12864-019-5455-1.
- International Wheat Genome Sequencing Consortium. 2018.** Shifting the limits in wheat

at research and breeding using a fully annotated reference genome. *Science* **361** eaar7191 DOI:10.1126/science.aar7191.

Kim EH, Lee Y, Kim HU. 2015. Fibrillin 5 is essential for plastoquinone-9 biosynthesis by binding to solanesyl diphosphate synthases in *Arabidopsis*. *The Plant Cell* **27**: 2956–2971 DOI:10.1105/tpc.15.00707.

Kim HU, Wu SS, Ratnayake C, Huang AH. 2001. *Brassica rapa* has three genes that encode proteins associated with different neutral lipids in plastids of specific tissues. *Plant Physiology* **126**:330–341 DOI:0.2307/4279891.

Kim EH, Lee DW, Lee KR, Jung SJ, Jeon JS, Kim HU. 2017. Conserved function of fibrillin5 in the plastoquinone-9 biosynthetic pathway in *Arabidopsis* and rice. *Frontiers in Plant Science* **8**:1197 DOI: 10.3389/fpls.2017.01197.

Kuntz M, Chen HC, Simkin AJ, Römer S, Bramley PM. 1998. Upregulation of two ripening-related genes from a non-climacteric plant (pepper) in a transgenic climacteric plant (tomato). *Plant Journal* **13**(3):351-61 DOI:10.1046/j.1365-313X.1998.00032.x.

Laizet Y, Pontier D, Mache, Régis, Kuntz M. 2004. Subfamily organization and phylogenetic origin of genes encoding plastid lipid-associated proteins of the fibrillin type. *Journal of Genome Science & Technology* **3**(1):19-28 DOI:info:doi/10.1166/glt.2004.038.

Lee DG, Ahsan N, Lee S-H, Kang KY, Lee JJ, Lee B-H. 2007. An approach to identify cold-induced low-abundant proteins in rice leaf. *Comptes Rendus Biologies* **330**(3):0-225 DOI:10.1016/j.crvi.2007.01.001

Leitner-Dagan Y, Ovadis M, Shklarman E, Elad Y, Rav David D, Vainstein A. 2006. Expression and functional analyses of the plastid lipid-associated protein CHRC suggest its role in chromoplastogenesis and stress. *Plant Physiology* **142**(1):233-244 DOI: 10.2307/20205918.

Ling HQ, Zhao S, Liu D, Wang J, Sun H, Zhang C. 2013. Draft genome of the whe
at A-genome progenitor *Triticum urartu*. *Nature* **496**(2):87-90 DOI:10.1038/nature1199
7.

Lohscheider JN, Río Bártulos C. 2016. Plastoglobules in algae: a comprehensive compa
rative study of the presence of major structural and functional components in comple
x plastids. *Marine Genomics* **28**:127-136 DOI:10.1016/j.margen.2016.06.005.

Lundquist PK, Poliakov A, Bhuiyan NH, Zybailov B, Sun Q, van Wijk KJ. 2012. T
he functional network of the Arabidopsis plastoglobule proteome based on quantitativ
e proteomics and genome-wide coexpression analysis. *Plant Physiology* **158**: 1172– 1
192 DOI:10.1104/pp.111.193144.

Monte E, Ludevid D, Prat S. 1999. Leaf C40. 4: a carotenoid-associated protein involv
ed in the modulation of photosynthetic efficiency? *The Plant Journal* **19**:399–410 D
OI: 10.1046/j.1365-313x.1999.00537.x.

Newman LA, Hadjeb N, Price CA. 1989. Synthesis of two chromoplast-specific proteins
during fruit development in capsicum annum. *Plant Physiology* **91**(2):455-458 DOI:1
0.1104/pp.91.2.455.

Ning P, Liu C, Kang J. 2017. Genome-wide analysis of wrky transcription factors in w
heat (*Triticum aestivum* L.) and differential expression under water deficit condition.
Peerj **5**(5):e3232 DOI:10.7717/peerj.3232.

Paux E, Legeai F, Guilhot N, Adam-Blondon AF, Alaux M. 2008. Physical mapping i
n large genomes: accelerating anchoring of BAC contigs to genetic maps through in
silico analysis. *Functional & Integrative Genomics* **8**(1):29-32 DOI:10.1007/s10142-00
7-0068-1.

Philippa B, Ricardo RG, Cristobal U. 2016. expVIP: a customisable RNA-seq data ana
lysis and visualisation platform. *Plant Physiology* **170**(4): 2172-2186 DOI: 10.1104/pp.1
5.01667

- Pozueta-Romero J, Rafia F, Houlné G, Cheniclet C, Carde JP, Schantz ML, Schantz R. 1997.** A ubiquitous plant housekeeping gene, PAP, encodes a major protein component of bell pepper chromoplasts. *Plant Physiology* **115**: 1185–1194 DOI:10.1104/pp.115.3.1185.
- Pradhan S, Babar MA, Robbins K, Bai G, Mason RE, Khan J, Shahi D, Avci M, Guo J, Maksud HM, Bhatta M, Mergoum M, Asseng S, Amand PS, Gezan S, Baik BK, Blount A, Bernardo A. 2019.** Understanding the Genetic Basis of Spike Fertility to Improve Grain Number, Harvest Index, and Grain Yield in Wheat Under High Temperature Stress Environments. *Front Plant Science* **10**:1481 DOI:10.3389/fpls.2019.01481.
- Pruvot G, Cuiné S, Peltier G, Rey P. 1996.** Characterization of a novel drought-induced 34-kDa protein located in the thylakoids of *Solanum tuberosum* L. plants. *Planta* **198**(3): 471-479 DOI:10.1007/bf00620065.
- Rahimi Y, Bihamta MR, Taleei A, Alipour H, Ingvarsson PK. 2019.** Genome-wide association study of agronomic traits in bread wheat reveals novel putative alleles for future breeding programs. *BMC Plant Biology* **19**(1):541 DOI: 10.1186/s12870-019-2165-4.
- Rey P, Gillet B, Römer S, Eymery F, Massimino J, Peltier G, Kuntz M. 2000.** Over-expression of a pepper plastid lipid-associated protein in tobacco leads to changes in plastid ultrastructure and plant development upon stress. *The Plant Journal* **21**:483–494 DOI:10.1046/j.1365-313x.2000.00699.x.
- Ramírez-González RH, Borrill P, Lang D, Harrington SA, Brinton J, Venturini L, Davey M, Jacobs J, van Ex E, Pasha A, Khedikar Y, Robinson SJ, Cory AI, Florio I, Concia L, Juery C, Schoonbeek H, Steuernagel B, Xiang D, Ridout CJ, Chalhoub B, Mayer KFX, Benhamed M, Latrasse D, Bendahmane A;International Wheat Genome Sequencing Consortium, Wulff BBH, Appels R, Tiwari V, Datla R, Choulet E, Pozniak CJ, Provart NJ, Sharpe AG, Paux E, Spannagl M, Bräuti**

- qam A, Uauy C. 2018 The transcriptional landscape of polyploid wheat. *Science* **361**(6403): 6403-6089 DOI:10.1126/science.aar6089.
- Simkin AJ, Moreau H, Kuntz M, Pagny GI, Lin C, Tanksley S, James MC. 2008. An investigation of carotenoid biosynthesis in *Coffea canephora* and *Coffea arabica*. *J. Plant Physiology* **165**(10):1087-1106 DOI: 10.1016/j.jplph.2007.06.016.
- Singh DK, McNellis TW. 2011. Fibrillin protein function: the tip of the iceberg? *Trends in Plant Science* **16**(8):432-441 DOI:10.1016/j.tplants.2011.03.014.
- Singh DK, Maximova SN, Jensen PJ, Lehman BL, McNellis TW. 2010. FIBRILLIN4 is required for plastoglobule development and stress resistance in apple and Arabidopsis. *Plant Physiology* **154**(3):1281-1293 DOI:10.1104/pp.110.164095.
- Ting JTL, Wu SS, Ratnayake C, Huang AH. 1998. Constituents of the tapetosomes and elaioplasts in *Brassica campestris* tapetum and their degradation and retention during microsporogenesis. *The Plant Journal* **16**:541–551 DOI:10.1046/j.1365-313x.1998.00325.x.
- Vidi PA, Kanwischer M, Baginsky S, Austin JR, Csucs G, Dörmann P, Kessler F, Bérthelin C. 2006. Tocopherol cyclase (VTE1) localization and vitamin E accumulation in chloroplast plastoglobule lipoprotein particles. *Journal of Biological Chemistry* **281**:11225–11234 DOI:10.1074/jbc.M511939200.
- Willems, E., Leyns, L., and Vandesompele, J. 2008. Standardization of real-time PCR gene expression data from independent biological replicates. *Analytical Biochemistry* **379**(1), 127-129.
- Yang Y, Sulpice R, Himmelbach A, Meinhard M, Christmann A, Grill E. 2006. Fibrillin expression is regulated by abscisic acid response regulators and is involved in abscisic acid-mediated photoprotection. *Proceedings of the National Academy of Sciences of the United States of America* **103**(15): 6061-6066 DOI:www.pnas.org/cgi/doi/10.1073/pnas.0501720103.

Youssef A1, Laizet Y, Block MA, Maréchal E, Alcaraz JP, Larson TR, Pontier D, Gaffé J, Kuntz M.. 2010. Plant lipid-associated fibrillin proteins condition jasmonate production under photosynthetic stress. *The Plant Journal* **61**(3):436-445 DOI:10.1111/j.1365-313X.2009.04067.x.

Ytterberg AJ, Wijk PKJV. 2006. Protein profiling of plastoglobules in chloroplasts and chromoplasts a surprising site for differential accumulation of metabolic enzymes. *Plant Physiology* **140**(3):984-997 DOI:10.2307/20205663.

Figure 1

Phylogenetic relationship of the *TaFBN* genes, exon-intron *TaFBN* gene structure and functional domain analysis of TaFBN proteins.

(a) A phylogenetic tree inferred using the neighbor-joining method in MEGA7, with bootstrap values of 1000, was constructed to determine whether the exon-intron distribution patterns correlated with the phylogenetic classification of *TaFBN* (the same phylogenetic tree is also shown in Figs 2 and 4). **(b)** The coding sequences (CDS) of exons are indicated by yellow boxes, introns are represented by lines, and blue boxes indicate untranslated regions (UTRs). **(c)** Conserved domains of TaFBN proteins were identified using the Conserved Domain Database (CDD) of NCBI against the Pfam v30.0 database (<https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi>).

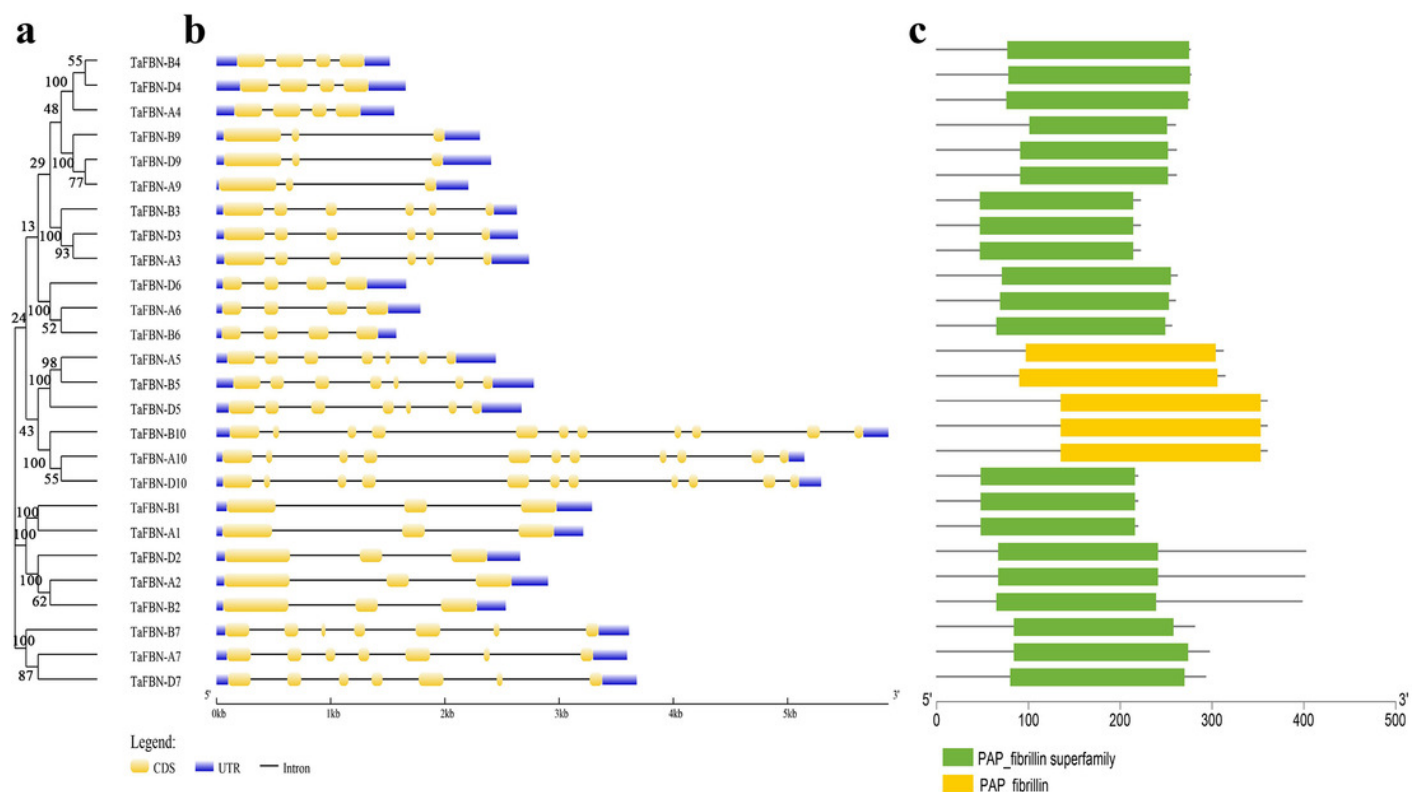


Figure 2

Motif distribution of proteins encoded by *Triticum aestivum* FBN genes.

(a) The phylogenetic tree of TaFBN genes was constructed using the neighbor-joining method in MEGA7, with bootstrap values of 1000. **(b)** Conserved Motifs were predicted using Multiple Em for Motif Elicitation (MEME) (<http://alternate.meme-suite.org/>). Box length indicates the number of amino acids in the motif.



Figure 3

Unrooted phylogenetic tree of all the *Triticum aestivum*, *Oryza sativa*, *Sorghum bicolor*, *Zea mays*, *Panicum hallii*, *Panicum virgatum*, *Setaria italica*, *Hordeum vulgare*, *Arabidopsis thaliana*, [i] *Brassica oler*

The phylogenetic tree was inferred using the neighbor-joining method in MEGA7, with bootstrap values of 1000. The FBN proteins are clustered into 11 subgroups, which are shown in different colors. Circles and stars indicate dicotyledons and monocotyledons plants, respectively. In addition, different colors represent different species.

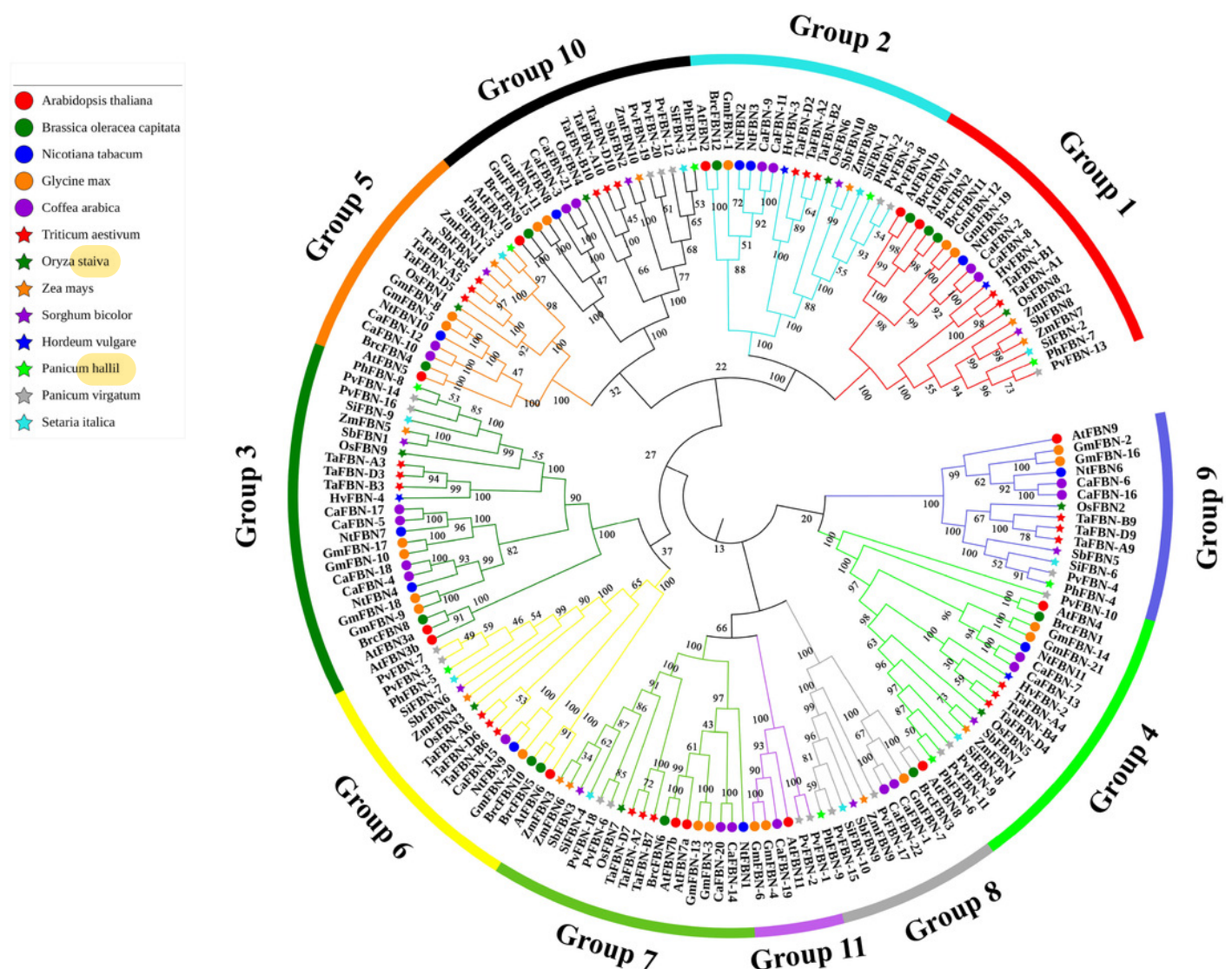


Figure 4

The analysis of *cis*-regulatory elements of *TaFBN* gene promoters.

(a) A phylogenetic tree inferred using the neighbor-joining method in MEGA7, with bootstrap values of 1000, was constructed to determine whether the exon-intron distribution patterns correlated with the phylogenetic classification of *TaFBN*. (b) Promoter sequences (2000 bp) upstream of genes were chosen for *cis*-regulatory element analysis using the PlantCARE online tool (<http://www.dna.affrc.go.jp/PLACE/>). Besides, each color indicates a *cis*-regulatory element.



Figure 5

Synteny and gene duplication in *TaFBN* genes.

A total of 26 *TaFBN* genes were unevenly located on 11 chromosomes. Seven segmentally duplicated gene pairs were identified in the wheat genome. Duplicated gene pairs are connected with green lines.

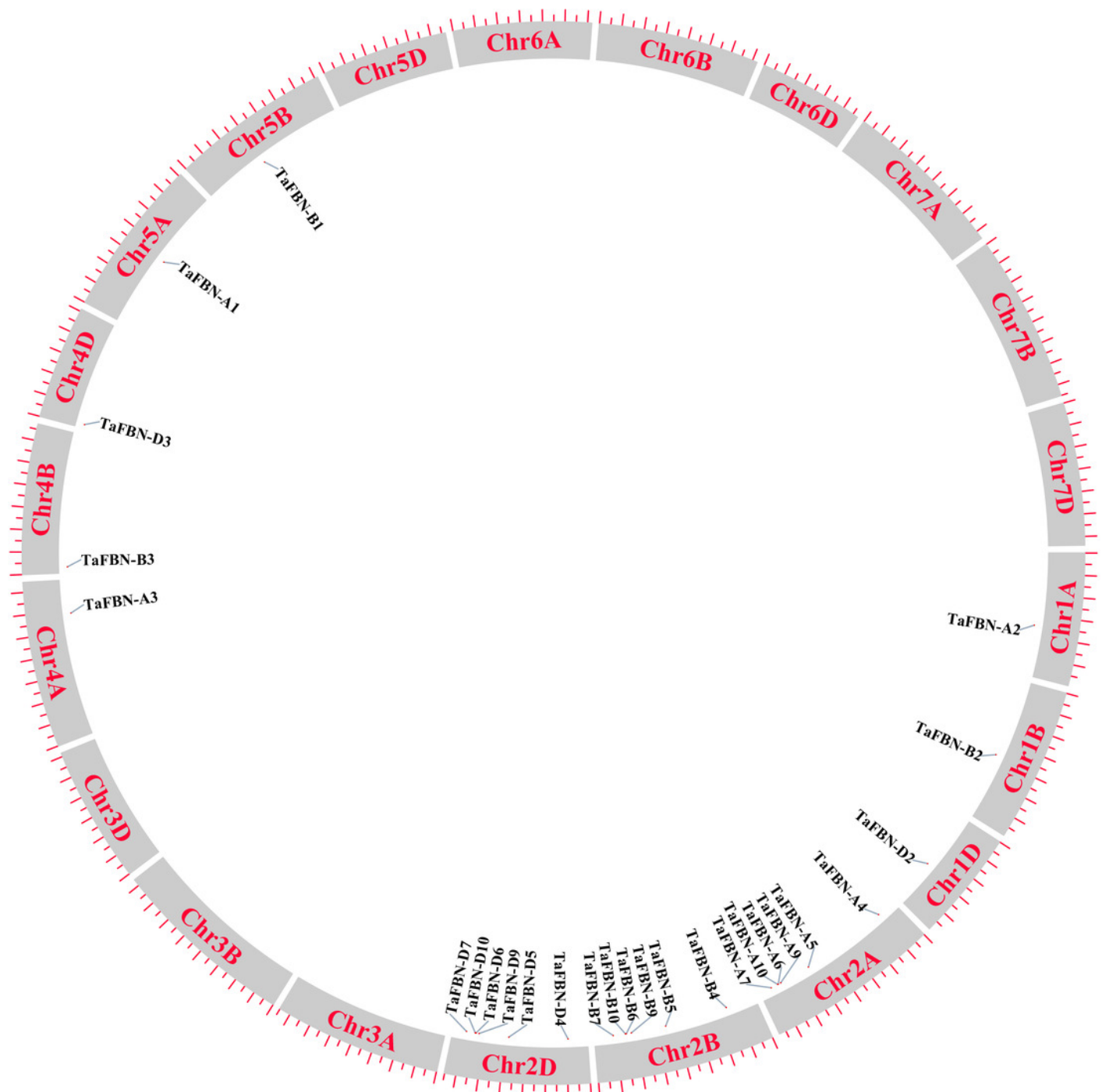


Table 1 (on next page)

Fibrillin (FBN) gene family in Triticum aestivum.

1 Table 1:

Gene name	Sequence ID	Chromosome	Genomic position	Intron number	Gene length (aa)	Molecular weight (kDa)	pI	Predicted pfam domain	Subcellular prediction by PC	Grand average of hydropathicity	Signal peptides	Chloroplast transit peptides
TaFBN-A1	TraesCS5A02G164600.1	Chr5A	353189098-353192310	2	314	33.06	7.77	PAP_fibrillin	Chloroplast	0.039	NA	Y
TaFBN-B1	TraesCS5B02G162100.1	Chr5B	299020240-299020330	2	312	32.94	7.77	PAP_fibrillin	Chloroplast	0.056	NA	Y
TaFBN-A2	TraesCS1A02G193500.1	Chr1A	350749390-350752293	2	360	38.27	4.79	PAP_fibrillin	Chloroplast	-0.261	NA	Y
TaFBN-B2	TraesCS1B02G208500.1	Chr1B	378397002-378399661	2	360	38.33	4.83	PAP_fibrillin	Chloroplast	-0.294	NA	Y
TaFBN-D2	TraesCS1D02G197400.1	Chr1D	278512124-278514657	2	360	38.28	4.79	PAP_fibrillin	Chloroplast	-0.253	NA	Y
TaFBN-A3	TraesCS4A02G272000.1	Chr4A	583754471-583757208	5	261	28.59	9.34	PAP_fibrillin	Extracellular	-0.33	NA	Y
TaFBN-B3	TraesCS4B02G042000.1	Chr4B	28717109-28719740	5	260	28.48	9.61	PAP_fibrillin	Chloroplast	-0.318	NA	Y
TaFBN-D3	TraesCS4D02G039200.1	Chr4D	16799419-16802059	5	261	28.55	9.21	PAP_fibrillin	Chloroplast	-0.325	NA	Y
TaFBN-A4	TraesCS2A02G145900.1	Chr2A	90688741-90690297	3	275	28.99	8.95	PAP_fibrillin	Chloroplast	-0.244	NA	Y
TaFBN-B4	TraesCS2B02G171300.1	Chr2B	144596063-144597581	3	276	29.35	9.51	PAP_fibrillin	Chloroplast	-0.267	NA	Y
TaFBN-D4	TraesCS2D02G150500.1	Chr2D	93046450-93048107	3	277	29.41	9.51	PAP_fibrillin	Chloroplast	-0.277	NA	Y
TaFBN-A5	TraesCS2A02G300200.1	Chr2A	515959001-515961447	6	262	28.96	9.16	PAP_fibrillin	Chloroplast	-0.213	NA	Y
TaFBN-B5	TraesCS2B02G316500.1	Chr2B	451833336-451836114	6	260	28.67	9.28	PAP_fibrillin	Chloroplast	-0.178	NA	Y
TaFBN-D5	TraesCS2D02G298100.1	Chr2D	380429694-380432365	6	256	28.43	9.36	PAP_fibrillin	Chloroplast	-0.2	NA	Y
TaFBN-A6	TraesCS2A02G431000.1	Chr2A	684246511-684248296	3	219	23.78	8.8	PAP_fibrillin	Extracellular	-0.044	NA	Y
TaFBN-B6	TraesCS2B02G452300.1	Chr2B	646214215-646215789	3	219	23.75	8.73	PAP_fibrillin	Chloroplast	0.003	NA	Y
TaFBN-D6	TraesCS2D02G428800.1	Chr2D	540824383-540826044	3	219	23.82	8.74	PAP_fibrillin	Chloroplast	-0.031	NA	Y
TaFBN-A7	TraesCS2A02G487900.1	Chr2A	722519297-722522892	6	297	32.55	5.73	PAP_fibrillin	Chloroplast	-0.231	NA	Y
TaFBN-B7	TraesCS2B02G515500.1	Chr2B	710281451-710285064	6	281	30.92	6.06	PAP_fibrillin	Chloroplast	-0.247	NA	Y

TaFBN-D7	TraesCS2D0 2G488200.1	Chr2D	587697352- 587701032	6	293	32.03	5.35	PAP_fibrilli n	Chloroplast	-0.171	NA	Y
TaFBN-A9	TraesCS2A0 2G413700.1	Chr2A	670791911- 670794116	2	222	24.21	6.74	PAP_fibrilli n	Extracellular	-0.106	NA	Y
TaFBN-B9	TraesCS2B0 2G432500.1	Chr2B	621664679- 621666987	2	222	24.27	7.9	PAP_fibrilli n	Extracellular	-0.123	NA	Y
TaFBN-D9	TraesCS2D0 2G410900.1	Chr2D	525935293- 525937697	2	222	24.24	7.9	PAP_fibrilli n	Extracellular	-0.136	NA	Y
TaFBN-A10	TraesCS2A0 2G434800.1	Chr2A	686874975- 686880122	10	401	43.44	9.31	PAP_fibrilli n	Extracellular	-0.16	NA	Y
TaFBN-B10	TraesCS2B0 2G455900.1	Chr2B	650399573- 650405456	10	402	43.59	9.19	PAP_fibrilli n	Extracellular	-0.182	NA	Y
TaFBN-D10	TraesCS2D0 2G432600.1	Chr2D	544665056- 544670351	10	398	43.12	9.11	PAP_fibrilli n	Extracellular	-0.152	NA	Y

Figure 6

Expression of *TaFBNs* in various tissues and developmental stages.

(a) Tissue-specific Expression of the *TaFBN* gene family in different wheat tissues; (b) Expression pattern of the *TaFBN* gene family at different developmental stages; (c) Heatmap was created in TBtools software and based on the expression data, The color scale represents relative expression levels, with red indicating higher levels of expression blue indicating lower expression levels.

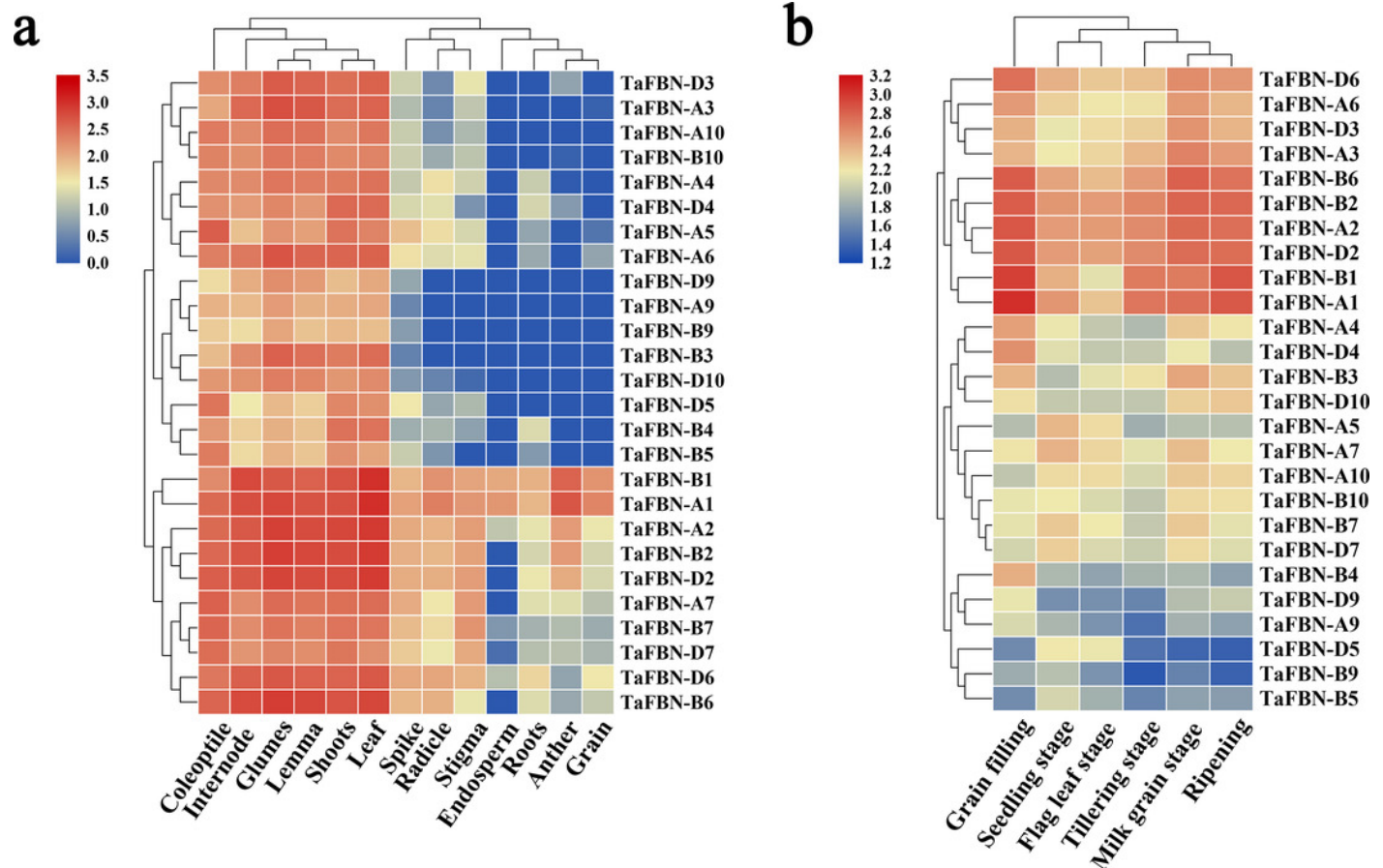


Figure 7

Heat map of expression profiles of *TaFBNs* in *Triticum aestivum* under biotic and abiotic stress.

Expression levels are indicated in different colors, with red indicating higher expression levels and blue indicating lower expression levels.

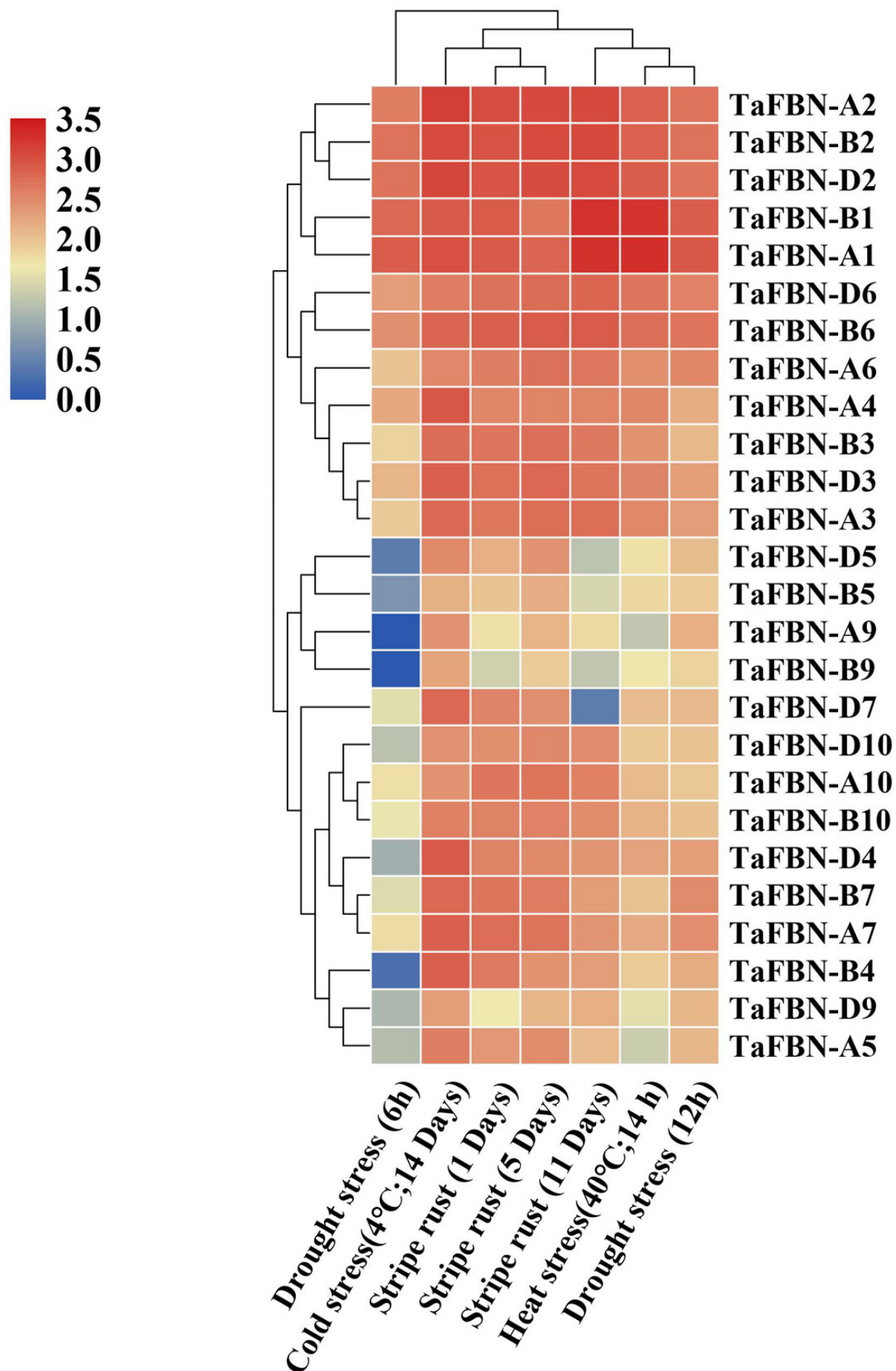


Figure 8

Expression analysis of TaFBN genes in different tissues and under drought stress using qRT-PCR.

(a) the relative expression levels of TaFBN genes in different tissues. (b) the relative expression levels of TaFBN genes in leaves after drought treatment for 1h and 6h. each treatment contains three biological replicates .

