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Article

Genome-Wide Identification and Functional Characterization of the Dof Family in Dendrobium officinale

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Abstract: The Dof gene family represents a class of plant-specific transcription factors that play crucial regulatory roles in various biological processes, including plant growth, development, and responses to abiotic stress. However, genome-wide identification and functional characterization of the Dof gene family remain unexplored in Dendrobium officinale. In this study, we performed a genome-wide identification and functional analysis of the DoDof gene family. A total of 28 Dof family members were identified and named DoDof1-28 based on genome annotation data. Phylogenetic analysis classified these genes into four major groups (A-D) and further subdivided them into nine subfamilies. Gene structure analysis revealed that most DoDofs lack introns, with no distinct specificity observed among different subfamilies and considerable diversity within the same subfamily. Sequence alignment analysis demonstrated that all DoDof proteins contain a conserved Dof domain consisting of 52 amino acids, which includes a C2-C2 zinc finger motif and a DNAbinding domain. MEME analysis revealed that the conserved motif composition exhibits a certain degree of conservation among DoDof proteins, but significant differences exist across subfamilies. Expression pattern analysis demonstrated that DoDofs have exhibited diverse expression profiles across different developmental stages, tissues, and under abiotic stresses (such as low temperature, salinity, and drought) in D. officinale, suggesting their potential roles in plant development and stress responses. Subcellular localization analysis indicated that DoDof15, DoDof22, and DoDof24 are localized exclusively in the nucleus. Yeast one-hybrid assays revealed that DoDof22 binds to the promoter of the ABA receptor DoPYL9, while DoDof15 and DoDof24 bind to the promoter of the bHLH transcription factor DobHLH68. These results suggest that DoDof proteins may regulate the growth, development, and stress response processes of D. officinale by binding to the promoters of target genes. This study provides critical insights into the functional roles of Dof transcription factors in Orchidaceae family and establishes a theoretical foundation for molecular breeding and stress resistance improvement in D. officinale.

Keywords: Dendrobium officinale; Dof transcription factor; gene expression; yeast one-hybrid

1. Introduction

Transcription factors (TFs) are nuclear-localized proteins that regulate various biological processes, including growth, development, and stress responses, by binding to specific DNA sequences and activating or repressing the expression of downstream target genes at the transcriptional level [1]. Among these TF families, the biological functions of many transcription factor families, such as WRKY, MYB, and NAC, have been extensively studied. The Dof (DNA-binding with one finger) family is a class of plant-specific transcription factors [2] that play crucial roles in plant development and stress responses [3-5]. In contrast to the WRKY and MYB superfamilies, the Dof family has a relatively limited number of members, typically not exceeding 50 in higher plants. For example, 36 and 30 members have been identified in Arabidopsis thaliana and Oryza sativa, respectively [6, 7]. Notably, the Dof transcription factor has not been found in some lower plants, such as Cyanidioschyzon merolae and Thalassiosira pseudonana [8], suggesting that the Dof family may have been acquired during plant evolution and plays an important role in the transition from lower to higher plants.

The Dof proteins have a relatively small molecular weight and consist of four key regions: an Nterminal oligomerization site, a Dof domain, a nuclear localization signal, and a transcriptional activation domain [9]. Notably, the N-terminal region contains a zinc finger domain composed of 52 amino acid residues [10]. This domain contains four cysteine residues that coordinate a Zn²⁺ ion, forming a zinc finger loop [11]. These four cysteines are essential for the stability of the zinc finger loop and play a crucial role in binding metal ions and maintaining the conformation required for DNA binding. Mutations in any of these four cysteine residues can disrupt the zinc finger structure, preventing the protein from binding to the promoter DNA of target genes and thereby abolishing its regulatory activity [12]. The N-terminal zinc finger domain is highly conserved across species, from lower plants such as algae to higher plants like Arabidopsis thaliana and Oryza sativa, indicating that this domain is crucial for the biological functions of Dof proteins. Studies have shown that the zinc finger domain of Dof specifically recognizes the core sequence 5'-(T/A)AAAG-3' in the upstream regions of target genes [13]. Due to the short length of this sequence, many gene promoters contain this binding site [6]. However, not all genes containing this binding site are target genes of Dof transcription factors; most of these sites may be non-functional and cannot be recognized or bound by Dof proteins. The C-terminal region of Dof transcription factors has a transcriptional regulatory function. The amino acid sequence of the C-terminal transcriptional regulatory region is highly variable and lacks conservation, exhibiting functional diversity [14, 15]. This region can act as either a transcriptional activator or repressor, modulating the activity of target gene promoters and thereby regulating transcriptional levels to fulfill its biological roles. For example, Dof1 (MNB1a) acts as a transcriptional activator through its C-terminal region, activating the expression of light-regulated genes and participating in light signal transduction and tissue-specific gene expression in Zea mays. In contrast, Dof2, despite its high sequence similarity with Dof1 in the Dof domain, acts as a transcriptional repressor due to differences in its C-terminal region [16].

Dof proteins are considered dual-functional transcription factors that play crucial regulatory roles in plant dormancy, germination, development, and various physiological processes. Dormancy and germination, which are strictly regulated by the coordinated expression of multiple genes, are complex processes in the plant life cycle [17, 18]. Dof proteins regulate the expression of genes associated with plant dormancy and germination, which is of great significance for analyzing plant life processes [19, 20]. For example, the RGL2-DOF6 complex activates the expression of GATA12 to regulate seed dormancy in Arabidopsis thaliana [21]. Dof proteins are also involved in the growth and development of plant tissues, such as root growth, hypocotyl elongation, and the development and senescence of leaves and floral organs [22-25]. For instance, PbDof9.2 encodes a nuclear-localized protein that regulates the expression of the FT gene by suppressing the activity of the PbTFL1a and PbTFL1b promoters, thereby delaying the flowering time in Pyrus bretschneideri [26]. Notably, Dof transcription factors are involved in the response to abiotic stress in plants, enhancing their stress tolerance. For example, ThDOF1.4 improves salt and osmotic stress tolerance by increasing proline

levels and enhancing reactive oxygen species (ROS) scavenging capacity in Tamarix hispida [27]. The proteins encoded by SICDF1-5 are nuclear-localized and exhibit diverse transcriptional activation in Solanum lycopersicum. These genes are induced under osmotic, salt, heat, and low-temperature stress, and overexpression of SICDF1 or SICDF3 enhances drought and salt tolerance in A. thaliana [28]. These studies demonstrate that Dof proteins play a crucial role in plant stress responses and may provide valuable genetic resources and strategies for plant breeding. The research remains notably limited on the functions of Dof genes. Most Dof proteins have yet to be identified or characterized across various plant species, leading to an incomplete understanding of the regulatory networks and hindering their potential application in plant breeding and related fields.

The Orchidaceae family is one of the largest families of flowering plants. D. officinale, a member of the Dendrobium genus within the Orchidaceae family, is a perennial herbaceous plant and a valuable traditional medicinal material. This study identified and analyzed the Dof family and the composition of conserved motifs, as well as the expression patterns under different developmental stages, tissues, and stress treatments. Additionally, potential DoDof target genes were identified and analyzed, and the binding of DoDof proteins to these genes was examined using yeast one-hybrid analysis. A comprehensive study of the DoDof family and its target genes provides valuable insights into the functions, clarifies future research directions, and offers clues for the application of DoDof transcription factors in orchids.

2. Results

2.1. Identification and Classification of Dof Family in D. officinale

A total of 28 Dof family members were identified and named DoDof1-28 based on the annotation file of the D. officinale genome. These genes are mainly distributed across different scaffolds, with three pairs of genes located on the same scaffold. For example, DoDof2 and DoDof3 are located on scaffold716, and DoDof24 and DoDof25 are located on scaffold1013 (Table S1). The coding sequence (CDS) lengths of the 28 DoDof proteins range from 330 to 1551 bp, with 23 genes shorter than 1000 bp (Table S1). To further analyze the family classification, we conducted a phylogenetic analysis of the 28 DoDof proteins from D. officinale and the 36 AtDof proteins from A. thaliana using the neighbor-joining method (Table S2). These Dof proteins can be classified into four groups (A-D), which were further subdivided into nine subfamilies: A, B1, B2, C1, C2.1, C2.2, C3, D1, and D2. As with other plants, group A contains only one subfamily, while groups B, C, and D have at least two subfamilies. Group C has the most subfamilies, including C1, C2.1, C2.2, and C3 (Figure 1). Group A has the fewest members, only containing DoDof20, while Group C has the most members, with 12 members (Figure 1). Gene structure analysis revealed that the DoDofs exhibit significant structural diversity. For instance, within the C3 subfamily, one member contains one intron, another contains two introns, and yet another lacks an intron (Figure 2). Most DoDofs lack introns, with all members of subfamilies A, B2, C2.2, and D2 devoid of introns (Figure 2). Among all genes, 15 genes have no introns, 10 genes have one intron, and only three genes have two introns (Figure 2). These findings indicate that the gene structure of DoDofs is not conserved.

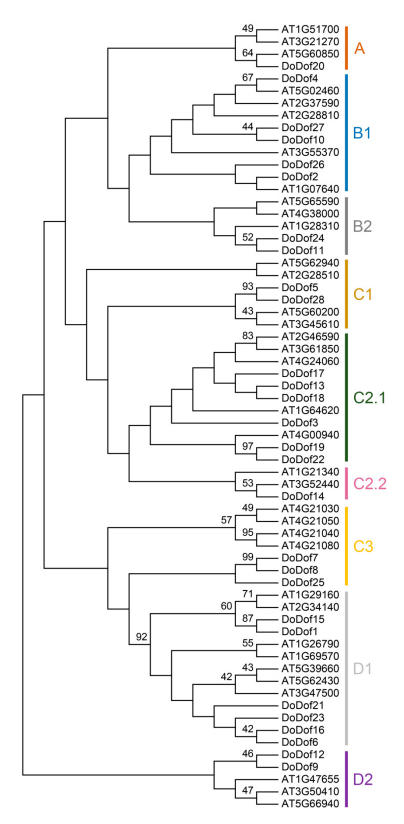


Figure 1. Phylogenetic tree of Dof proteins in *D. officinale* and *A. thaliana*. The phylogenetic tree was constructed using the neighbor-joining (NJ) method with 1000 bootstrap replications based on full-length protein sequences from 28 DoDofs and 36 AtDofs by MEGA 7.0.



Figure 2. Gene structure analysis of *DoDof* genes in *D. officinale*. The intron and exon information of *DoDof* genes were obtained by whole genome sequencing GFF file of *D. officinale*. *DoDof* genes structure were mapped using GSDS 2.0 software.

2.2. Sequence Alignment and Gene Structure Analysis of DoDof Proteins

Sequence alignment analysis of DoDof proteins revealed that contain a conserved domain composed of 52 amino acids, which includes a conserved CX2CX21CX2C (C2-C2) motif and a DNA-binding domain (Figure 3). Within the C2-C2 zinc finger domain, there are four Cys residues that covalently bind to Zn²+ to form a single zinc finger C2C2 domain, which plays a crucial role in DNA binding (Figure 3). The C2C2 domain cannot function when Cys is replaced by His [12]. Adjacent to the zinc finger domain is a DNA-binding motif, which is present in most DoDof proteins and contains two conserved amino acids, Tyr (Y) and Trp (W). However, the Tyr (Y) is mutated to His (H) in DoDof7 and DoDof8 (Figure 3). The MEME online website was used to predict the conserved motifs and further analyze the motif characteristics of DoDof proteins. Interestingly, the motif composition of DoDof proteins shows a certain degree of conservation in that all DoDofs containing motif1 (Figure 4). Some motifs are unique to specific subfamilies, for example, motif2 and motif7 are only present in the C3 subfamily, while motif3, motif4, motif6, and motif8 are only found in the D1 subfamily. These motifs may play important roles in the functioning of genes, leading to the diversity of functions within the Dof gene family.

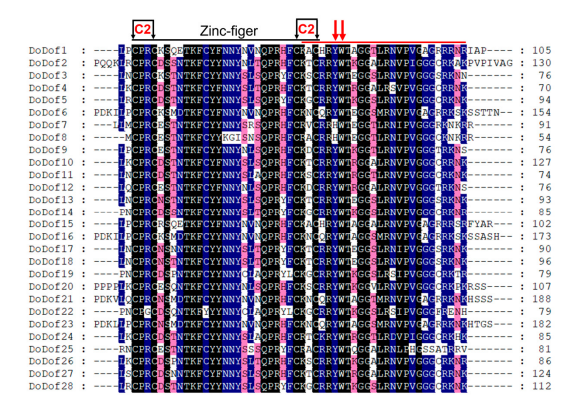


Figure 3. Multiple sequence alignment of DoDof proteins in *D. officinale*. Dof domain were aligned using ClustalX 2.1 and export the file in FASTA format, amino acid conserved residues were displayed using Genedoc 3.0. Diferent colors indicate diferent levels of conservation in residues, black, blue and pink indicate 100%, 75%, and 50% identical residues, respectively. Black arrows indicate cysteine residues that are likely contributing to the formation of a C2–C2 Zn finger. A red bar marks a region outside the Zn finger that plays a role in DNA binding. Red arrows highlight conserved aromatic amino acid residues that are essential for DNA interaction.

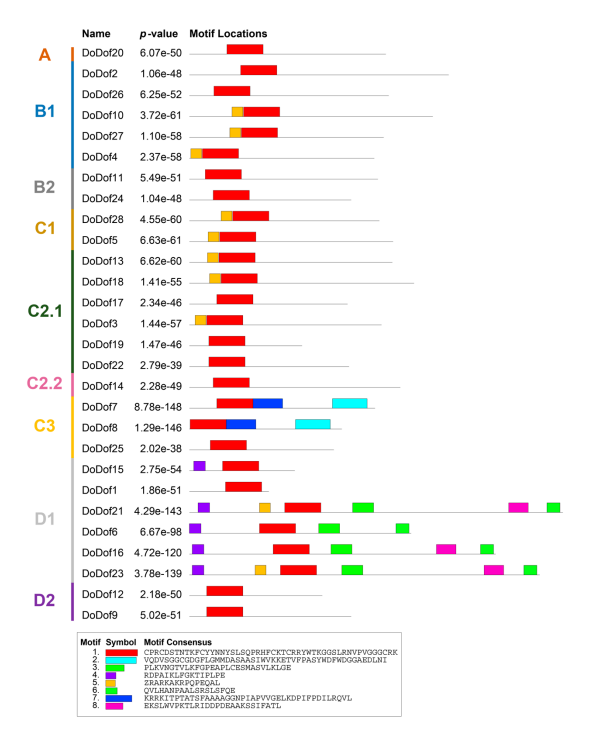


Figure 4. Conserved motif analysis of DoDof protein in *D. officinale*. DoDof protein sequence were imported into MEME online website for conserved domain analysis. The different subfamilies of the DoDof genes are shown in different colors on the left. The sequence of conserved motifs is presented at the bottom.

2.3. Expression Patterns of DoDof Genes in Different Developmental Stages and Tissues

We analyzed the expression patterns of DoDofs in different developmental stages and tissues, including protocorm-like bodies (PLBs), shoots derived from PLB, plantlets, and the rhizomes, stems, and leaves of adult plants. The qPCR results revealed that, except for DoDof25, which was not detected, all other genes were expressed, with DoDof4 and DoDof27 having relatively high expression levels in plantlets (Figure 5). The expression levels of DoDof3, DoDof8, DoDof12, DoDof18, DoDof19, DoDof22, and DoDof24 were highest in PLBs and significantly decreased as PLBs developed into shoots (Figure 5). The expression levels of DoDof4, DoDof5, DoDof7, DoDof15, DoDof23, and DoDof27 were highest in plantlets and relatively low in PLBs and shoots, suggesting

that these genes may play important roles in plantlets development (Figure 5). Notably, the expression levels of DoDof6, DoDof9, DoDof10, DoDof11, DoDof14, DoDof16, DoDof20, DoDof21, and DoDof26 were highest in leaves, implying that these genes are involved in leaf development of D. officinale (Figure 5). The expression levels of DoDof2, DoDof17, and DoDof28 were highest in stems and lowest in leaves (Figure 5). These results indicate that the expression patterns of DoDofs are diverse and they may play important roles in the organ development of D. officinale.

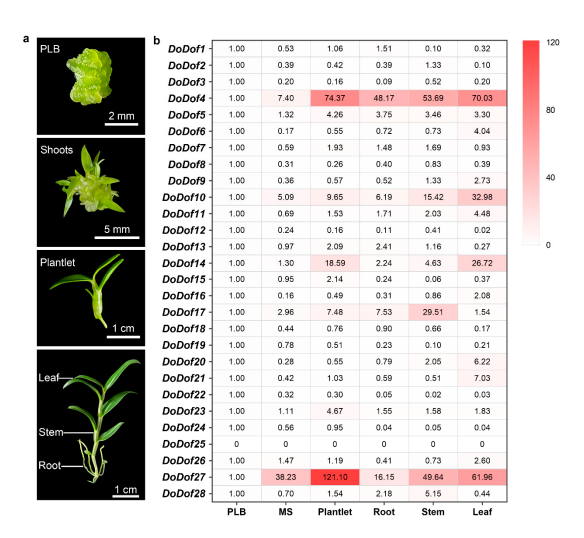


Figure 5. *DoDof* genes expression analysis of *D. officinale* in different development stages and tissues. (a) The three developmental stages of *D. officinale* are protocorm-like body (PLB), shoots (MS) and plantlet. The three organs of the adult plant (about 10cm) are root, stem and leaf. (b) Heatmap of expression pattern of 28 *DoDof* genes by qRT-PCR. Values are the mean values of Ct for three technical repeats.

2.4. Expression Patterns of DoDof Genes Under Different Stress Treatments

Dof transcription factors are widely involved in stress response processes in plants. The expression patterns of DoDof genes were analyzed under salt, low temperature, 15% PEG-6000, and mannitol treatments. All 28 DoDofs showed altered expression under low-temperature stress, with the expression levels of 24 genes significantly decreased, while the expression of DoDof7, DoDof8, DoDof9, and DoDof23 significantly increased (Figure 6). Under salt treatment, the expression of DoDof1, DoDof9, DoDof23, and DoDof27 was significantly upregulated, while those of DoDof3, DoDof4, DoDof6, DoDof8, DoDof11, DoDof13, DoDof14, DoDof17, DoDof18, DoDof22, DoDof24, DoDof25, and DoDof28 were significantly downregulated (Figure 6). Notably, the expression of DoDof25 was significantly reduced under salt stress and was undetectable in all other treatments (Figure 6). Although both PEG (osmotic stress) and mannitol (dehydration stress) are used to

simulate drought stress treatments, DoDofs exhibit distinct expression trends under these two treatments, with most of the genes showing opposite expression patterns. Compared to untreated plants, the expression of DoDof2, DoDof6, DoDof10, DoDof12, DoDof13, DoDof16, DoDof17, DoDof19, DoDof20, DoDof21 and DoDof28 were significantly upregulated or remained unchanged under PEG treatment, while these genes were significantly downregulated under mannitol treatment (Figure 6). It is noteworthy that DoDof25 was significantly downregulated under salt treatment and undetected under low temperature, PEG and mannitol treatments (Figure 6). These results indicate that DoDofs play a significant role in the response to stress and may primarily act as negative regulators in D. officinale.

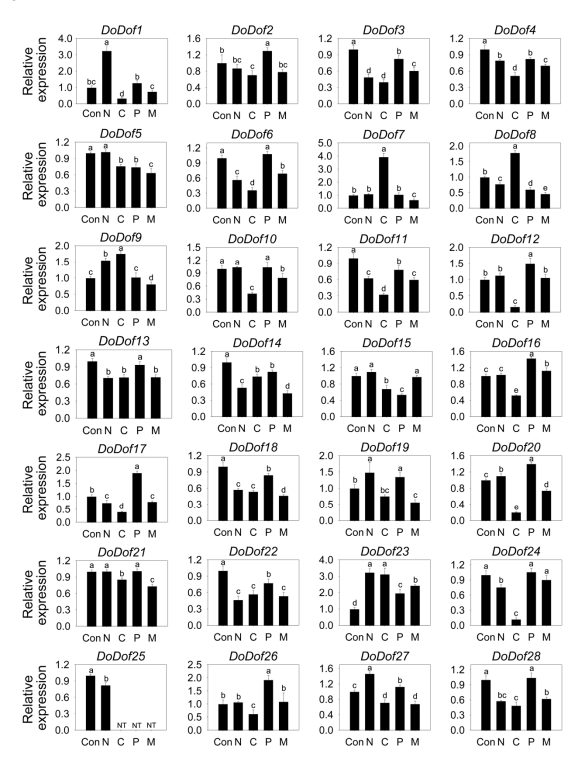


Figure 6. Expression pattern of *DoDof* genes in *D. officinale* under stress treatment. Seedlings untreated, 250 mM NaCl treatment, 4 °C treatment, 15% polyethylene glycol-6000 treatment and 300mM Mannitol treatment are denoted by the letters Con, N, C, P and M, respectively. Each data bar represents the mean \pm standard deviation (SD) of three replicates. ND, not detected. Different lowercase letters indicate statistically significant differences (p < 0.05), while the same letters indicate no significant difference.

2.5. Subcellular Localization Analysis of DoDof15, DoDof22, and DoDof24

Transcription factors perform their transcriptional regulatory roles within the nucleus [29]. Subcellular localization predictions of DoDof proteins also showed that these proteins are localized in the cell nucleus using WoLF PSORT online software (Table S3). To further analyze the localization of DoDof proteins, we selected DoDof20 from Group A, DoDof24 from Group B, DoDof22 from Group C, and DoDof15 from Group D for subsequent analysis. However, DoDof20 could not be successfully cloned, so DoDof15, DoDof22, and DoDof24 were selected for further investigation. The DoDof-YFP transient expression vectors were constructed using yellow fluorescent protein (YFP) as the fluorescent marker. When these vectors were transiently expressed in protoplasts of A. thaliana, the yellow fluorescence of DoDof15-YFP, DoDof22-YFP, and DoDof24-YFP primarily appeared in the nuclear region and overlapped with the red fluorescence of the nuclear localization marker NLS-mCherry, indicating that DoDof15, DoDof22, and DoDof24 are localized in the nucleus. (Figure 7a). As a positive control, the yellow fluorescence of 35S::YFP was observed in the cell membrane, cytoplasm, and nucleus, showing a lack of specific localization (Figure 7b).

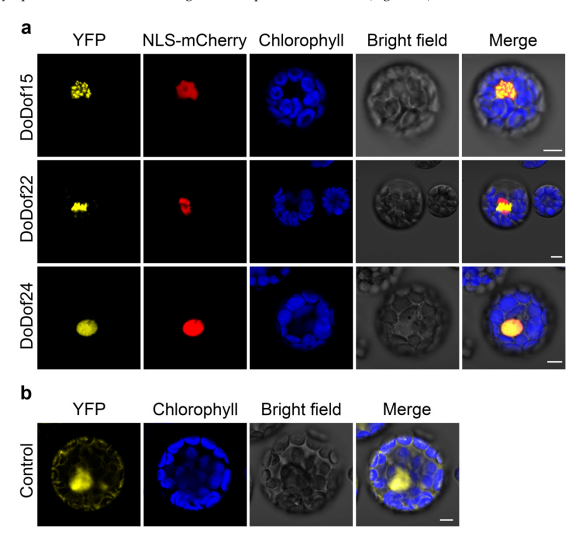


Figure 7. Subcellular localization of three DoDof proteins in *Arabidopsis* protoplasts. (a) Localization of DoDof15-YFP, DoDof22-YFP, and DoDof24-YFP fusion proteins with the nuclear marker NLS-mCherry. (b) The control YFP was co-transfected with empty YFP and NLS-mCherry plasmids. Scale bars = $5 \mu m$.

2.6. Binding Analysis of Candidate Target Genes for DoDof15, DoDof22, and DoDof24

Dof transcription factors bind to the core sequence 5'-(T/A)AAAG-3' in the promoters of target genes, thereby regulating gene expression and achieving biological functions such as regulating plant growth and development, and responding to abiotic stress [30-32]. To date, the target genes regulated by DoDof transcription factors remain unidentified. The 1000 bp upstream sequences of the start codons of all genes were obtained through whole-genome data in D. officinale. Using the core sequence 5'-(T/A)AAG-3' as a probe, all promoters were scanned to identify genes containing the core sequence 5'-(T/A)AAAG-3' in their promoter regions. A total of 27,164 genes were found to have at least one 5'-(T/A)AAAG-3' core sequence in their promoter regions. Among these, 23,491 genes contain two or more 5'-(T/A)AAAG-3' core sequences, and 18,734 genes have at least three 5'-(T/A)AAAG-3' core sequences in their promoter sequences (Table S4). To improve the accuracy of identifying candidate target genes, only those containing at least three 5'-(T/A)AAAG-3' core sequences were selected for further analysis. The bHLH transcription factor family exists in all eukaryotes and is involved in key processes of plant growth, development, and interaction with the environment [33]. Among these candidate target genes, the promoter of DobHLH68 (Dca022885) contains eleven 5'-(T/A)AAAG-3' core sequences (Figure 8a). Abscisic acid (ABA) is one of the most important plant hormones regulating stress responses, primarily exerting its precise regulatory effects through the ABA signaling pathway [33, 34]. The ABA receptor protein PYL is a critical component of the ABA signaling pathway [35] and plays a significant role in stress responses [36-38]. Among the candidate target genes identified, the promoter region of the ABA receptor protein gene DoPYL9 (Dca025790) contains ten 5'-(T/A)AAAG-3' core sequences (Figure 8a). We analyzed the binding of DoDof15, DoDof22, and DoDof24 to the promoters of DobHLH68 and DoPYL9 in a yeast system, respectively. The negative control pGADT7 + proDobHLH68 and other combinations could grow normally in SD/-T/-L and SD/-T/-L/-H media. The growth of pGADT7 + proDobHLH68 was significantly inhibited, and even when DoDof22 was present, the growth of yeast was still inhibited in media containing 60 mM 3-amino-1, 2, 4-triazole (3-AT). However, yeast growth was better when DoDof15 and DoDof24 were present (Figure 8b and Figure S1). These results suggest that DoDof15 and DoDof24 can bind to the promoter of DobHLH68 (proDobHLH68). The growth of pGADT7 + proDoPYL9 was significantly inhibited in media containing 140 mM 3-AT. In the presence of DoDof22, yeast growth was significantly improved, while no significant change was observed in the presence of DoDof15 and DoDof24 (Figure 8b and Figure S1), which suggests that DoDof22 binds to the promoter of DoPYL9 (proDoPYL9).

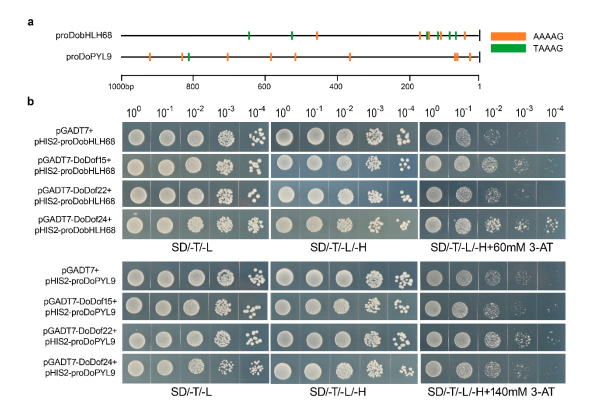


Figure 8. Validation of three DoDof proteins binding core sequence 5'-(T/A)AAAG-3'. (a) The number of core elements contained in the *DobHLH68* and *DoPYL9* promoters 1000 bp. (b) Yeast one-hybrid verification of DoDof15, DoDof22 and DoDof24 binding to *DobHLH68* and *DoPYL9* promoters, respectively. SD/-T/-L: a yeast culture medium without tryptophan and leucine; SD/-T/-L/-H: a yeast culture medium lacking tryptophan, leucine, and histidine; SD/-T/-L/-H + 3-AT: a yeast culture medium excluding tryptophan, leucine, and histidine, supplemented with 3-AT at specific concentrations (60 mmol/L for DobHLH68 and 140 mmol/L for DoPYL9).

3. Discussion

3.1. Dof Transcription Factors Are Ubiquitous in Plants and Possess a Conserved Zinc Finger Domain

The first Dof protein, ZmDof1, was identified in maize, and its binding domain includes a C2C2 zinc finger motif [39]. Since this discovery, an increasing number of Dof family members have been identified across diverse plant species, ranging from single-celled algae, and mosses to flowering plants, such as A. thaliana [7], Glycine max [40], Brassica rapa L. ssp. pekinensis [41], Cucumis sativus [42], and Betula platyphylla [43]. Notably, only one Dof transcription factor has been identified in Chlamydomonas reinhardtii, Volvox carteri, and Micromonas pusilla [8, 44]. In contrast, more complex plant species exhibit greater diversity, with the moss Physcomitrella patens containing 19 Dof genes and the gymnosperm Pinus taeda harboring 10 Dof genes [8]. Extensive studies have identified and characterized numerous Dof genes in angiosperms [9]. In the case of D. officinale, a member of the orchid family, 28 DoDof genes have been characterized. These findings suggest that Dof transcription factors are widely present in green plants and may play roles in functions specific to these plants. The N-terminus of these Dof proteins contains a conserved zinc finger domain, which has four cysteines. These four cysteines bind to a Zn2+ to form a zinc finger structure, which is necessary for the specific recognition of the core sequence 5'-(T/A)AAAG-3' upstream of target genes. Among the 28 DoDof genes, the N-terminus of their protein sequences all contain a zinc finger domain (Figure 3). The Dof domain is not only a DNA-binding domain but also exhibits proteinprotein interaction activity. Dof proteins can interact with other transcription factors, such as MYB, bZIP, and WRKY [9]. For example, the Dof protein SAD interacts with the R2R3MYB protein (GAMYB) to activate the transcription of endosperm-specific genes in Hordeum vulgare cv. Bomi

[45]. These results indicate that the zinc finger domain is conserved in Dof transcription factors and plays a key role in the biological functions of this gene family.

3.2. Dof Transcription Factors Regulate Leaf Development in Plant

Leaves are one of the important organs in plants and serve as the primary sites for photosynthesis [46, 47]. Stomata are distributed on the underside of leaves, regulating the exchange of gases and water vapor [48, 49]. Dof transcription factors play a crucial role in regulating stomatal development. SCAP1 encodes a Dof transcription factor, and the scap1 mutant exhibits irregular guard cells, lacking the ability to control stomatal aperture, including CO2-induced stomatal closure and light-induced stomatal opening. For example, SCAP1 is a key regulator directly regulating stomatal function in A. thaliana [50]. The promoters of AtDof2.4 and AtDof5.8 exhibit specific activity in leaf primordia [51], suggesting their involvement in leaf initiation in A. thaliana. The activation tagging mutant Dof5.1-D exhibits an upward-curling leaf phenotype, while overexpression of the DNA-binding domain of Dof5.1 results in a downward-curling leaf phenotype, indicating that Dof5.1 regulates leaf axial patterning in A. thaliana [52]. Furthermore, Dof5.8 encodes a Dof transcription factor, and its overexpression suppresses the formation of higher-order veins in cotyledons and leaves in A. thaliana [53]. Studies have demonstrated that Dof transcription factors are not only involved in leaf development but also play a role in leaf senescence. For instance, the Dof transcription factor CDF4 promotes leaf senescence by upregulating the transcription of ABA biosynthesis genes nced2,3, thereby increasing endogenous ABA levels. Additionally, CDF4 represses the expression of the CAT2 gene, inhibiting the clearance of H₂O₂ and further accelerating leaf senescence [54]. Dof2.1 is a JA-induced gene that is closely related to leaf senescence. Overexpression of Dof2.1 promote leaf senescence in A. thaliana. Mechanistically, Dof2.1 activates the MYC2 promoter and enhances MYC2 expression to drive leaf senescence [55]. Notably, the high expression levels of DoDof6, DoDof9, DoDof10, DoDof11, DoDof14, DoDof16, DoDof20, DoDof21, and DoDof26 in the leaves suggest that these genes may be involved in leaf development or senescence and play significant roles in the regulation of leaf function in D. officinale.

3.3. Dof Transcription Factors Respond to Abiotic Stress in Plants

Dof transcription factors not only regulate in plant development but also play pivotal roles in plant stress responses to adverse conditions. Extensive research has demonstrated that Dof transcription factors can be induced by various abiotic stresses such as salt, drought, and low temperatures. For instance, most of StDof genes are upregulated under drought and salinity conditions in potatoes [14]. Two cold-induced genes, OsDof1 and OsDof9, have been identified in Oryza sativa L., with OsDof1 contributing to enhanced cold stress tolerance [56]. The Dof gene CDF3 plays a significant role of abiotic stress responses in A. thaliana. It is strongly induced by drought, extreme temperatures, and abscisic acid (ABA) treatment. Mutants of CDF3 exhibit heightened sensitivity to drought and low-temperature stress, whereas its overexpression improves tolerance to drought, cold, and osmotic stress in transgenic plants [57]. Research has revealed that Dof transcription factors are involved in stress response signaling pathways, such as ABA and ROS pathways, thereby enhancing plant stress tolerance. These transcription factors promote the accumulation of ABA by activating the expression of ABA biosynthesis genes and suppressing the expression of ABA catabolism genes. For example, the Dof transcription factor CDF4 increases endogenous ABA levels by upregulating the transcription of the ABA biosynthesis gene nced2,3 [54]. Similarly, DAG1 directly represses the expression of the ABA catabolism gene CYP707A2, effectively increasing ABA content [58]. These findings suggest that Dof transcription factors likely promote the accumulation of ABA, activating the ABA signaling pathway and enabling plants to adapt to adverse environments such as drought, high salinity, and low temperatures. Under stress conditions, a large amount of ROS is accumulated in plants. While excessive ROS can damage cellular components, they also serve as vital signaling molecules [59]. Dof transcription factors have two functions in regulating ROS: promoting ROS accumulation and clearing ROS. For example, Dof transcription factors inhibit

the expression of the CAT2 gene, thereby inhibiting the clearance of H2O2 [54]. ROS plays a crucial role in the perception of abiotic and biotic stresses in plants, integrating different environmental signals, and activating stress response networks, which helps to establish defense mechanisms and stress resistance in plants [59]. The Dof transcription factor RsCDF3 directly binds to the promoters of the key plasma membrane-bound enzymes RsRbohA and RsRbohC that produce ROS in Raphanus sativus L., inhibiting their transcription, maintaining the homeostasis of ROS in cells, and thereby enhancing cold resistance [60]. In this study, the expression levels of all 28 DoDof genes were altered under low-temperature treatment, with four genes significantly upregulated and the rest downregulated. The expression levels of DoDof1, DoDof9, DoDof23, and DoDof27 were significantly increased under salt stress. Similarly, some DoDofs were induced by PEG-6000 and mannitol (Figure 6). These studies suggest that Dof transcription factors may be involved in the stress response of D. officinale, indicating a conserved function in abiotic stress responses across plant species.

3.4. Dof Transcription Factors Show Diversity in Recognizing and Binding to Core Sequences

The single zinc finger domain of Dof transcription factors can specifically recognize and bind to the 5'-(T/A)AAAG-3' sequence in the promoters of target genes, thereby regulating their expression and fulfilling the functions of Dof transcription factors. For instance, AtDof5.1 recognizes and binds to the TAAAGT motif, thus regulating the transcription of target genes in A. thaliana [52]. Dof transcription factors generally have a high affinity for the AAAG promoter sequence, and binding is weakened or prevented when AAAG is mutated [61, 62]. However, this does not imply that all Dof transcription factors can bind to the 5'-(T/A)AAAG-3' sequence; the sequences flanking this binding site may influence the binding function [10]. Although the promoters of DobHLH68 and DoPYL9 both contain the 5'-(T/A)AAAG-3' sequence, the sequences at the ends of this binding site are not identical, which may contribute to the binding specificity of Dof transcription factors. Additionally, the C-terminal region of Dof proteins is crucial for protein-protein interactions and exhibits specificity. Dof transcription factors are likely to interact with other transcription factors to achieve the necessary promoter specificity. For example, the Dof transcription factor BPBF interacts with the MYB transcription factor GAMYB to activate the promoter of target genes, regulating of seed development and germination [63]. In this study, we found that the promoters of DobHLH68 and DoPYL9 both contain abundant 5'-(T/A)AAAG-3' sequences. However, among the three selected DoDof genes, only DoDof15 and DoDof24 bound to the promoter of DobHLH68 in yeast, while DoDof22 bound to the promoter of DoPYL9 (Figure 8b). These research results indicate that the binding of Dof proteins to the 5'-(T/A)AAAG-3' motif may be affected by the flanking sequences the motif and interacting proteins, and their regulatory roles may be a complex and diverse process on target genes. This indicates that more in-depth research is needed to clarify the biological functions of Dof in plant development and stress responses.

4. Materials and Methods

4.1. Plant Materials and Growth Conditions

The D. officinale plants used in this study were cultivated at the South China Botanical Garden, Chinese Academy of Sciences, Guangzhou, China. Additionally, tissue-cultured D. officinale seedlings are preserved in the laboratory. The experimental materials included protocorm-like bodies (PLBs), shoots (MS), plantlets (approximately 1.5 cm tall), and adult plants (about 10 cm tall), all of which were obtained under controlled tissue culture conditions. The culture room was maintained at 25°C with a 12-hour light/12-hour dark photoperiod and 50 µmol m⁻² s⁻¹ light intensity. The basic culture medium was 1/2 MS medium [64], supplemented with 0.1% activated charcoal, 2% sucrose, and 0.6% agar (pH 5.4). To investigate stress responses, the plantlets (about 1 cm tall) were subjected to various abiotic stress treatments according to our previously established protocols [65]. The treatment conditions were as follows: 250 mM NaCl (Guangzhou Chemical Reagent Factory, Guangzhou, China) to simulate salt stress; 15% polyethylene glycol-6000 (PEG-6000) (Solarbio

Science & Technology Co., Ltd., Beijing, China) to simulate osmotic stress; 300 mM mannitol (Macklin Biochemical Technology Co., Ltd., Shanghai, China) to simulate drought stress; and 4°C low-temperature stress. Each stress treatment lasted 6 hours, while the control group seedlings were cultured under normal conditions. All experiments were repeated three times. Samples were collected and immediately frozen in liquid nitrogen for 15 minutes after treatment, followed by RNA extraction for subsequent analysis. The PLBs, MS, plantlets, and roots, stems, and leaves from adult D. officinale plants were processed similarly for qRT-PCR analysis.

For subcellular localization experiments, an appropriate quantity of Columbia-type A. thaliana seeds was placed in centrifuge tubes with distilled water and incubated in the dark at 4°C for 2-3 days. Subsequently, the seeds were sown on a substrate mixture of nutrient soil and perlite (2:1, v/v) and transferred to a growth chamber maintained at 22 ± 2 °C with a 16-hour light/8-hour dark photoperiod, light intensity of 80 μ mol/m²/s, and relative humidity of 80%. When the Arabidopsis reached 4-6 weeks of age, protoplasts were isolated from leaves using previously described methods [66].

4.2. Identification of DoDof Transcription Factors and Construction of Phylogenetic Tree

The genomic data of D. officinale were downloaded from the National Center for Biotechnology Information (NCBI) Genomic Database (https://www.ncbi.nlm.nih.gov/) under the project PRJNA262478 [67]. The D. officinale genome database was searched using the keyword "Dof" to identify genes containing this term in their annotations. BlastX alignment was performed on NCBI, along with phylogenetic analysis, to remove duplicate or incorrect DoDof sequences. Ultimately, 28 Dof proteins were identified from the D. officinale genome. The amino acid sequences of the DoDof proteins were then submitted to ProtParam (https://web.expasy.org/protparam/) [68] to calculate their physicochemical properties and to WoLF PSORT (https://wolfpsort.hgc.jp/) for subcellular localization prediction.

Multiple sequence alignment of the full-length amino acid sequences of DoDof proteins was performed using Clustal X 2.0 software [69]. Based on the alignment results, a phylogenetic tree was constructed using the neighbor-joining (NJ) method [70] in MEGA 7 software [71]. The authenticity of the tree was tested by performing 1000 bootstrap replications, resulting in the construction of a phylogenetic tree of Dof proteins from D. officinale and A. thaliana (Table S2).

4.3. Bioinformatics Analysis of DoDof Proteins

Based on the FASTA alignment file obtained from Clustal X 2.0 software, multiple sequence alignment of the full-length amino acid sequences of DoDof proteins was performed using GENEDOC 3.0 software [72] with default parameters. The phylogenetic tree of the DoDof family proteins was constructed using the neighbor-joining method based on the multiple sequence alignment, and a Newick Tree file was exported. The gene annotation information of the DoDof family members was extracted from the GFF3/GTF file of D. officinale and submitted to the GSDS2.0 online website (https://gsds.gao-lab.org/) for gene structure visualization [73]. The DoDof proteins amino acid sequences were submitted to the MEME online website (https://memesuite.org/meme/tools/meme) for conserved motif analysis, with the number of conserved motifs set to 8 and other parameters maintained at default values. The conserved motif information was subsequently visualized [74].

4.4. RNA Extraction, cDNA Synthesis, and qRT-PCR Analysis

Total RNA was extracted from PLBs, shoots, plantlets, roots/stems/leaves of adult plants, and stress-treated plantlets using the HiPure HP Plant RNA Mini Kit R4165-02 (Magen Biotechnology Co., Ltd., Guangzhou, China). RNA purity and concentration were determined using the Nanodrop 2000C spectrophotometer (Thermo Scientific, Wilmington, DE, USA). First-strand cDNA synthesis was performed using the All-in-One First-Strand Synthesis MasterMix MD80101 (Magen

Biotechnology Co., Ltd., Guangzhou, China), and the cDNA was quantified and stored at -20°C for subsequent analysis. The qRT-PCR primers were designed (Table S5) and synthesized by Sangon Biotech Co., Ltd. (Shanghai, China). Using ACTIN (NCBI accession number: JX294908) as the reference gene [75], the relative expression levels of DoDof genes were determined across different developmental stages, tissues, and stress treatment samples in D. officinale. The qRT-PCR reactions were performed according to the protocol of Magen Universal SYBR qPCR Mix MD70101 (Magen Biotechnology Co., Ltd., Guangzhou, China) on the Roche LightCycler 480 system (Roche, Basel, Switzerland). The thermal cycling conditions consisted of an initial denaturation at 95°C for 60 s, followed by 40 cycles of 95°C for 10 s and 60°C for 10 s. The gene relative expression levels were calculated using the $2-\Delta\Delta$ Ct method [76] to determine the relative differences in gene expression between samples. The experimental data were presented as the average of three biological replicates, and graphs were created using Sigmaplot 15.0 software (Systat Software Inc., San Jose, California, USA), followed by significance analysis.

4.5. Subcellular Localization Analysis

To investigate the subcellular localization of DoDof proteins, primers (Table S6) were designed to amplify the full-length coding sequences of DoDof15, DoDof22, and DoDof24 genes. The pSAT6-EYFP-N1 vector [77] was digested with EcoRI, and the digested product was purified. The PCR-amplified target genes were ligated into the linearized vector using homologous recombination. The recombinant plasmids were transformed into Escherichia coli DH5α competent cells (Weidi Biotechnology Co., Shanghai, China). After transformation, single colonies were selected for colony PCR screening, and positive clones were subsequently sequenced. The correctly sequenced strains were cultured in a liquid medium for amplification. The recombinant plasmids (DoDof15-YFP, DoDof22-YFP, and DoDof24-YFP) were extracted and co-transformed with the nuclear localization marker NLS-mCherry into mesophyll protoplasts using polyethylene glycol (PEG)-mediated transformation [66]. The transformed protoplasts were incubated in darkness for 8-10 hours. Fluorescence expression was observed using a Leica TCS SP8 STED 3× super-resolution confocal microscope (Leica Camera AG., Solms, Germany). The excitation and emission wavelengths were 514 nm and 527 nm for YFP, and 587 nm and 610 nm for the nuclear marker mCherry, respectively.

4.6. Analysis of the Core Sequence 5'-(T/A)AAAG-3' in the Promoter of D. officinale

To evaluate the distribution characteristics and potential regulatory roles of Dof recognition sequences in the genome of D. officinale, the upstream 1000 bp sequences of all genes were extracted as promoter regions using Tbtools software [78]. Subsequently, the Dof recognition sequences 5'-(T/A)AAAG-3' was identified within the extracted sequences using regular expression matching, specifically targeting sequences beginning with either T or A followed by AAAG. The occurrence frequency of this core sequences in the promoters of each gene was counted, and its distribution pattern was recorded for subsequent analysis.

4.7. Promoter Self-Activation Detection and Yeast One-Hybrid Assay

For yeast one-hybrid assays, specific primers were designed (Table S7) to amplify the promoter fragments of DobHLH68 and DoPYL9 (proDobHLH68 and proDoPYL9), which were subsequently cloned into the pHIS2 vector to construct recombinant plasmids pHIS2-proDobHLH68 and pHIS2-proDoPYL9. These plasmids were transformed into Saccharomyces cerevisiae strain Y187 yeast competent cells (Weidi Biotechnology Co., Shanghai, China) following the manufacturer's protocol. Well-growing colonies were selected, dissolved in ddH2O, and serially diluted to 10-1, 10-2, 10-3, 10-4, and 10-5. A 10µl of the diluted cell suspension was spread onto SD/-T/-H medium containing 3-AT at concentrations of 0, 20, 40, 60, 80, 100, 120, 140, 160, and 180 mM (Coolaber Biotechnology Co., Ltd., Beijing, China) to assess the resistance of pHIS2-proDobHLH68 and pHIS2-proDoPYL9 to 3-AT. The

most suitable concentration for 3-AT during the yeast one-hybrid assay was determined based on optimal growth (Figure S1).

The coding sequences of DoDof15, DoDof22, and DoDof24 were amplified and inserted into the pGADT7 vector to generate effector constructs pGADT7-DoDof15, pGADT7-DoDof22, and pGADT7-DoDof24. The plasmids pHIS2-proDobHLH68 and pHIS2-proDoPYL9 were co-transformed with three effector vectors into Y187 yeast competent cells using the lithium acetate method (Weidi Biotechnology Co., Shanghai, China). Transformant strains were initially selected on SD/-Trp/-Leu (SD/-T/-L) medium, and positive clones were transferred onto SD/-Trp/-Leu/-His (SD/-T/-L)-H) medium containing 3-AT. Plates were incubated at 29°C for 3-4 days, and yeast growth was monitored and recorded. Negative controls consisted of yeast co-transformed with pHIS2-proDobHLH68 and pGADT7, or pHIS2-proDoPYL9 and pGADT7.

4.8. Statistical Analysis

The qRT-PCR experimental data were analyzed using SigmaPlot 15.0 software, with results presented as means \pm standard deviation (SD) of three biological replicates. Statistical significance was determined by one-way analysis of variance (ANOVA) followed by Duncan's multiple range test between pairs of groups. Two groups with significant differences (p < 0.05) are denoted by different letters, while two groups with non-significant differences are indicated by the same letter.

5. Conclusions

We identified Dof transcription factors in D. officinale for the first time. This is also the first exploration of the conserved domains, expression patterns, and potential biological functions of Dof transcription factors in orchids. The genome contains 28 DoDof genes in D. officinale, all of which possess the C2-C2 conserved domain characteristic of the Dof family. These genes exhibit diversity in both gene structure and expression patterns. The expression pattern analysis suggests that the DoDof genes are likely involved in plant development and response to stress conditions. We identified potential target genes of Dof transcription factors, discovering that the promoters of 13,811 genes contain three or more core sequences of 5'-(T/A)AAAG-3'. Yeast one-hybrid assays indicate that there may be diversity in the recognition and binding of these core sequences. We found that DoDof22 binds to the promoter of the ABA receptor protein DoPYL9, suggesting that DoDof22 is involved in the ABA signaling pathway, regulating the stress response in D. officinale. Furthermore, DoDof15 and DoDof24 could bind to the promoter of DobHLH68 in yeast. Research indicates that the bHLH transcription factors primarily participate in the secondary metabolism of plants, which implies that DoDof may regulate the expression of DobHLH68, thereby influencing the synthesis of secondary metabolites and enhancing the plant's stress resistance.

Supplementary Materials: The following supporting information can be downloaded at the website of this paper posted on Preprints.org.

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