Abstract: The tree peony (Paeonia section Moutan DC.) is the candidate flower in China, with abundant germplasm resources and high ornamental value. However, the short and concentrated flowering period severely restricted the improvement of the economic value of tree peonies. Based on the full-length transcriptome database of tree peonies, the PoVIN3 (GenBank ID: OP341879), involved in the flowering regulation of tree peonies were identified and cloned for the first time. The PoVIN3 was also characterized by bioinformatics methods, quantitative real-time PCR (qRT-PCR), and the establishment of a transgenic system. The expression levels of PoVIN3 in seven different petals developmental stages were the highest at the initial flowering stage of the variant cultivar of Paeonia ostii ‘Fengdan,’ the initial decay stage of the normal flowering Paeonia ostii ‘Fengdan,’ and the half opening stage of the late flowering Paeonia suffruticosa ‘Lianhe.’ Tissue-specific expression analysis showed that the relative expression levels of PoVIN3 were the highest in sepal of both normal flowering Paeonia ostii ‘Fengdan’ and the late flowering Paeonia suffruticosa ‘Lianhe,’ and the highest expression was in stamens of early flowering mutant Paeonia ostii ‘Fengdan.’ In addition, the flowering time of pCAMBIA2300-PoVIN3 transgenic plants was significantly earlier than that of the wild-type, indicating that PoVIN3 could promote plant flowering. The results provide a theoretical basis for exploring the role of PoVIN3 in the regulation of flowering in tree peonies.

Keywords: tree peony; PoVIN3; flowering regulation; expression analysis; functional analysis

1. Introduction

The tree peony (Paeonia section Moutan DC.) is a famous traditional flower in China. It has a high ornamental, cultural, and economic value with big flowers, bright colors, and elegant appearance, and is renowned as a symbol of Chinese civilization [1]. The research on ornamental characteristics of tree peony has made continuous progress from breeding varieties to cultivation technology, which has been widely cultivated worldwide, and now contains as many as 2200 ornamental varieties [2]. However, the flowering time of tree peonies is short and concentrated, and a widespread social consensus and regret with the short period of flowering and viewing. Under natural conditions, it takes 50–60 days from flower bud initiation to petal decay, and the flowering time is 3–5 days for a single flower (a single flower from the full blooming stage to the decay stage) and 10–15 days for the various group (the flowering period from the first to the last flower of the same variety). Most tree peonies are middle–flowering varieties, with smaller numbers of early– and late–flowering varieties [3,4]. The flowering period determines the commercial value of tree peonies [5,6]. Consequently, a thorough understanding of the genetic basis and molecular regulatory mechanisms related to the regulation of tree peony flowering will help to meet the demand of people for prolonged ornamental traits in tree peonies.

Flowering is an important process for higher plants to achieve the transition from vegetative to reproductive growth. The MADS–box transcription factor named the FLOWERING LOCUS C (FLC) is a key gene in the Arabidopsis thaliana flowering regulatory network.
and plays a pivotal role in flowering regulation [7]. There are many genes controlling flowering in the autonomous pathway and vernalization pathway, and their functions were realized by regulating the expression of FLC. FLC inhibited flower formation mainly by acting on the flowering integration genes FLOWERING LOCUS T (FT) and SUPPRESSOR of OVEREXPRESSION of CO 1 (SOC1). FT and SOC1 integrated signals from flowering pathway signals and are called floral pathway integrators. FLC inhibited their expression by binding to the CArG box of the SOC1 promoter and part of the FT first intron (containing the CArG box), respectively [8,9], thereby inhibiting floral meristem determining genes and thus delaying flower formation. Vernalization mainly causes changes in the chromatin structure of FLC so as to relieve its inhibitory effect on flowering [10].

VERNALIZATION INSENSITIVE 3 (VIN3) has been identified as an essential gene for triggering vernalization and inhibiting FLC, and the induction of VIN3 expression was the earliest event of vernalization. Together with VERNALIZATION 1 (VRN1) and VERNALIZATION 2 (VRN2), VIN3 forms a complex that acts on histones on FLC chromatin to deacetylate and methylate them, thereby inhibiting FLC expression, and the plant exhibits early flowering [11–13]. At low temperatures, VIN3 inhibited FLC expression by recognizing the time of low-temperature treatment during vernalization and acting together with VRN1, VRN2, and VRN5 [14]. VIN3 had the characteristic of sensing the time course of low temperature. The expression of VIN3 could only be induced under long-term low-temperature treatment, and VIN3 could not be detected after the plant returned to normal temperature, but FLC was still inhibited. Therefore, the role of VIN3 is to identify the time of low-temperature treatment in the vernalization process and establish the suppression of the expression of FLC [13]. VIN3 encodes a PHD–finger protein, which can be involved in the methylation and deacetylation of nucleosome histones, causing the remodeling of chromatin structure [15]. PHD domain is a class of conserved C4CH3 (Cys4–His–Cy3) structure composed of about 60 amino acid residues, belonging to the “cross-brace” zinc finger protein family, which can change the spatial structure of chromatin to cause gene silencing [16,17]. Upon induction of VIN3, which is essential for PHD–PRC2 nucleation, H3K9 and H3K27 on the histones of FLC chromatin were dimethylated and trimethylated, thus showing genetic silencing and plants showing early flowering [18,19].

According to previous reports on the VIN gene family in Arabidopsis thaliana and Poaceae, this study focused on the key gene VIN3 in upstream of the vernalization pathway in the flowering network. The PoVIN3 related to flowering regulation was cloned from a tree peony for the first time. The sequence of PoVIN3 was analyzed by biological software, and the expression of PoVIN3 in tree peonies was detected by qRT–PCR, which provided a theoretical basis for studying the molecular mechanism of tree peony flowering regulation.

2. Results
2.1. Cloning and Sequence Analysis of PoVIN3

The target fragment was obtained by using the petal cDNA of FD as a template and named PoVIN3. The gene sequence was submitted to GenBank (OP341879) in NCBI. The complete open reading frame (ORF) of PoVIN3 is 1740 bp in total (Figure 1A,B) and encodes a polypeptide with 579 amino acids. The predicted molecular mass of PoVIN3−encoded protein was found to be 64.76 kD, and the isoelectric point (pI) was 7.62. The instability coefficient was 49.66 and was classified as unstable. In addition, the total average hydrophilicity was −0.400, indicating that the protein was hydrophilic (Figure 1C). The analysis of conserved domains showed that the protein contained two domains, including the PHD-SF (Accession: PFAM07227) superfamily region present at amino acids 60–179 and the 286–343 amino acids containing the FN3 (Accession: Smart100060) superfamily conserved domain (Figure 1D). The predicted protein secondary domain shows that the protein amino acids mainly exist as alpha helices (178 amino acids), extended strands (74 amino acids), random coils (297 amino acids), and beta helices (30 amino acids) (Figure 1E). Furthermore, according to the template 1zlgA model, the confidence score (C-score) value is −2.03 (Figure 1F).
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Figure 1. Cloning and bioinformatics analysis of PoVIN3. (A) PCR amplified product of PoVIN3, and M means DNA maker DL2000. (B) Nucleotide sequence and its encoded amino acid sequence. (C) Hydrophilic analysis of PoVIN3-encoded protein. (D) Analysis of conserved domain of PoVIN3. (E) Secondary structure prediction of PoVIN3-encoded protein. (F). Tertiary structure prediction of PoVIN3-encoded protein.

2.2. Phylogenetic Tree Analysis

In order to study the evolutionary relationship between PoVIN3 of tree peony and VIN3 in other plants, VIN3 coding amino acid sequences of different plants were selected from NCBI for comparative analysis. The PoVIN3-encoded protein was obtained from Camellia sinensis (accession number: XP_028064280), Ricinus communis (accession number: XP_015572530), Ziziphus jujuba var. spinosa (accession number: XP_048320875), Vitis Riparia (accession number: XP_034700197), Pistacia Vera (accession number: XP_031256809), and Populus Trichocarpa (accession number: XP_024446061) had high homology of 100%, 98%, 98%, 98%, and 96%, respectively (Figure 2A).
Figure 2. Multiple sequence alignment (A) and phylogenetic relationship (B) of PoVIN3−encoded protein and VIN3−encoded protein from other plant species. The red asterisk represented the PoVIN3−encoded protein.

The phylogenetic tree of 28 different plants was constructed, including *Populus trichocarpa* (XP_024446061, XP_024447843), *Ziziphus jujube* (XP_015898363), *Pistacia vera* (XP_031256809, XP_031256810), *Populus alba* (XP_034921479, XP_034921480, XP_034913370), *Actinidia chinensis* var. *chinensis* (PSS01860), *Gossypium hirsutum* (XP_016690001, XP_040949013), *Camellia sinensis* (XP_028064280), *Salix suchowensis* (KAG5225613), *Ricinus communis* (XP_015572530), *Morella rubra* (KAB1201411), *Prunus avium* (XP_021807197), *Gossypium arboretum* (KHG19667), *Prunus persica* (XP_024423658), *Mangifera indica* (XP_044464405, XP_044464404, XP_044464406), *Tripterygium wilfordii* (XP_038712221, XP_038712219, XP_038712220), *Carya illinoensis* (XP_042977516, XP_042977519, XP_042977518), *Heterocistus* (XP_021667538, XP_021667537, XP_021666091, XP_021666090), *Morus notabilis* (XP_024029764, EXB37057), *Juglans regia* (XP_018822838), *Citrus sinensis* (XP_006475725), *Citrullus lanatus* (XP_024033728), *Juglans microcarpa* × *Juglans regia* (XP_041005313), *Jatropha curcas* (XP_012078661), *Herrania unbratia* (XP_021275899, XP_021275900), *Manihot esculenta* (XP_021597269), *Cannabis sativa* (XP_030488315, XP_0304883160), *Durio zibethinus* (XP_022729825), and *Vitis riparia* (XP_034700197). Compared with the PoVIN3−encoded protein, the relationship between the VIN3−encoded protein of *Vitis riparia* and that of *Salix suchowensis* was close. VIN3−like protein (XP_021597269) from *Manihot esculenta* was phylogenetically closest to PoVIN3−encoded protein (Figure 2B).

2.3. Expression Pattern of PoVIN3

In order to determine the spatiotemporal expression patterns of PoVIN3, the expression levels were investigated by qRT−PCR within samples taken from various tissues and petals at different developmental stages of VC, FD, and LH. The qRT−PCR results demonstrated that the expression levels of PoVIN3 in LH were significantly higher than that in VC and FD at different developmental stages (CE, BS, IF, HO FB, ID, and DS) (Figure 3A). In VC, PoVIN3 had a high expression level in the early stage of flower development (CE, BS, and IF), and the expression level was significantly higher in the IF stage ($p < 0.05$). The expression levels were significantly lower in the HO and FB stages and increased again in the decline process (ID and DS) but still lower than that in the early stage (Figure 3A). During FD flower development to flower blooming, PoVIN3 expression levels decreased
first and then increased, reached a significantly lower level at IF, and then increased sharply at ID, reaching a significantly higher level (p < 0.05) (Figure 3A). The PoVIN3 expression levels of LH were different from that of the former two. With the progress of flower development, PoVIN3 expression levels gradually increased and reached a significantly higher level at HO (p < 0.05). The expression level of PoVIN3 decreased abruptly in the FB stage and increased abruptly in the ID stage. When the petals decayed, the PoVIN3 expression level decreased to a significantly lower level (p < 0.05) (Figure 3A).

PoVIN3 was expressed in all tissues collected from the three tree peony cultivars, including bracts, sepals, carpels, stamens, leaves, and petals. However, the expression level in the leaves of FD was at the lower limits of detection (Figure 3B). It is worth noting that the expression of PoVIN3 was mainly concentrated in stamens, while the lowest expression level was found in petals (Figure 3B).

2.4. Subcellular Localization

Prediction of subcellular localization suggested that PoVIN3—encoded protein was localized in the nucleus. A fusion expression vector pCAMBIA2300—PoVIN3—GFP was transiently transformed in N. benthamiana to determine the subcellular localization of PoVIN3—encoded protein, while the pCAMBIA2300—GFP was a control. Subsequently, confocal scanning microscopy was used to observe the Agrobacterium tumefaciens—infected leaves of N. benthamiana containing the gene of interest. This study found that the pCAMBIA2300—GFP emitted green fluorescence throughout the cells under the excitation of 488 nm wavelength, while the strong fluorescence signals were distributed in the nucleus for the pCAMBIA2300—PoVIN3—GFP, consistent with the prediction. Thus, indicating that PoVIN3 was indeed nuclearly localized (Figure 4).
Figure 4. Subcellular localization of GFP fusions of PoVIN3.

2.5. Ectopic Expression of PoVIN3 Accelerate Flowering in Arabidopsis Thaliana

The spatiotemporal expression pattern of PoVIN3 suggested that it might play a key role in the regulation of flowering time and flower development in plants. To prove that PoVIN3 can promote the early flowering of Arabidopsis thaliana, four groups of transgenic Arabidopsis thaliana plants over-expressing pCAMBIA2300–PoVIN3–1, pCAMBIA2300–PoVIN3–2, pCAMBIA2300–PoVIN3–3, and pCAMBIA2300–PoVIN3–4, respectively, were generated (Figure 5A). All four groups of transgenic plants showed an early flowering phenotype compared with the control plants transformed with an empty vector and WT plants, and the flowering time was significantly advanced by 5.67–7.67 days (p < 0.05) (Figure 5B). At the same time,40–day-old leaves of T2 plants were harvested, and the results of RT–PCR analysis of PoVIN3 also proved that the transgenic lines obtained specific amplified bands of the expected size; by contrast, no specific bands were detected in either WT or pCAMBIA2300 plants (Figure 5C). These results suggest that PoVIN3 plays an important role in the flowering process of plants under long-day conditions.

2.6. pCAMBIA2300-PoVIN3 Overexpressed Tree Peony Petals

To test the possible role of PoVIN3 in tree peony petals, we overexpressed PoVIN3 in tree peony petals by transient expression technique. During dehydration, Paeonia suffruticosa ‘Luoyanghong’ petals infected with the pCAMBIA2300–PoVIN3 overexpression vector generally showed a slightly longer length than petals infected with pCAMBIA2300 empty vector, but there was no significant difference (p > 0.05) (Figure 6A(a,e)). After 12 h of dehydration, the relative width of petals infected with pCAMBIA2300–PoVIN3 was about 52.42% ± 1.12% of its initial value (Figure 6A(b,e)), and the relative fresh weight of all petals was about 44.24% ± 1.02% of its initial value, 5.18% as compared with the empty vector control (41.95% ± 1.06%) (p > 0.05) (Figure 6A(c,e)). During 12 h of rehydration, petals infected with pCAMBIA2300–PoVIN3 returned to 89.51% ± 7.28% (the relative length of petals), 71.73% ± 8.25% (the relative width of petals), and 73.93% ± 1.16% (the relative fresh weight of petals) of their initial value. However, only 85.26% ± 0.89% (the relative length of petals), 71.01% ± 1.54% (the relative width of petals), and 65.14% ± 1.19% (the relative fresh weight of petals) of their initial value were infected with pCAMBIA2300 empty vector. Similarly, qRT–PCR showed that there was an extremely high expression in the petals of Paeonia suffruticosa ‘Luoyanghong’ after 9 h dehydration (Figure 6A(d)).
The effects of gene overexpression and dehydration were also tested on petal discs. In the process of dehydration, the relative fresh weight of petal discs infected with the pCAMBIA2300–PoVIN3 overexpression vector was slightly higher than that infected with pCAMBIA2300 empty vector, but there was no significant difference ($p > 0.05$) (Figure 6B(a,d)). However, after 12 h of rehydration, petal discs infected with pCAMBIA2300 empty vector could recover to 91.02% ± 0.73% of their initial value, which was significantly higher than that infected with overexpressed bacteria ($p < 0.05$) (Figure 6B(b)). The relative expression of PoVIN3 in the overexpressed infected petal discs during dehydration was higher than that in the empty vector control and reached the peak at 12 h after dehydration (Figure 6B(c)).

Notably, in the infection process of fresh cut flowers of Paeonia suffruticosa ‘Luoyanghong,’ it was found that the fresh cut flowers infected with pCAMBIA2300–PoVIN3 reached FB 7 h earlier than that of empty vector infection, which was consistent with the results obtained from the stable expression of Arabidopsis thaliana, indicating that PoVIN3 may advance the flowering time of tree peony (Figure 6C(a)), but there was no significant difference in the relative fresh weight between them ($p > 0.05$) (Figure 6C(b)). The relative expression level of PoVIN3 in the fresh-cut flowers of Paeonia suffruticosa ‘Luoyanghong’ was detected. It was found that the relative expression level of PoVIN3 in the petals infected with overexpression reached the highest at 42 h after transient expression, and a certain amount of PoVIN3 was still expressed at 70 h (Figure 6C(c)).
Figure 6. Transient expression of PoVIN3 was overexpressed in tree peony petals, discs, or fresh-cut flowers. (A) Tree peony petals were infiltrated with Agrobacterium tumefaciens containing pCAMBIA2300 control and pCAMBIA2300-PoVIN3 for dehydration (0, 3, 6, 9, 12 h) and rehydration for 12h. The relative length (a), relative width (b), relative fresh weight (c), relative expression (d), and phenotype (e) of petals were analyzed. (B) Phenotype and recovery of PoVIN3-overexpressed petal discs. The relative fresh weight (a), relative area (b), relative expression (c), and phenotype (d) of petal discs were analyzed. (C) Overexpression infection in fresh cut flowers of tree peony. The phenotype (a), relative fresh weight (b), and relative expression (c) of fresh cut flowers were analyzed. Error bars represent standard error (SE). Different lowercase letters (a–d) indicate significant differences at $p < 0.05$, * means $p < 0.05$, and *** means $p < 0.001$. 
3. Discussion

Vernalization of plants is of great significance to the development of economic benefits, and many plants need to be exposed to the cold for a period of time, that is, undergo the vernalization process, to induce flowers, as this indicates the end of winter and avoids mortality from premature flowering in the fall [20]. Based on the full—length transcriptome database of tree peonies previously constructed in our laboratory, genes related to vernalization were screened. Primers were designed to clone the target gene. PoVIN3 (GenBank ID: OP341879) was cloned from a tree peony for the first time by RT—PCR, which was also the first time to obtain key genes in the vernalization pathway from ornamental plants, and its expression characteristics and functions were preliminarily studied.

Sequence analysis showed that the vernalization—related gene PoVIN3 had high homology with the corresponding gene in Camellia sinensis. PoVIN3 contains two domains, namely, the PHD—SF superfamily and the FN3 superfamily. This was consistent with the structure of VIN3 in Arabidopsis thaliana [13], Brassica oleracea L. var. capitata L. [21], and Brassica campestris ssp. chinensis var. pupurea Hort. [22]. PHD—finger protein is one of 14 known zinc—binding motifs, which are found in more than 400 eukaryotic proteins and are highly conserved during evolution [23]. Three homologous genes of VIN3 were cloned from Triticum monococcum L., namely TmVIL1(VIN3—like1), TmVIL2, and TmVIL3, and also four VEL family genes (VIN3 homologous gene family) were obtained from Oryza Sativa (OsVIL1, OsVIL2, OsVIL3, and OsVIL4). They all encode proteins with the zinc—binding motif of Cys4 (Cys4—His—Cys3), which is conserved in the PHD domain [15]. Studies have shown that the PHD domain is involved in many functions, including protein methylation, acetylation, phosphorylation, ubiquitination, and so on [15,24]. In this study, the results of subcellular localization showed that PoVIN3—encoded protein was localized in the nucleus, indicating that PoVIN3 had the function of encoding protein. Therefore, we speculated that PoVIN3 might act on FLC through the PHD domain, thereby shutting down FLC expression and inhibiting FLC function.

In this study, it was found that the expression trend of PoVIN3 was not the same among the three tree peony varieties, and the overall expression trend was that the late—flowering Paeonia suffruticosa ‘Lianhe’ was higher than the early—flowering and normal flowering Paeonia ostii ‘Fengdan’ at different stages of flower development, indicating that PoVIN3 may have variety specificity. The results were similar to the expression pattern of BcVIL1 (VIN3—LIKE1 belongs to the VIN3 family gene) in non—heading Chinese cabbage [25]. VIN3 is a protein produced only in cold winters, and its cold—inducing ability is related to the degree of vernalization [26]. Numerous experiments have shown that VIN3 plays a role in mediating proper vernalization in Arabidopsis [27,28]. In the process of mango flowering induction, it was also confirmed that VIN3 could be expressed or up—regulated only after prolonged low—temperature induction [29]. Although the changes of PoVIN3 expression patterns during cold vernalization were not involved in this study, the expression of PoVIN3 in tree peonies from flower formation to decay was demonstrated, which provided a new idea for the study of this gene. In addition, we demonstrated that there were significant differences in the expression of PoVIN3 in various organs and tissues of tree peonies. As expected, PoVIN3 expression was abundant in various floral organs and tissues. These results suggested that PoVIN3 remains active during floral organ differentiation and growth. In contrast to our results, the expression of rice VIN3 was high in leaves, moderate in seedlings, roots, and presbyteria, and low in stems and young flowers [30]. The lower expression of VIN3 in floral tissues may be due to the fact that rice plants do not require vernalization, unlike plants such as Arabidopsis, cereal, and tomato, which show a strong flowering response to vernalization [31,32].

Previous studies have confirmed that the function of VIN3 was to inhibit the expression of FLC, a key gene for vernalization, so as to promote flowering [33,34]. The tree peony is a traditional Chinese flower, and the research on the mechanism of regulating tree peony flowering has always been the focus of researchers [35,36]. In this study, PoVIN3 was cloned from the petals of Paeonia ostii ‘Fengdan’ at the full blooming stage. In order to
investigate the function of PoVIN3, an important gene in the vernalization pathway that regulates flowering, we constructed a pCAMBIA2300–PoVIN3 overexpression vector and transformed it into Arabidopsis Thaliana. According to the statistics of the flowering time of wild-type and transgenic Arabidopsis thaliana under long-day conditions, the flowering time of the transgenic Arabidopsis thaliana was significantly advanced by 5.67–7.67 days ($p < 0.05$) (Figure 5B). It was speculated that PoVIN3 might promote flowering. Based on the current experimental studies, further studies on the relationship between PoVIN3 and other transcription factors and its role in the vernalization process of peony should be the focus of future work. Due to the large reference genome and many repetitive sequences in tree peony, the tree peony ontology cannot be used for functional verification of genes at present. Therefore, the model plant Arabidopsis Thaliana was selected to verify the possible function of PoVIN3 in this experiment. This method is often used for functional gene verification in tree peonies [37]. In addition, the methods for verifying the gene function of Arabidopsis thaliana transformation are also common in plants with large genomes and imperfect genetic transformation systems, such as herbaceous peonies [38] and cotton [39]. Therefore, it is reliable to select Arabidopsis Thaliana as the receptor for verifying PoVIN3 function in this study.

The progress of genetic transformation in tree peonies has been hampered by long-term regeneration schemes, which are required for successful ontological genetic transformation in tree peonies, and the low efficiency of regeneration systems further aggravates the establishment of genetic transformation systems in tree peonies. At present, the identification of functional genes in tree peonies rely on model plants such as Arabidopsis thaliana, but it takes a long time to analyze the function of introduced transgenes [40]. Therefore, using transient expression of the gene to determine gene function was an effective alternative method while avoiding the problems of low transformation efficiency and long generation time of transgenic plants. The establishment of efficient and rapid gene function analysis methods in plants has become a research hotspot in recent years [41]. A transient expression system could overexpress or silence target genes in plant cells in a short period of time, which greatly accelerates the process of gene function research. In rose [42], kiwifruit [43], lily [44], grape [45], strawberry [46], and petunia [47]. The transient expression system was established in plants by the Agrobacterium tumefaciens–mediated method. In this study, the transient transformation system of tree peony overexpression was also established by Agrobacterium tumefaciens. The results laid a foundation for the verification of the function of PoVIN3 in tree peony and also provided research ideas and a technical basis for the verification of other important functional genes in tree peonies.

Based on the above results, PoVIN3 may play a role in regulating flowering time. This is the first time to clone and obtain the cDNA ORF coding region of vernalization-related gene PoVIN3 in tree peony, which has great reference significance for further study of vernalization-mediated flowering pathway in tree peony and can lay a foundation for improving the ornamental and economic value of tree peony.

4. Materials and Methods

4.1. Plant Materials and Growth Conditions

For the analysis of gene expression patterns, plant materials were obtained from the tissues (including bracts, sepals, carpels, stamens, leaves, and petals) of a variant cultivar of Paeonia ostii ‘Fengdan’ that exhibits earlier flowering (VC), a normal flowering Paeonia ostii ‘Fengdan’ (FD), and a late flowering Paeonia suffruticosa ‘Lianhe’ (LH). All the species used as test material was planted in the experimental field of Henan University of Science and Technology (112°28′36.34″ E, 34°39′30.34″ N). In order to profile the changes in gene expression during different stages of flower development, petals from the same plant of three varieties at seven different developmental stages were sampled. These different flower developmental stages included the color-exposure stage (CE), blooming stage (BS), initial flowering stage (IF), half opening stage (HO), full blooming stage (FB), initial decay stage (ID), decay stage (DS). All samples were collected in April 2020.
The genetic transformation experiments used the seedlings of *Arabidopsis thaliana* cultivar ‘Columbia’ (Col−0) as the recipient plants, and the subcellular localization analysis used *N. benthamiana* plants. The transformed plant was cultivated in a tissue culture room (16 h:8 h, light:dark, 25 °C). The resulting transgenic plants were grown in a greenhouse (16 h:8 h, light:dark, 22 °C, 120–150 μmol·m−2·s−1 light intensity, 25–75% relative humidity).

4.2. Nucleic Acid Extraction and cDNA Synthesis

RNAPrep Pure Polysaccharide Polyphenol plant total RNA Extraction Kit (TianGen Biotech, Beijing, China) was used to extract total RNA from the petals at different developmental stages and different tissues (including bracts, sepals, carpels, stamens, leaves, and petals) of VC, FD, and LH. The quality and concentration of nucleic acid were determined by 1% agarose gel electrophoresis and NanoDrop One (Thermo Fisher Scientific, Waltham, MA, USA). Subsequently, 100 ng of total RNA was reversely transcribed into cDNA according to the PrimeScript™ II 1st Strand cDNA Synthesis Kit Reverse Transcription Kit (TaKaRa, Osaka, Japan).

4.3. Cloning of PoVIN3 and Construction of Expression Vectors

Using NCBI (http://www.ncbi.nlm.nih.gov) (accessed on 25 November 2021) and the full-length transcriptome database of tree peony based on Isoform-sequencing (Iso−seq), previously constructed in our laboratory, to screen potential PoVIN3 genes. Primer 5.0 software was used to design specific primers (Table 1), and the cDNA of the petals of FD at the full blooming stage was used as a template to amplify candidate genes. After ligation with the pMD−18T vector, the cells were transformed into *E. coli* DH5α competent cells and the positive clones were screened for sequencing verification.

Table 1. Primer sequence in this study.

| Primer Name | Sequence (5′-3′) | Use |
|-------------|-----------------|-----|
| PoVIN3      | F: ATGACGGATAATCCAAAGAC  
             | R: TTAATGCCATAGCTTACTGCT | Cloning the full−length of ORF |
| PoVIN3−qRT  | F: CAGGGATACCTGGGGGTTT  
             | R: TCTGCTGACAATGCCACG | Quantitative Real−time PCR |
| Tbulin−α    | F: CCGTCAACTTTTCCCCACCTG  
             | R: CCTCACTCGGTCAAGCCAGA | |
| pCAMBIA2300−PoVIN3−GFP | F: GGAGAGGACAGGGTACCATGACGGATAATCCAAA-GAC  
                          | R: GATCCCCCGGTACCATTCCATGCACTTACTGCAAGAC | Subcellular localization |
| pCAMBIA2300−PoVIN3 | F: GGAGAGGACAGGGTACCATGACGGATAATCCAAA-GAC  
                          | R: GGATCCCCCGGTACCCTTAATGCCCATAGCTTACTGCA | |
| AtVIN3      | F: CCGTAAAGACTGGCGAACAG  
             | R: CCGCACGAGTAACCCTGAT | Functional verification |

Plant expression vectors pCAMBIA2300 and pCAMBIA2300−GFP were digested with restriction enzyme *Kpn* I am using correctly sequenced positive cloned plasmids as templates. The overexpression vector pCAMBIA2300−PoVIN3 and the subcellular localization vector pCAMBIA2300−PoVIN3−GFP were obtained by seamless cloning technology, and the plasmid extracted from the correct bacterial solution was screened and sequenced to transform *Agrobacterium* GV3101 for subsequent tests.

4.4. Bioinformatics Analysis of PoVIN3

The sequencing results were analyzed by DNAMAN software. The amino acid composition of PoVIN3−encoded protein was analyzed by the online software Translate (http://web.expasy.org/translate/) (accessed on 10 January 2022). The physical and
chemical properties, hydrophobicity, transmembrane structure, and phosphorylation sites of the protein were analyzed by ProtParam (https://web.expasy.org/protparam/) (accessed on 10 January 2022), Protoscale (https://web.expasy.org/protoscale/) (accessed on 10 January 2022), TMHMM (http://www.cbs.dtu.dk/services/TMHMM-2.0/) (accessed on 10 January 2022), and NetPhos (http://www.cbs.dtu.dk/services/NetPhos/) (accessed on 10 January 2022). The conserved domains of PoVIN3 using the NCBI Conserved Domain Database (CDD) (http://www.ncbi.nlm.nih.gov/cdd/) search tools (accessed on 10 January 2022). SOPMA (https://npsa-prabi.ibcp.fr/cgi-bin/secpred_sopma.pl) (accessed on 10 January 2022) and I–TASSER (https://zhanggroup.org//I-TASSER/) (accessed on 10 January 2022) were used to predict the secondary structure, and three-dimensional homologous modeling of the protein, respectively. Other plants VIN3 sequences were collected from GenBank (http://www.ncbi.nlm.nih.gov/Genbank/) (accessed on 10 January 2022) and processed with default parameters using Clustal Omega (https://www.ebi.ac.uk/Tools/msa/clustalo/) (accessed on 10 January 2022) [48–50]. The phylogenetic tree was constructed by MEGA 7.0 software employing the Neighbor-Joining (NJ) method.

4.5. Analysis of Subcellular localization

The subcellular localization analysis used the 5–week-old seedlings of N. benthamiana plants. The constructed plasmid of pCAMBIA2300–PoVIN3–GFP was expanded in LB liquid medium containing 50 µg·mL⁻¹ kanamycin (kan) and then transformed into the lower epidermis of N. benthamiana leaves by the Agrobacterium tumefaciens-mediated leaf disk transformation method. The infected N. benthamiana was incubated in the dark for 2–3 h and then incubated in the light/dark time of 16 h /8 h at 25 °C for 3 days. On the day of observation, the DAPI dye solution was injected into the back of infected N. benthamiana leaves, and after 3 h of culture under low light, the green fluorescent protein in leaves was observed under confocal scanning microscopy (OLYMPUS, Tokyo, Japan).

4.6. Arabidopsis Thaliana Transformation and Analysis of Flowering Time

The overexpression vector pCAMBIA2300–PoVIN3 was transformed into Arabidopsis thaliana Col–0 via the Agrobacterium tumefaciens-mediated method, and kan was used as a screening marker. Transgenic plants transformed with the empty vector pCAMBIA2300, and wild-type Arabidopsis thaliana (WT) were used as controls. The first generation (T₁) of transgenic seedlings were screened by 50 µg·mL⁻¹ kan and transplanted to the greenhouse for growth. The second generation (T₂) of transgenic seeds was obtained by self–crossing. All transgenic lines were verified by phenotypic observation and PCR analysis.

Flowering time was analyzed by recording the number of days from the sowing of T₂ seeds to the opening of the first flower. In addition, the data are shown as mean values ± SE (standard error). Statistical differences were analyzed by One–Way ANOVA (SPSS 19.0) at the significance level of p < 0.05.

4.7. Establishment of Transient Expression System of Tree Peony

Overnight cultures of Agrobacterium tumefaciens harboring pCAMBIA2300 empty vector and pCAMBIA2300–PoVIN3 were grown at 28 °C in liquid LB medium supplemented with 50 µg·mL⁻¹ kan and 20 µg·mL⁻¹ Rifampin. Agrobacterium tumefaciens cells were centrifuged at 6000 rpm for 10 min at 4 °C and suspended in the infiltration buffer composed of 10 mM MgCl₂, 200 mM acetosyringone, and 10 mM MES/KOH pH 5.6 to a final optical density reached 1.8–2.0 at 600 nm.

Tree peony petals were collected from Paeonia suffruticosa ‘Luoyanghong’ (Petals purplish red, base with inky purple spots) at HO. the outermost whorl of the flowers was discarded, and the petals were divided into two forms: whole petals and disc for subsequent processing. Using a hole punch to remove a point five–centimeter–diameter disc from the center of the petals. The fresh tree peony petals or discs were randomly divided into several equal parts, wrapped with gauze, and immersed in different bacterial
suspension solutions. For vacuum infiltration, tree peony petals or discs were infiltrated under vacuum at 0.07 MPa for 10 min. After releasing the vacuum, the petals and discs were washed four times in deionized water and placed in sterile petri dishes (9 cm in diameter) with 5–10 mL of deionized water, respectively. Subsequently, they were held at 8 °C for 2–3 days, then at 23 °C for 1 day. The tree peony petals and discs were dehydrated for 12 h and then placed in deionized water for rehydration for 12 h. The relative length, relative width, and relative fresh weight of all petals and discs were determined at 3 h intervals. qRT–PCR was used to measure the relative expression levels of the target gene.

Similarly, the flowers of *Paeonia suffruticosa* ‘Luoyanghong’ at BS with 5 cm pedicels were cut off and placed in 40 mL of bacterial suspension solution, and the buds were wrapped with gauze and soaked in 10 mL of bacterial suspension solution. The buds were vacuumized to 0.07 MPa for 30 min and then slowly exhaled. There were 5 flowers per treatment, single flower repeat. The aspirated petals were rinsed with deionized water to remove excess bacterial suspension solution. The pedicels were still immersed in the corresponding bacterial suspension solution and cultured in the dark at 8 °C for 2–3 days, then transferred to 23 °C for 1 day. Then, the fresh-cut tree peony flowers were cultured under normal light and sampled every 7 h. The relative expression levels of target genes and the flowering time of fresh—cut flowers of tree peonies were detected compared with the control.

**4.8. Quantitative Real–Time PCR Analysis of PoVIN3**

Total RNA samples collected from different tissues and petals of developmental stages were used to analyze the spatial and temporal expression patterns of *PoVIN3*, and the expression levels were investigated by quantitative real–time PCR (qRT–PCR). The tubulin-α gene of tree peony was used as the reference gene. Specific primers for quantitative real–time PCR (qRT–PCR) were designed according to the CDS sequence of *PoVIN3* (Table 1). The qRT-PCR was performed on the Light Cycler 96 System (Roche, Germany) using TB Green Premix Ex Taq II (Takara, Japan). Each sample was assayed in four biological replicates, and relative expression levels were analyzed using the $2^{-\Delta\Delta CT}$ method [51]. SPSS 19.0 and Origin 8.5 software was used for statistical analysis. Duncan’s multiple comparison tests were used to determine significant differences between samples. Error bars in all figures represent the standard error from the mean.

**5. Conclusions**

In conclusion, *PoVIN3*, a key gene of the vernalization pathway, was first identified and described in *Paeonia ostii* ‘Fengdan’. The current findings have revealed details about the potential features of *PoVIN3*. *PoVIN3* was the highest at the initial flowering stage of the variant cultivar of *Paeonia ostii* ‘Fengdan,’ the initial decay stage of the normal flowering *Paeonia ostii* ‘Fengdan,’ and the half opening stage of the late flowering *Paeonia suffruticosa* ‘Lianhe.’ Heterologous overexpression of *PoVIN3* in *Arabidopsis thaliana* further clarified its functions. The flowering time was advanced in transgenic *Arabidopsis thaliana* plants hosting *PoVIN3*. *PoVIN3* may act as positive regulatory factor affecting flowering time, thus promoting plant flowering.

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