Genome-wide identification and characterization of OVATE family proteins in Betula luminifera reveals involvement of BIOFP3 and BIOFP5 genes in leaf development

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Ovate family proteins (OFP) are plant-specific transcription factors involved in regulating morphologies of the lateral organs, plant growth and development. However, the functional roles of OFP genes in Betula luminifera, an important timber tree species, are not well studied. In this study, we identified 20 BIOFP genes and analyzed their phylogenetic relationship, gene structure, conserved motifs, and cis-elements. Further, expression analysis indicates that BIOFP genes were up-regulated in leaves on the one-year-old branch compared to leaves on the current-year branch and bract, except BIOFP7, BIOFP11, BIOFP14 and BIOFP12. The overexpression of BIOFP3 and BIOFP5 in Arabidopsis thaliana not only resulted in a slower growth rate but also produced sawtooth shape, flatter and darker green rosette leaves. Further investigation showed that the leaf thickness of the transgenic plants was more than double that of the wild type, which was caused by the increasement in the number and size of palisade tissue cells. Furthermore, the expression analysis also indicated that the expressions of several genes related to leaf development were significantly changed in the transgene plants. These results suggested the significant roles of BIOFP3 and BIOFP5 in leaf development. Moreover, protein-protein interaction studies showed that BIOFP3 interacts with BKNAT5, and BIOFP5 interacts with BKNAT5, BIBLH6 and BIBLH7. In conclusion, our study demonstrates that BIOFP3 and BIOFP5 were involved in leaf shape and thickness regulation by forming a complex with BKNAT5, BIBLH6 and BIBLH7. In addition, our study serves as a guide for future functional genomic studies of OFP genes of the B. luminifera.

KEYWORDS
Betula luminifera, ovate family protein, expression analysis, protein-protein interaction, transmission electron microscopy
Introduction

The OVATE Family Proteins (OFP) are unique plant-specific transcription factors containing a ~70 amino acids long conserved C-terminal OVATE domain (DUF623) (Schmitz et al., 2015). Based on the similarity of the OVATE domain, OFPs were identified in monocots, dicots, bryophytes and lycophytes (Liu et al., 2014). The first OFP gene was cloned by Liu et al. (2002) from tomatoes, and they reported that a point mutation in the OFP gene resulted in pear-shaped tomatoes, and overexpression causes abnormal phenotypes like slower plant growth, exserted stigmas, and altered floral and vegetative architecture. Later, the function of OFP genes was studied in many plant species and identified that OFP genes exhibit various roles in the different stages of a plant’s growth and development. For example, in Arabidopsis thaliana, an overexpression study of AtOFP1 exhibits round and curly leaves and kidney-shaped cotyledons. Moreover, the expression of AtOFP1 also inhibits the expression of AtGA20ox1, an essential gene for gibberellin biosynthesis, and prevents cell elongation (Wang et al., 2007). Besides, Wang et al. (2011) reported that in A. thaliana, AtOFPs act as transcriptional repressors regulating the morphology of cotyledons and true leaves, flower formation, and seed development. Further, overexpression of OsOFP2 in rice leads to a decrease in plant height and changes in leaf morphology and seed shape (Schmitz et al., 2015). Likewise, genetic transformation experiments revealed that rice OsOFP19 could affect architecture and seed shape by regulating the BR signalling pathway (Yang et al., 2018).

OFP proteins form complexes with other proteins and directly or indirectly affect the transcription of target genes to regulate plant growth and development. For example, A. thaliana OFPs can interact with TALE (three-amino-acid-loop-extension) superfamily members to form a complex and play an essential regulatory role in growth and development (Hackbusch et al., 2005). Further, AtOFP1 and AtBLH3 form a protein complex that regulates the transition time from vegetative growth to reproductive development of A. thaliana (Zhang et al., 2016). Moreover, AtOFP4 interacts with AtKNAT7 protein and acts as a transcriptional inhibitor to regulate the formation of the secondary cell wall of the inflorescence stem in A. thaliana (Li et al., 2011; Li et al., 2012). In addition, AtOFP5 interacts with AtBLH1 and AtKNAT3 and coordinate regulates embryo sac development (Pagnussat et al., 2007). Besides this, AtOFP1 also interacts with DNA double-strand break repair-related protein AtKu and jointly regulates the DNA double-strand break repair process (Wang et al., 2010). Further, rice OsOFP8 phosphorylates OsGSK2 protein and plays a role in growth and development by participating in the BR signalling pathway (Yang et al., 2016). Likewise, OsOFP19 forms a complex with OSH1 (a member of the KNOX subfamily protein) and DLT (DWARF AND LOW-TILLERING) protein. The complex of these three proteins plays a vital role in regulating BR signalling and determining cell division patterns (Yang et al., 2018).

Materials and methods

Plant materials

The material used in this study is B. luminifera 1-V-25-2 clone, planted in the Pingshan nursery of Zhejiang A & F University. The leaves of three different developmental stages were collected from a healthy plant and named Bt (bracts), YL1 (young leaf from the current-year branch), and YL2 (young leaf from the one-year-old branch). The sample was collected in April 2021. The A. thaliana Columbia-0 (Col-0) was used for gene transformation. Seeds of Col-0 were sterilized twice with 75% alcohol and grown on ½ MS media until leaves reached the four-leaf stage in an artificial incubator (at 25°C, light for 16 h, dark for 8 h, with an average light intensity of 12,000 lx). Then the seedlings were transplanted to the peat moss (peat soil: vermiculite: perlite is 3:1:1) and grew at 22°C under photoperiod (16h light/8h dark) in a growth chamber.

Identification of OFP genes

The Hidden Markov Model (HMM) profile of the OVATE domain (PF04844) was obtained from the Pfam database (Mistry et al., 2021). Putative OFP protein sequences were identified by
HMMER search against the B. luminifera genome and the publicly available B. pendula genome (Salojarvi et al., 2017). In cases of multiple putative proteins from the same gene locus, the most extended variant was kept for further analysis. Editsq module in the DNASTar package was used to predict the amino acid sequence, molecular weight, and isoelectric point of the deduced protein. The identified gene sequences are provided in Supplementary Table 1.

Gene structure, genome location and cis-elements analysis of BIOFPs

The gene structure and genomic location of BIOFPs were searched in the genomic database, and the localization of the BIOFP genes on the chromosomes was generated using TBtools (Chen et al., 2020). The PlantCARE webtool (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/) was used to analyze the cis-acting elements in the promoter sequence.

Phylogenetic analysis and conserved motif analysis of BIOFP proteins

The phylogenetic tree was constructed using MEGA-X using the Maximum likelihood method (Kumar et al., 2018). The aligned peptide sequences of OVATE domains from B. luminifera, B. pendula, A. thaliana, Oryza sativa and P. trichocarpa OFP proteins were used to generate phylogenetic analysis. The bootstrap value of 1000 replicates was used to evaluate the statistical significance. The conserved motifs were identified by the online program MEME (http://meme-suite.org/tools/meme) and visualized using the TBTools (Chen et al., 2020).

RNA isolation

According to the manufacturer’s instructions, the total RNA was extracted from tissues using PureLink™ Plant RNA Reagent (Invitrogen, Carlsbad, CA, USA). The concentration of RNA samples was analyzed by spectrophotometer (Nanodrop ND-2000), and purity and integrity were analyzed by agarose gel electrophoresis. Promega’s RQ1 RNase-Free DNase was used to remove the DNA contamination following the manufacturer’s instructions.

Real-time qRT-PCR

The cDNA was synthesized using Takara’s PrimeScript™ RT reagent Kit (Perfect Real Time) as per the manufacturer’s instructions. Oligo7.0 software was used to design fluorescent quantitative PCR primers (Supplementary Table 2). The TB green premix ex Taq II (Tli RNase H Plus) (Takara) was used for qRT-PCR amplification in the CFX96™ Real-Time PCR detection system (Bio-Rad, USA). All PCR reactions were performed in triplicate. The BIEFla gene was used as an internal reference for B. luminifera samples, and the Actin gene was used as an internal reference in qRT-PCR for Arabidopsis samples. All the primers used in qRT-PCR were listed in Supplementary Table 2. The relative expression levels were determined with CFX96 Manager™ software (version 1.6, Bio-Rad, USA) using the delta-delta Ct method (Livak and Schmittgen, 2001). GraphPad Prism (Version 5.01) was used to draw the graphs of the quantitative results.

Production of transgenic A. thaliana plants

For the construction of the overexpression vector, the open reading frame (ORF) of BIOFP3 and BIOFP5 genes were inserted into the pCAMBIA13011 vector, respectively. The generated vectors pCAMBIA13011-BIOFP3 and pCAMBIA13011-BIOFP5 were transferred to Agrobacterium tumefaciens (GV3101). Transformation of transgenic A. thaliana plants was performed using the floral dip method (Clough and Bent, 1998), and transgenic plant selection was performed on MS agar medium supplemented with hygromycin. The transgenic plants were transplanted to peat moss and grown as mentioned above for wild-type (WT) plants. The transgenic A. thaliana plants were also verified by PCR and qRT-PCR, and homozygous T3 generation plants were used for further analysis.

Phenotyping of transgenic plants

To observe the growth and leaf development of transgenic plants, the young (21-day-old) and mature (42-day-old) transgenic plants were photographed and compared with WT. The relative chlorophyll content of leaves was measured using a SPAD (soil plant analysis development) meter. To measure the leaf thickness and cell number, the cross-section of about 8 mm thick from rosette leaves of 42-day-old plants was obtained by microtome (Leica). The sections were stained with 0.02% toluidine blue for 5 minutes and rinsed with distilled water until the wash solution was clear. Finally, the sections were observed under a light microscope (Zeiss). At least 8 sections (8 replicates) of each sample were used to determine the leaf thickness and cell number.

Transmission electron microscopy

To measure cell size, the rosette leaves of a 42-day-old plant were fixed in 0.1 M phosphate buffer (pH 7.2) with 2.5%
glutaraldehyde at 4°C overnight. Then, the samples were washed with 0.1 M phosphate buffer and treated with 1.0% osmium tetroxide for 1h. After that, the samples were rinsed twice in deionized water and dehydrated in a graded ethanol series. After dehydration, the samples were infiltrated and embedded in epon-araldite. Ultra-thin sections (70 nm) were sliced with a diamond knife (Reichert-Jung, Germany), stained with uranyl acetate and lead citrate, and mounted on nickel grids for examination in the transmission electron microscope (FEI Titan G2 80-200 ChemiSTEM, USA). At least 10 photos for each sample were analyzed.

### Yeast two-hybrid assay

Y2H experiment was performed according to the manufacturer’s instructions of Matchmaker™ gold yeast two-hybrid system (Clontech). Briefly, the ORFs of BIOFPs were cloned into bait vector pGBK7, respectively. The ORFs of BIKNOXs and BIBLHs genes were constructed into prey vector pGADT7, respectively. The recombinant bait and prey vectors were transformed into the Y2H gold strain and Y187 vector pGADT7, respectively. The recombinant bait and prey in BiFC were listed in Supplementary Table 2. To observe and detect protein interaction signals, the primers used in Y2H were also listed in Supplementary Table 2.

### Bimolecular fluorescence complementation assay

The ORF of BIOFP3, BIOFP5, BIKNOX5, BIKNOX9, BIBLH6, BIKNOX5, BIKNOX9 and BIBLH7 were cloned into the pSAT1- nEYFP-C1 and pSAT1- cEYFP-C1-B vectors using ClonExpress® II One Step Cloning Kit (Vazyme). The recombinant vectors transiently co-expressed in Arabidopsis protoplast cells through PEG-mediated transformation (Yoo et al., 2007). The images of transformed protoplast cells were captured on a Laser confocal microscope (Carl Zeiss LSM880) to observe and detect protein interaction signals. The primers used in BiFC were listed in Supplementary Table 2.

### Statistical analysis

The significant differences in experimental data were compared by the Duncan algorithm. All calculations are based on Data Processing System Software (V15.10), and differences were considered significant at a p-value ≤ 0.05 level.

### Results

#### Identification of OFP genes in B. luminifera and B. pendula

Our lab has developed B. luminifera genome reads using PacBio’s RS II platform, and the assembled genome contained 9,332 contigs with an N50 of 247 kb (unpublished data). Using this contigs information, we have identified a total of 20 B. luminifera BIOFPs (BIOFP1-BIOFP20) genes using the HMM model. Further, we also identified the 20 BpOFPs of silver birch (B. pendula) using the published genome (Salojärvi et al., 2017). The 20 BIOFP proteins consist of 106-422 amino acid residues, with a molecular weight of 12.08-48.21 kDa and an isoelectric point of 4.38-9.79 (Table 1). Further, multiple sequence alignment indicated that all the 20 BIOFP proteins contain an OVATE domain, which was not highly conserved among these proteins (Supplementary Figure 1). The gene structure of BIOFP genes is also analyzed and identified that the ORF size in the genomic DNA ranged from 321 - 1269 bp. Out of 20 BIOFP genes, 18 BIOFPs genes are intron-free. While BIOFP19 and BIOFP20 have an intron of 2506 bp and 110 bp in length, respectively (Figure 1). Based on the published genome database of B. pendula (Salojärvi et al., 2017), we analyzed the distribution of BpOFPs genes on the chromosomes (Figure 2). The 16 BpOFP genes are mapped to 9 chromosomes of B. pendula, of which chromosomes 6 and 8 have the highest number of genes, i.e. three OFP. In addition, BpOFP5, BpOFP11, BpOFP15, and BpOFP17 were found to be located on unanchored contig796, contig419 and contig230 (Figure 2).

#### Phylogenetic analysis of birch OFP proteins

In order to analyze the phylogenetic relationship of OFP proteins, a phylogenetic tree containing 102 OFP proteins (BIOFPs, BpOFPs, AtOFPs, OsOFPs and PtOFPs) is constructed. All OFP proteins can be divided into eight subgroups, except Subgroup II and VIII. The 20 BIOFP proteins are placed into the same subgroup, Subgroup-I. Along with BIOFP3 and BIOFP5, BpOFPs were placed in the same subgroup, Subgroup-I. All OFP proteins are placed next to each other in all clades, indicating that these OFP proteins have a certain degree of differentiation during evolution. The two concerned BIOFPs of the present study, i.e., BIOFP3 and BIOFP5, were placed in the same subgroup, Subgroup-I. Along with BIOFP3 and BIOFP5, AtOFP6 and AtOFP19 from A. thaliana and OsOFP8 and OsOFP20 from O. sativa are grouped in the same subgroup, indicating their close homologous relationships (Liu et al., 2014). Further, most monocot OFP proteins (OsOFPs) are divided into six subgroups, except Subgroup II and VIII.
Prediction of conserved motifs in birch OFP proteins

Studying the conserved motifs can reveal the functional diversity of OFP proteins. Through conserved motif analysis, fifteen conserved motifs of BIOFPs were identified. Among these motifs, OVATE domains are present in the C-terminal of OFP proteins, while other motifs are upstream of the OVATE domain (Figure 3). Interestingly, the OsOFP23 in Subgroup III, OsOFP11 in Subgroup VI and BIOFP2, BpOFP2, PtOFP28,
AtOFP17 and AtOFP20 in the Subgroup VIII don’t contain the complete OVATE domain (Figure 3). These results suggest that the OVATE domain is not that conserved during evolution. Further, most OFP proteins in the same subgroup have similar motif distributions (Figure 3). These results show that the motifs of members within the same subgroup are conserved, which implies they may perform similar functions.

Analysis of cis-acting elements in the promoter region of BIOFP genes

The cis-acting elements play a vital role in regulating gene transcription during plant growth, development, and stress. Therefore, studying the cis-elements in BIOFP genes will help understand their role in plant growth and development. Hence, we analyzed the 1500 bp upstream region of the 20 BIOFPs genes. We classified the cis-acting elements into eight types based on the functions of cis-elements: light-responsive elements, induction-specific elements, plant tissue-specific elements, binding site-specific elements, transcription-related elements, process-specific elements, regulatory-specific elements, and environment-specific elements (Figure 4A). The promoter regions of BIOFPs genes contain at least six different types of cis-acting elements. Although the regulatory-specific cis-elements are the most abundant among the different cis-elements analyzed, the process-specific elements (plant hormone-related cis-acting elements) seem to be essential. Further, ABRE, ERE, GARE motif, TCA-element, TC-rich repeats, and TGA elements were found in abundance compared to other process-specific elements (Figure 4B).

Expression of BIOFP genes during leaf development in B. luminifera

The OFP genes are reported to be involved in the regulation of leaf development (Yang et al., 2016; Sun et al., 2020). So to identify the expression pattern of BIOFP genes in leaf development, we performed qRT-PCR in B. luminifera leaves with three developmental stages: Brt, YL1, and YL2. Most BIOFP genes have increased expression in YL2 except BIOFP7, BIOFP11, BIOFP14, and BIOFP12 (Figure 5). The BIOFP7 and BIOFP12 showed their highest relative expressions in Brt, and the BIOFP11 and BIOFP14 showed their highest expressions in YL1 (Figure 5). Interestingly, our qRT-PCR analysis showed that the BIOFP9 and BIOFP20 gene expression in YL2 was 228 and 184 times to the Brt, and 63 and 162 times to YL1, respectively. Further, BIOFP3 and BIOFP5 in YL2 were 6.9 and 4.4 times to the Brt and 9.6 and 6.2 times to the YL1, respectively (Figure 5).

Phenotype and anatomic analysis of Arabidopsis overexpressing BIOFP3 and BIOFP5

In our study, through phylogenetic and motif analysis, we identified that AtOFP6 and OsOFP8 are closely related to BIOFP3 and BIOFP5. Furthermore, sequence alignment of AtOFP6, BIOFP3 and BIOFP5 showed that conserved OVATE domain among these sequences (Supplementary Figure 2). Considering the function of AtOFP6 in leaf development (Wang et al., 2011), the role of BIOFP3 and BIOFP5 genes was first investigated through ectopic expression in A. thaliana plants. The transgenic plants were screened out and confirmed
using PCR detection on two reporter genes and BlOFP3/5 (Supplementary Figure 3), and then homologous T3 generations were used for phenotypic observation and anatomical analysis. The expression of BlOFP3 and BlOFP5 genes are also investigated by qRT-PCR in T3 generation transgenic plants. The results showed that BlOFP3 and BlOFP5 were strongly expressed in the transgenic plants (Supplementary Figure 3). Phenotype observation showed that overexpressing of BlOFP3 and BlOFP5 caused apparent differences in growth rate and leaf shape (Figure 6). The plant height of 42-day-old transgenic plants was 13.04 cm (35S::BlOFP3) and 13.08 cm (35S::BlOFP5), respectively, while the plant height of WT was 15.60 cm, which indicated that the growth of transgenic plants was slower (Figure 6E). Moreover, the rosette leaves of the two transgenic lines were flatter, darker green in colour, and a more obvious sawtooth appeared near the leaf stalk (Figures 6A, B). Measurement of the leaf area of 21-day-old and 42-day-old plants showed that the plants overexpressing BlOFP3 and BlOFP5 have considerably less leaf area (Figure 6C).

Furthermore, the SPAD values indicated that the relative chlorophyll content of leaves in transgenic plants is significantly higher than that in WT (Figure 6D). And the anatomic analysis

![Phylogenetic relationship and conserved motif analysis of BIOFPs from B. luminifera. The phylogenetic tree was constructed using aligned OVATE domains sequences of B. luminifera (Bl), B. pendula (Bp), A. thaliana (At), Oryza sativa (Os) and P. trichocarpa (Pt). All the eight subgroups are indicated in different colours. The conserved motifs (1-15) are indicated in 15 different colour boxes.](image)
revealed that the leaf thickness of plants overexpressing *BlOFP3* and *BlOFP5* is more than two times of the WT (Figures 7A, C). Meanwhile, an obvious increase in the number and size of palisade tissue cells is also observed in transgenic plants (Figures 7A, D, E). The increase in the size of palisade tissue cells was confirmed using transmission electron microscopy (TEM) (Figure 7B). Besides, the expression of *AtOFP6*, the endogenous homolog for *BlOFP3* and *BlOFP5*, was not altered in the plants overexpressing *BlOFP3* and *BlOFP5* (Supplementary Figure 4). These results suggest that *BlOFP3* and *BlOFP5* may play essential roles in leaf development by regulating the cell’s number and size in palisade tissue.

Expression analysis of genes associated with leaf development in transgenic *Arabidopsis*

To explore the molecular mechanism of *BlOFP3* and *BlOFP5* in leaf development, six genes, including KNAT2, KNAT6, AS1, CLV1, GA20ox1, PIN1 and CUC2 genes, have been proved as crucial genes in leaf development (Long et al., 1996; Belles-Boix et al., 2006; Wang et al., 2021; Zhou et al., 2022), were chosen for expression analysis in the transgenic *A. thaliana* plants. As shown in figure 8, the expressions of KNAT2 and KNAT6 were both drastically inhibited in plants overexpressing *BlOFP3* and *BlOFP5* (Figure 8). At the same time, the expression of CLV1, GA20ox1, PIN1 and CUC2 genes up-regulated in the transgenic plants (Figure 8). These results further support the essential roles of *BlOFP3* and *BlOFP5* in leaf development. It also implies that these two genes may perform redundant functions through similar pathways in the leaf development of *B. luminifera*.

Interaction analysis of *BlOFP3* and *BlOFP5* with TALE homeodomain protein

In previous studies, OFP proteins have been shown to interact with various TALE (KNOX and BLH) proteins, forming a TALE-OFP protein complex to regulate different biological processes (Hackbusch et al., 2005; Li et al., 2011; Liu and Douglas, 2015). In this study, the interactions between these two BIOFP proteins (*BlOFP3* and *BlOFP5*), eight BIKNOX proteins and nine BIBLH proteins were first analyzed using the Y2H method, respectively. As shown in figure 9, strong interactions between BIOFP3 and BIKNOX5, BIOFP3 and BIBLH6, BIOFP5 and BIKNOX5 as well as BIOFP5 and BIBLH6 were detected, and weak interactions between BIOFP3 and BIKNOX9, and BIOFP5 and BIBLH7 were also identified. However, in the present experiment, all other members of TALE
FIGURE 5
Expression analysis of BIOFP genes for leaf development. qRT-PCR analysis of 20 BIOFP genes in three developmental stages of B. luminifera leaves. The error bar represents the standard deviation from three biological replicates. The asterisk indicates significant difference compared to Brt at p<0.05 level.

FIGURE 6
Phenotype observation on transgenic plants overexpressing BIOFP3 and BIOFP5 in A. thaliana. (A) 21-day-old transgenic plants transformed with 35S::BIOFP3 (left), 35S::BIOFP5 (middle), and WT plants (right). (B) 42-day-old transgenic plants transformed with 35S::BIOFP3 (left), 35S::BIOFP5 (middle), and WT plants (right). (C) Leaf area of 21-day and 42-day-old plants. (D) SPAD values of plants. (E) Plant height of 42-day-old plants. Bar = 1cm. The asterisk indicates significant difference compared to WT at p<0.05 level.
homeodomain proteins did not show any interactions with BIOFP3 and BIOFP5 (Figure 9). This means there is interaction specificity exists between OFP and TALE homeodomain proteins.

Furthermore, the interactions obtained from Y2H were then verified by the BiFC assay in mesophyll protoplast cells of A. thaliana. Similar to the Y2H assay, the protoplast cells co-expressing BIOFP3-cEYFP with BIKNOX5-nEYFP, BIOFP5-cEYFP with BIKNOX5-nEYFP, as well as BIOFP5-cEYFP with BIBLH6-nEYFP exhibited intense fluorescence signals (Figure 10). In addition, a weak fluorescence signal was detected in the protoplast cells co-expressing BIOFP5-cEYFP with BIBLH7-nEYFP (Figure 10). These results were consistent with the Y2H interactions. However, in contrast to Y2H results, the fluorescence signal was absent in protoplast cells co-expressing BIOFP3-cEYFP with BIKNOX9-nEYFP and
BIOFP3-cEYFP with BBLH6-nEYFP (Figure 10). Based on the above results, we hypothesized that at least BKNOX5 is a common partner for BIOFP3 and BIOFP5, forming a functional complex to regulate leaf development in B. luminifera.

Discussion

OFPs, encoding a class of plant-specific transcription regulators, have been identified in many plants (Wang et al., 2010; Rodriguez et al., 2011; Tsaballa et al., 2011; Wang et al., 2011; Wang et al., 2016; Yang et al., 2016; Yang et al., 2018). This study identified and characterized 20 genes encoding OFP proteins in B. luminifera. As indicated by phylogenetic analysis, 20 OFP proteins in B. luminifera are distributed into eight subgroups based on their sequence similarity (Figure 3). The two studied BIOFPs of the present study, i.e., BIOFP3 and BIOFP5, showed close homologous relationships with AtOFP6 and AtOFP19 from A. thaliana. Domains and motifs are the key elements for the structure and functions of a protein or protein family (Moore et al., 2008; Forslund and Sonnhammer, 2012). In this study, 15 conserved motifs were identified, and most of these motifs showed similar distribution patterns to their homologs within the same subgroup (Figure 3). Moreover, the OVATE domain was also identified to be present at the end of 20 BIOFPs (Figure 3), which is consistent with the previous studies (Liu et al., 2014; Li et al., 2019). Since BIOFPs and BpOFPs share almost identical OVATE domain sequences, all BIOFPs and BpOFPs are positioned next to each other except BIOFP18 and BpOFP18. Divergent expansion might be considered the reason behind the presence of BIOFP18 and BpOFP18 in different subgroups (Liu et al., 2014). In the current study, hormone-specific cis-acting elements, including ABRE, TCA-element, TC-rich repeats, and TGA elements, which are associated with abscisic acid (ABA), salicylic acid (SA), gibberellin (GA) and auxin (IAA) responses, respectively, were found to be distributed in the promoter region of the BIOFPs genes (Figure 4). These hormone-responsive regulatory elements in the promoter regions suggest that BIOFP genes might be involved in response to various plant hormones and developmental processes (Liu and Douglas, 2015; Tang et al., 2018; Wu et al., 2018).
OFPs play diverse roles at different stages of plant growth and development (Wang et al., 2016; Knaap and Ostergaard, 2018). Regulation of leaf development is one of those important functions for plant OFP genes. For example, Wang et al. (2011) reported that overexpression of the AtOFP6 gene resulted in flat, thick and cyan rosette leaves in A. thaliana. In the current study, transgenic A. thaliana overexpressing BlOFP3 and BlOFP5 genes showed flat, thick leaves with higher relative chlorophyll content (Figure 6). Likewise, Zhou et al. (2019) found that tomato plants overexpressing SlOFP20 genes showed shorter compound leaves and contained more chlorophyll than wild plants. These observations indicate that these OFP genes, including BlOFP3 and BlOFP5, are essential regulators for leaf development. Moreover, in transgenic Arabidopsis plants overexpressing BIOFP3 and BIOFP5, the expressions of several essential leaf development genes (KNAT2, KNAT6, AS1, CLV1, GA20ox1, PIN1, and CUC2) were considerably changed (Figure 8). As a critical TF in plant growth and development, the class I KNOX genes (KNAT2, KNAT6) maintain the function of shoot apical meristem (SAM) by regulating other key genes involved in leaf development. Prior studies have reported that AS1 and AS2 form a protein complex that inhibits the expression of KNAT2, resulting in regular leaf morphology (Li et al., 2016; Li et al., 2017). In the current study, the expression analysis of KNAT2 and KNAT6 in transgenic plants has shown an antagonistic relationship with AS1 expression (Figure 8). There is an antagonism that exists between auxin and KNOX too. At the same time, auxins are transported by PIN1. The interaction among AS1, auxin and KNOX1 may directly affect the process of leaf initiation as well as leaf morphology (Hay et al., 2006). Likewise, the GA20ox1 is involved in forming the boundary between SAM and the primary leaf primordia, which results from inhibition of the synthesis of GA by KNOX (Rosin et al., 2003; Hay and Tsiantis, 2009; Nakayama et al., 2014). Our study identified that the GA20ox1 expression is also significantly changed in transgenic lines. At the same time, CLV1 promotes stem cell activity in the SAM. Similarly, CUC2 genes are also involved in leaf primordia initiation and affect the leaves’ morphology (Hasson et al., 2011). In the current study, CUC2 expression is also found to be up-regulated in the transgenic lines (Figure 8). These results further demonstrated that BIOFP3 and BIOFP5 probably regulate leaf development through various molecular pathways.

The TALE homeodomain proteins are fundamental regulators of plant leaf development and meristem function (Blanco et al., 2009). These homeodomain TFs is conserved in animals, plants, and fungi. In plants, TALE proteins are comprised of two classes, KNOTTED1-LIKE homeobox (KNOX) and BEL1-
like homeobox (BELL), which are involved in the determination and morphological development of leaves, stems, and inflorescences (Kumar et al., 2007; Hamant and Pautot, 2010; Hay and Tsiantis, 2010). The KNOX/BELL homo- and hetero-dimerization play an essential role in general TALE protein function (Hackbusch et al., 2005). Further, Hackbusch et al. (2005) also demonstrated that KNOX and BELL proteins interact with nine AtOFP proteins, regulating plant meristem and leaf development. Many other studies also showed that OFP proteins could interact with KNOX and BLH to regulate plant growth and development (Pagnussat et al., 2007; Li et al., 2011). Therefore, it is well-established that OFPs could act as the central point to interact with multiple proteins to form different regulatory complex to accurately and efficiently regulate plant growth and development. In our study, the physical interaction between BIOFP3 and BIKNOX5 as well as BIOFP5 and BIKNOX5 was detected by Y2H and BiFC experiments (Figures 9, 10). This implies that BIKNOX5 should be a significant component for the complex formed by BIOFP3 or BIOFP5. Besides, several other TALE proteins, including BIKNOX5, BIKNOX9, BIBLH6 and BIBLH7 proteins, were identified to interact with BIOFP3 or BIOFP5 by either Y2H or BiFC (Figures 9, 10). Although no consistent results were obtained from Y2H and BiFC, it is possible that BIOFP3 or BIOFP5 could interact with these TALE proteins to form various complexes, which may be associated with different biological processes in B. luminifera, but further experiments are needed to analyze the underlying molecular mechanism.

Conclusion

In the current study, we identified 20 BIOFP genes and analyzed the phylogenetic analysis, conserved motifs and cis-elements. The qRT-PCR analysis reveals that most BIOFP genes were up-regulated in YL2 compared to YL1 and bract. The overexpression of BIOFP3 and BIOFP5 genes in A. thaliana produces sawtooth shape, flatter, darker green rosette leaves, indicating their involvement in leaf development. Moreover, TEM results showing a significant increase in the number and size of palisade tissue cells in transgenic plants, which resulted in an increase in leaf thickness. Besides, expression analysis also indicated that the expressions of several genes related to leaf development were considerably changed in transgenic plants. Furthermore, physical interactions between BIOFP3, BIOFP5 and several TALE proteins were also identified. On the basis of the interaction assay, we hypothesized that at least BIKNOX5 is a...
common interacting protein for BIOFP3 and BIOFP5, forming a functional complex to regulate leaf development in *B. luminifera*.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

Author contributions

EL and HH designed the research. PB, FN, WY and HZ performed the experiments. JY performed the BiFC assay and revised the manuscript. PB, FN, WY, HZ, SC and XH analyzed the data. PB, EL and HH wrote the manuscript with comments from all authors. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.950936/full#supplementary-material

References

Belles-Boix, E., Hamant, O., Witäk, S. M., Morin, H., Traas, J., and Pautot, V. (2006). KNAT6: An arabidopsis homeobox gene involved in meristem activity and organ separation. *Plant Cell* 18 (8), 1900–1907. doi: 10.1105/tpc.106.041988

Blanco, F., Salinas, P., Cecchini, N. M., Jordana, X., Van Hummelen, P., Alvarez, M. E., et al. (2009). Early genomic responses to salicylic acid in arabidopsis. *Plant Mol. Biol.* 70, 79–102. doi: 10.1007/s11103-009-9458-1

Cai, M., Huang, H., Ni, F., Tong, Z., Lin, E., and Zhu, M. (2018). RNA-Seq analysis of differential gene expression in betula luminifera xylem during the early stages of tension wood formation. *PeerJ* 6, e5427. doi: 10.7717/peerj.5427

Chen, C., Chen, H., Zhang, Y., Thomas, H. R., Frank, M. H., He, Y., et al. (2020). TBtools: An integrative toolkit developed for interactive analyses of big biological data. *Mol. Plant* 13, 1194–1202. doi: 10.1016/j.molp.2020.06.009

Clough, S. J., and Bent, A. F. (1998). Floral dip: A simplified method for agrobacterium-mediated transformation of arabidopsis thaliana. *Plant J.* 16, 735–743. doi: 10.1046/j.1365-313x.1998.00343.x

Forslund, K., and Sonnhammer, E. L. J. E. G. (2012). “Evolution of protein domain architectures,” in *Evolutionary genomics: methods in molecular biology*, vol. 856. Ed. M. Anasimova (Totowa, New Jersey, USA: Humana Press), 187–216.

Hackbusch, J., Richter, K., Muller, I., Salamini, F., and Uehig, J. F. (2005). A central role of arabidopsis thaliana ovate family proteins in networking and subcellular localization of 3-aa loop extension homeodomain proteins. *PNAS* 102, 4908–4912. doi: 10.1073/pnas.0501181102

Hamant, O., and Pautot, V. (2010). Plant development: A TALE story. *C. R. Biol.* 333 (4), 371–381. doi: 10.1016/j.crvi.2010.01.015

Hasson, A., Plessis, A., Bein, T., Adroher, B., Grigg, S., Tsiantis, M., et al. (2011). Evolution and diverse roles of the CUP-SHAPED COTYLEDON genes in arabidopsis leaf development. *Plant Cell* 23 (1), 54–68. doi: 10.1105/tpc.110.081448

Hay, A., Barkoulas, M., and Tsiantis, M (2006). ASYMMETRIC LEAVES1 and auxin activities converge to repress *BREVIPEDICELLUS* expression and promote leaf development in *Arabidopsis*. *133*, 20, 3955–3961. doi: 10.1242/dev.02545

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Supplementary material

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Supplementary figure 1

OVATE domain alignment of BIOFP proteins.

Supplementary figure 2

Sequence alignment of AIOFP6, BIOFP3 and BIOFP6.

Supplementary figure 3

Transgenic Arabidopsis plants confirmation with PCR and qRT-PCR. (A) PCR amplification of BIOFP3 and BIOFP5 genes in transgenic lines and WT. (B) PCR amplification of hygromycin and GUS in WT and 35S::BIOFP3 transgenic lines. (C) PCR amplification of hygromycin and GUS in WT and 35S::BIOFP5 transgenic lines. (D) Expression analysis of BIOFP3 in 35S::BIOFP3 transgenic lines by qRT-PCR. (E) Expression analysis of BIOFP5 in 35S::BIOFP5 transgenic lines by qRT-PCR. The size of the PCR products indicated in Kb. The asterisk indicates significant difference compared to WT at p<0.05 level.

Supplementary figure 4

Expression analysis of AIOFP6 in transgenic plants.

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10.3389/fpls.2022.950936
Hay, A., and Tsiantis, M. (2009). A KNOX family tale. *Curr. Opin. Plant Biol.* 12 (5), 593–598. doi:10.1016/j.pbi.2009.06.006

Hay, A., and Tsiantis, M. (2010). KNOX genes: Versatile regulators of plant development and diversity. *Development* 137 (19), 3153–3165. doi:10.1242/dev.030049

Knaap, E. V. D., and Ostergaard, L. (2018). Shaping a fruit: Developmental pathways that impact growth patterns. *Semin. Cell Dev. Biol.* 79, 27–36. doi:10.1016/j.semcdb.2017.09.028

Kumar, R., Kushalappa, K., Godt, D., Pidkowich, M. S., Pastorelli, S., Hepworth, S. R., et al. (2007). The arabidopsis BEL1-LIKE HOMEODomain proteins SAW1 and SAW2 act redundantly to regulate KNOX expression spatially in leaf margins. *Plant Cell* 19 (9), 2719–2735. doi:10.1105/tpc.106.048769

Kumar, S., Stecher, G., Li, M., Knyaz, C., and Tamura, K. (2018). MEGA X. Molecular evolutionary genetics analysis across computing platforms. *Mol. Biol. Evol.* 35, 1547–1550. doi:10.1093/molbev/msy096

Li, E., Bhargava, A., Qiang, W., Friedmann, M. C., Forneris, N., Savidge, R. A., et al. (2012). The class II KNOX gene KNAT7 negatively regulates secondary wall formation in arabidopsis and is functionally conserved in populus. *New Phytol.* 194, 102–115. doi:10.1111/j.1469-8137.2011.04016.x

Li, H., Dong, Q., Zhu, X., Zhao, Q., and Ran, K. (2019). Genome-wide identification, expression, and interaction analysis for ovate family proteins in peach. *Mol. Reprod. Dev.* 4 (4), 3755–3764. doi:10.1002/mrd.21048

Li, Z., Li, B., Liu, J., Guo, Z., Liu, Y., Li, Y., et al. (2016). Transcription factors AS1 and AS2 interact with LHP1 to repress KNOX genes in arabidopsis. *J. Integr. Plant Biol.* 58 (12), 959–970. doi:10.1111/jipb.12485

Li, X. Y., Lin, E. P., Huang, H. H., Niu, M. Y., Tong, Z. K., and Zhang, J. H. (2018). Molecular characterization of SQUAMOSA PROMOTOR BINDING PROTEIN-LIKE (SPL) gene family in betula luminafrina. *Front. Plant Sci.* 9, 608. doi:10.3389/fpls.2018.00608

Li, Z., Li, B., Shen, W. H., Huang, H., and Dong, A. (2017). TCP transcription factors interact with AS2 in the repression of class I KNOX genes in arabidopsis thaliana. *Plant J.* 71 (1), 99–107. doi:10.1111/tpj.13135.2012.04973.x

Li, Y., and Douglas, C. J. (2015). A role for Ovate FAMILY PROTEIN1 (OPF1) and OPF2 in a BLH6-KNAT7 multi-protein complex regulating secondary cell wall formation in arabidopsis thaliana. *Plant Signal Behav.* 10, e1033126. doi:10.1080/15592324.2015.1033126

Liu, D., Sun, W., Yuan, Y., Zhang, N., Hayward, A., Liu, Y., et al. (2014). Phylogenetic analyses provide the first insights into the evolution of OVATE family protein1 interaction with BLH3 regulates transition timing from vegetative to reproductive phase in arabidopsis. *Comm. Integr. Biol.* 7, 1–16. doi:10.4172/2151-9685-7-1

Wang, S., Chang, Y., and Ellis, B. (2016). Overview of Ovate FAMILY PROTEINS, a novel class of plant-specific growth regulators. *Front. Plant Sci.* 7, 417. doi:10.3389/fpls.2016.00417

Wang, S., Chang, Y., Guo, J., and Chen, J. G. (2007). Arabidopsis ovate family protein 1 is a transcriptional repressor that suppresses cell elongation. *Plant J.* 50, 858–872. doi:10.1111/j.1365-313X.2007.01396.x

Wang, S., Chang, Y., Guo, J., Zeng, Q., Ellis, B. E., and Chen, J. G. (2011). Arabidopsis ovate family proteins, a novel transcriptional repressor family, control multiple aspects of plant growth and development. *Proc. Natl. Acad. Sci. U. S. A.* 108, 20525–20530. doi:10.1073/pnas.1108360108

Wang, S., Chen, J., Yu, H., and Tsiantis, M. (2021). A KNOX family tale. *Integr. Plant Biol.* 6, e1006118. doi:10.1093/ipb/diaa005

Wang, H., Kong, F., and Zhou, C. (2021). From genes to networks: The genetic control of leaf development. *Integr. Plant Biol.* 6, 1181–1196. doi:10.1093/tpj/jdp088

Wu, S., Zhang, B., Keyhaninejad, N., Rodriguez, G. R., Kim, H. J., Chakrabarti, M., et al. (2018). A common genetic mechanism underlying morphological diversity in fruits and other plant organs. *Nat. Commun.* 9, 4734. doi:10.1038/ncomms10716

Yang, C., Ma, Y., He, Y., Tian, Z., and Li, J. (2018). OsOFP19 modulates plant architecture by integrating the cell division pattern and brassinosteroid signaling. *Plant J.* 93, 489–501. doi:10.1111/tpj.13793

Yang, C., Shen, W., Li, Y., Tian, Z., and Li, J. (2016). OVATE family protein 8 positively mediates brassinosteroid signaling through interacting with the GSK3-like kinase in rice. *Pl. Sci. Genet.* 12, e006118. doi:10.1371/journal.pgen.1006118

Yoo, S. D., Cho, Y. H., and Sheen, J. (2007). Arabidopsis mesophyll protoplasts: A versatile cell system for transient gene expression analysis. *Nat. Protoc.* 2, 1565–1572. doi:10.1038/nprot.2007.199

Zhang, L., Zhang, X., Lu, H., Chen, J., Wang, S., Wang, H., et al. (2016). Ovate family protein1 interaction with BLH3 regulates transition timing from vegetative to reproductive phase in arabidopsis. *Biochem. Biophys. Res. Commun.* 470, 492–497. doi:10.1016/j.bbrc.2016.01.135

Zhou, S., Cheng, X., Li, F., Peng, F., Hu, G., Chen, G., et al. (2019). Overexpression of SIOPFP20 in tomato affects plant growth, chlorophyll accumulation, and leaf senescence. *Front. Plant Sci.* 10, 510. doi:10.3389/fpls.2019.01510

Zhou, R., Fan, M., Zhao, M., Jiang, X., and Liu, Q. (2022). Overexpression of LkNOX1 from *Lilium tajovitae* in *Nicotiana benthamiana* affects the development of leaf morphology. *Plant Signal Behav.* 17, 1. doi:10.1080/15592324.2022.2031783