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Overexpression of MdZAT5, an C2H2-Type Zinc Finger Protein, Regulates Anthocyanin Accumulation and Salt Stress Response in Apple Calli and Arabidopsis

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Abstract: Zinc finger proteins are widely involved and play an important role in plant growth and abiotic stress. In this research, MdZAT5, a gene encoding C2H2-type zinc finger protein, was cloned and investigated. The MdZAT5 was highly expressed in flower tissues by qRT-PCR analyses and GUS staining. Promoter analysis showed that MdZAT5 contained multiple response elements, and the expression levels of MdZAT5 were induced by various abiotic stress treatments. Overexpression of MdZAT5 in apple calli positively regulated anthocyanin accumulation by activating the expressions of anthocyanin biosynthesis-related genes. Overexpression of MdZAT5 in Arabidopsis also enhanced the accumulation of anthocyanin. In addition, MdZAT5 increased the sensitivity to salt stress in apple calli. Ectopic expression of MdZAT5 in Arabidopsis reduced the expression of salt-stress-related genes (AtNHX1 and AtABI1) and improved the sensitivity to salt stress. In conclusion, these results suggest that MdZAT5 plays a positive regulatory role in anthocyanin accumulation and negatively regulates salt resistance.

Keywords: apple; MdZAT5; anthocyanin; salt tolerance

1. Introduction

Among the numerous external natural environmental factors, light (one of the most important) affects the entire plant life cycle [1–3]. Light is an essential factor in anthocyanin synthesis [4]. Previous studies have shown that the longer the light exposure and the greater the light intensity, the more conducive to anthocyanin formation [5,6]. Additionally, salt is one of the most important environmental stresses, which limits the growth and development of plants and poses a serious threat to global agriculture [7,8]. Salt stress significantly inhibits the growth of apple plants, mainly manifesting in slow growth, leaf wilting, and other symptoms, which even leads to plant death in serious cases [9]. Salt tolerance varies amongst different apple varieties. For example, Red fruit Begonia is affected by salt stress at 0.5 mg/g NaCl, and Mulus zhumei and Midget Crabapple appear to be affected by salt stress at 5.5 mg/g NaCl [10]. In order to adapt to complex and changing environmental factors, complex regulatory mechanisms have gradually evolved in plants [11–13]. At present, many transcription factors have been found to regulate anthocyanin accumulation and respond to salt stress, such as MYB, bHLH, WRKY, ZFP, DREB/CBF, NAC, and AP2/ERF [14–21].

Zinc finger proteins are among the most widely distributed proteins in eukaryotes [22,23]. The Cys2/His2 (C2H2)-type zinc finger proteins, also known as TFIIIA zinc
finger protein, are the most common protein in the zinc finger gene family [23]. TFIIIA transcription factor was first found in *Xenopus laevis* [24]. In plants, C2H2-type zinc finger proteins have a highly conserved amino acid sequence (QALGGH) located at the junction of the zinc finger and the DNA α Spiral zone [25]. In addition to the typical zinc finger domain that binds to DNA, most C2H2-type zinc finger proteins also contain three characteristic motifs [17,26]. One is the ERF-associated amphiphilic repression (EAR) motif (also known as DLN-box)—a short hydrophobic transcriptional repressor domain near the C-terminal. The second is a nuclear localization signal (NLS)—mainly related to subcellular localization. The third is an L-box—possibly related to protein interactions.

The C2H2-type zinc finger proteins play an important role in the growth and development of plants [27–29]. In *Arabidopsis*, JAGGED (JAG) can regulate cell differentiation and flower morphological development [30]. ZINC FINGER of *ARABIDOPSIS THALIANA* 6 (ZAT6) plays an important role in the regulation of anthocyanin under hydrogen peroxide treatment [18]. ZINC FINGER PROTEIN 5 (ZFP5) can affect root hair development by directly regulating root hair development-related genes [31]. In addition, it also plays an important role in abiotic stress [32,33]. *AtZAT18* positively regulates plant drought tolerance, while its mutant decreased the tolerance to drought stress in *Arabidopsis* [34]. *Triticum aestivum* predicted that Dof zinc finger protein (TaZNF) regulates salt tolerance [35]. Ectopic expression of *GmZFP3* in *Arabidopsis* plays a negative regulatory role in drought response [36]. *Oryza sativa* drought-responsive zinc finger protein 1 (OsDRZ1) overexpression in rice can improve plant drought tolerance by accumulating more proline and scavenging ROS [37]. A multi-stress-responsive gene (OsMSR15) in *Oryza sativa* L. shows positive regulation in cold, drought, and heat stress conditions at various developmental stages of rice [38]. MdZAT10 positively regulates JA-induced leaf senescence by interacting with MdBT2 (BTB-TAZ 2) and negatively regulates plant drought tolerance in apple [39,40].

So far, C2H2-type zinc finger proteins have been widely cloned and identified in *Arabidopsis*. However, ZAT5 has been rarely reported in other species, and its function in apple is poorly understood. In this assay, we identified a C2H2-type zinc finger protein transcription factor MdZAT5 in apple and characterized its roles in regulating apple calli and transgenic *Arabidopsis* anthocyanin accumulation and salt stress.

2. Results

2.1. Identification and Bioinformatics Analysis of the MdZAT5 Gene in Apple

*MdZAT5* (MD03G1128800) was identified as the closest apple homolog of the C2H2-type zinc finger transcription factor *AtZAT5* (At2G28200) on the NCBI database. Its full-length cDNA was 969 bp long and encoded 322 aa. As shown in Figure 1A, it contained two conserved zinc finger domains. We predicted the secondary structure of MdZAT5 protein and found that it was mainly random coils (69.88%), followed by alpha-helices (14.29%), extended-strands (12.73%), and beta-turns (3.11%) (Figure 1B). Based on the prediction results of the secondary structure, the tertiary structure was predicted (Figure 1C).

2.2. Phylogenetic and Conserved Motif Analysis of ZAT5 Proteins from Different Plants

In order to analyze the phylogenetic relationship between MdZAT5 protein and ZAT5 proteins of other species, an evolutionary tree was constructed using MEGA_X software. It was found that the ZAT5 protein of apple (*Malus domestica*) showed the closest evolutionary relationship with pear (*Pyrus × bretschneideri* and *Pyrus ussuriensis × Pyrus communis*) (Figure 2A). In addition, we compared the ZAT5 protein sequences of apple and the other nine species and found that they all contained two conserved zinc finger domains and an EAR motif (Figure 2B).
Figure 1. Basic information about the *MdZAT5* sequence. (A) Conserved sequence of *MdZAT5* protein. The blue rectangle indicates the zinc finger domain. The numbers represent the length of amino acids. (B,C) Predicted the secondary and tertiary protein structures of *MdZAT5*, respectively. The numbers denote the length of amino acids.

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Figure 2. Phylogenetic tree, amino acid sequence alignment, and conserved motifs analysis. (A) The phylogenetic tree of ZAT5 proteins from 18 different plants. (B) Comparison of amino acid sequences of ZAT5 proteins from 10 different plants. The red triangle represents *MdZAT5*. The red box represents a conserved domain. They all have two conserved zinc finger domains and an EAR motif.
2.3. Tissue Expression Pattern of MdZAT5

To further explore the potential biological function of MdZAT5 in apple, we detected its expression in five different organ tissues of apple (roots, stems, leaves, flowers, and fruits) by qRT-PCR. MdZAT5 was expressed in organs and tissues, among which the expression was the highest in flowers, followed by stems and leaves, and lower in roots and fruits (Figure 3A). The constructed ProMdZAT5::GUS vector was genetically transformed into Arabidopsis to obtain ProMdZAT5::GUS transgenic Arabidopsis. GUS staining showed that MdZAT5 was expressed in roots, stems, leaves, flowers, and fruits (Figure 3B), and the results were consistent with Figure 3A.

![Figure 3. Tissue expression analysis of MdZAT5. (A) The relative expression level of MdZAT5 in different tissues (roots, stems, leaves, flowers, and fruits) by qRT-PCR. (B) Tissue expression analysis of MdZAT5 via GUS staining in transgenic Arabidopsis. Different lowercase letters represent a significant difference ($p < 0.05$). Data are the mean ± SD of three independent replicates.](image)

2.4. Cis-Elements Analysis of MdZAT5 Promoter Sequence and Expression Patterns of MdZAT5

We analyzed MdZAT5 cis-elements in promoter 2000 bp upstream using the PlantCARE online software. There are various response elements in the MdZAT5 promoter (Table 1), such as cis-elements in response to plant hormones: abscisic responsive element (ABRE) and MeJA responsive element (CGTCA-motif). A large number of light response elements is known in the MdZAT5 promoter, such as Box 4, G-Box, GATA-motif, GT1-motif, MRE, and TCCC-motif. In addition, there is an ARE element that responds to hypoxia in the MdZAT5 promoter.

![Table 1. Cis-elements analysis of MdZAT5 promoter regions.](image)
To gain further insight into the expression patterns of *MdZAT5* with multiple abiotic stressors, we measured the expression of the *MdZAT5* gene under NaCl (150 mM), PEG6000 (10%), temperature (4 °C), and ABA (100 µM) treatments. Under treatment with 150 mM NaCl, the expression of *MdZAT5* reached its highest at 12 h, and the overall trend was first decreasing and then increasing (Figure 4A). In response to 10% PEG 6000, the expression of *MdZAT5* reached its highest at 12 h (Figure 4B). Under low temperature conditions (4 °C), *MdZAT5* showed a downward trend first and then an upward trend, reaching its maximum at 12 h (Figure 4C). The expression of *MdZAT5* showed an upward trend with the treatment of 100 µM ABA over time (Figure 4D). These results showed that *MdZAT5* expression was caused by different stressors, which meant that *MdZAT5* played an important regulatory role in the process of stress response.

2.5. The Abiotic Stress Response of *MdZAT5*

Based on the analysis of *cis*-acting elements and expression patterns, we treated Pro*MdZAT5::GUS* transgenic *Arabidopsis* with different treatment conditions. Compared with the control, 150 mM NaCl, 6% PEG6000, and high light could significantly promote GUS activity, but at 4 °C, it was lightly stained. Moreover, we found that 100 µM ABA caused no significant changes in leaf color relative to the control (Figure 5A). In addition, we also observed that the staining was deeper in older leaves and less intense in younger leaves, indicating that *MdZAT5* was expressed higher with leaf age. We obtained Pro*MdZAT5::GUS* transgenic calli, and their staining results were consistent with that of Pro*MdZAT5::GUS* transgenic *Arabidopsis* (Figure 5A–C).

![Figure 4](image-url)  
**Figure 4.** The expression pattern of *MdZAT5* under different treatment conditions. The relative expression of *MdZAT5* in 150 mM NaCl (A), 10% PEG6000 (B), 4 °C (C), and 100 µM ABA (D), respectively. Different lowercase letters represent a significant difference (p < 0.05). Data are the mean ± SD of three independent replicates.
2.6. Overexpression of MdZAT5 in Apple Calli and Arabidopsis Promoted the Accumulation of Anthocyanin

To study the function of MdZAT5, we constructed the MdZAT5 overexpression vector and transferred it into apple calli. Wild-type calli (WT) and transgenic apple calli (MdZAT5-OVX) were cultured under high light for 18 days. As shown in Figure 6A, MdZAT5-OVX accumulated more anthocyanins, while WT accumulated less anthocyanins. At the same time, we used a spectrophotometer to quantitatively determine the content of anthocyanin. The results also showed that the anthocyanin content of MdZAT5-OVX was higher than that of WT (Figure 6B). In addition, the expression levels of flavonoid structural genes in WT and MdZAT5-OVX were analyzed by qRT-PCR. The results showed that compared with WT, the expression levels of anthocyanin biosynthesis-related genes (MdANR, MdCHS, MdDFR, MdF3H, and MdUFGT) increased by different degrees (Figure 6C).

In addition, we genetically transformed the constructed MdZAT5 overexpression vector into Arabidopsis (Columbia ecotype) and obtained three overexpression lines (OE1, OE2, and OE3) (Figure 6F). Comparing the anthocyanin content of Col-0 and MdZAT5-OE, it was found that under high light, the anthocyanin content accumulated by the three overexpression lines was significantly higher than that of Col-0 (Figure 6G).

2.7. MdZAT5 Increased Sensitivity to Salt Stress in Transgenic Apple Calli and Arabidopsis

To further explore the function of MdZAT5 under abiotic stress, the 16-day-old WT and MdZAT5-OVX were transferred to 100 mM NaCl MS medium. As shown in Figure 7A, the growth rate of MdZAT5-OVX was much lower than that of WT in 100 mM NaCl. Meanwhile, the fresh weight of MdZAT5-OVX was significantly lower than that of WT, and its MDA content and relative electronic conductivity were significantly higher than that of WT (Figure 7B–D). Therefore, the overexpression of MdZAT5 negatively regulates the salt resistance of apple calli.

Figure 5. GUS staining in ProMdZAT5::GUS transgenic calli and Arabidopsis. The ProMdZAT5::GUS transgenic Arabidopsis (A) and calli (B), treated with 24 °C, 150 mM NaCl, 6% PEG, 4 °C, 100 µM ABA, and high light. (C) The GUS activity of MdZAT5 of (B). Different lowercase letters represent a significant difference (p < 0.05). Data are the mean ± SD of three independent replicates.

Figure 6. (A) The growth rate of WT and transgenic apple calli (OE1, OE2, and OE3) (Figure 6F). Comparing the anthocyanin content of Col-0 and MdZAT5-OE, it was found that under high light, the anthocyanin content accumulated by the three overexpression lines was significantly higher than that of Col-0 (Figure 6G).
Figure 6. Overexpression of *MdZAT5* in apple calli and *Arabidopsis* promoted anthocyanin accumulation. The phenotypes (A) and anthocyanin content (B) of WT and *MdZAT5-OVX*. Expression analysis of *MdZAT5* (D) and genes involved in anthocyanin biosynthesis-related genes (*MdANR*, *MdCHI*, *MdCHS*, *MdDFR*, *MdF3H*, and *MdUFGT*) (C) in WT and *MdZAT5-OVX*. The phenotypes (E) and anthocyanin content (G) of Col-0 and *MdZAT5-OE*. (F) Expression analysis of *MdZAT5* in Col-0 and *MdZAT5-OE*. Different lowercase letters represent a significant difference (*p* < 0.05). Data are the mean ± SD of three independent replicates.
Meanwhile, the fresh weight of MdZAT5-OVX was significantly lower than that of WT, and its MDA content and relative electronic conductivity were significantly higher than that of WT (Figure 7B–D). Therefore, the overexpression of MdZAT5 negatively regulates the salt resistance of apple calli.

Figure 7. MdZAT5 enhanced the sensitivity to salt in apple calli. (A) The phenotypes of WT and MdZAT5-OVX with 100 mM NaCl. Fresh weight (B), MDA content (C), relative electronic conductivity (D) of WT, and MdZAT5-OVX. Different lowercase letters represent a significant difference ($p < 0.05$). Data are the mean ± SD of three independent replicates.

Furthermore, Arabidopsis seedlings (Col-0 and MdZAT5-OE) were grown on MS medium for 3 days and then transferred to 150 mM NaCl MS medium for 14 days. In the control condition, there was no significant difference in the number of lateral roots or primary root lengths. However, the number of lateral roots of MdZAT5-OE was greater than that of Col-0, and the length of primary roots was lower than that of Col-0 under the salt treatment (Figure 8B,C). In addition, we conducted salt tolerance tests on Col-0 and transgenic plants for 14 days. As shown in Figure 8D, Col-0 grew normally, while MdZAT5-OE had yellow and wilting leaves. Simultaneously, the content of MDA in MdZAT5-OE was significantly higher than that in Col-0 plants under salt stress. In order to further study the role of MdZAT5 in the signal pathway of salt stress, we detected the expression levels of AtNHX1 and AtABI1 in Arabidopsis by qRT-PCR. The results showed that the expression levels of AtNHX1 and AtABI1 in MdZAT5-OE Arabidopsis were significantly lower than those in Col-0 (Figure 8F,G).

2.8. Ectopic Expression of MdZAT5 Increased ROS Accumulation under Salt Stress

$H_2O_2$ levels were measured using diaminobenzidine (DAB) staining and $O_2^-$ using nitro blue tetrazolium (NBT). In the control group, no significant difference was observed between WT and MdZAT5-OE. Under salt treatment, the staining of MdZAT5-OE was deeper, while that of Col-0 was weaker (Figure 9A,B). At the same time, we quantitatively determined $H_2O_2$ and $O_2^-$, and the results were consistent with their staining results. The contents of $H_2O_2$ and $O_2^-$ of MdZAT5-OE were significantly higher than those of Col-0 under salt stress, indicating that MdZAT5 increased sensitivity to salt stress in Arabidopsis (Figure 9C,D).
Figure 8. MdZAT5 enhanced the sensitivity to salt in transgenic Arabidopsis. (A) The phenotypes of Arabidopsis seedlings treated with MS medium, 150 mM NaCl treatment. Lateral root numbers (B) and primary root length (C) in Col-0 and MdZAT5-OE. (D) Phenotypes of Arabidopsis treated with 150 mM NaCl after 14 days and MDA content (E). The expression level of AtNHX1 (F) and AtABI1 (G) in Col-0 and MdZAT5-OE. Different lowercase letters represent a significant difference ($p < 0.05$). Data are the mean ± SD of three independent replicates.
were confirmed to perform biological functions. In this study, Arabidopsis [43–45]. Different treatments (salt, drought, cold, ABA, and high light) induced transgenic calli (Figure 5). Under high light, overexpression of ProMdZAT5::GUS up-regulated under high light treatment, consistent with the result of GUS staining of number of light-responsive elements (Table 1). Light is the most important external factor in flowers (Figure 3A). We analyzed promoter sequences of fruits, stems, leaves, and seeds of plants [47]. Here, the expression of were confirmed to perform biological functions. In this study, many abiotic stressors (Figures 4 and 5). Previous studies have shown that bidopsis regulated and biotic stress responses in plants [37,41,42]. In recent years, many studies have studied function of C2H2-type zinc finger proteins. Their functions have been widely characterized in Arabidopsis and rice. However, little has been reported in apple. Here, we isolated ZAT5 from apple and found that both ZAT5 and AtZAT5 contained two highly conserved zinc finger domains and an EAR motif, indicating that it has the conserved function of this family (Figures 1A and 2B). The EAR motif, a short hydrophobic region, has been shown to function as repressor, e.g., ZAT6, STZ/ZAT10, ZAT11, and ZAT12 in Arabidopsis [43–45]. Different treatments (salt, drought, cold, ABA, and high light) induced the expression of MdZAT5, suggesting that MdZAT5 may be involved in the tolerance of many abiotic stressors (Figures 4 and 5). Previous studies have shown that AISTZ/ZAT10 is induced by the same treatments [44,46]. MdZAT5 transgenic apple calli and Arabidopsis were confirmed to perform biological functions. In this study, MdZAT5 was involved in anthocyanin synthesis and salt stress response in apple.

Anthocyanin, a polyphenol water-soluble plant pigment, exists widely in flowers, fruits, stems, leaves, and seeds of plants [47]. Here, the expression of MdZAT5 was highest in flowers (Figure 3A). We analyzed promoter sequences of MdZAT5 and found a large number of light-responsive elements (Table 1). Light is the most important external factor regulating anthocyanin synthesis [4]. ProMdZAT5::GUS transgenic Arabidopsis was deeply up-regulated under high light treatment, consistent with the result of GUS staining of ProMdZAT5::GUS transgenic calli (Figure 5). Under high light, overexpression of MdZAT5 actively regulated anthocyanin accumulation in apple calli and Arabidopsis, further indicating that MdZAT5 plays an important role in plant response and adaptation to high light. Under high light, AtZAT12 promotes the increase of anthocyanin and chlorophyll content [48]. In Petunia, ZPT2-1 participates in anthocyanin synthesis [49]. Few studies have

3. Discussion

C2H2-type zinc finger proteins played key roles in regulating plant growth, abiotic, and biotic stress responses in plants [37,41,42]. In recent years, many studies have studied the function of C2H2-type zinc finger proteins. Their functions have been widely characterized in Arabidopsis and rice. However, little has been reported in apple. Here, we isolated ZAT5 from apple and found that both ZAT5 and AtZAT5 contained two highly conserved zinc finger domains and an EAR motif, indicating that it has the conserved function of this family (Figures 1A and 2B). The EAR motif, a short hydrophobic region, has been shown to function as repressor, e.g., ZAT6, STZ/ZAT10, ZAT11, and ZAT12 in Arabidopsis [43–45]. Different treatments (salt, drought, cold, ABA, and high light) induced the expression of MdZAT5, suggesting that MdZAT5 may be involved in the tolerance of many abiotic stressors (Figures 4 and 5). Previous studies have shown that AISTZ/ZAT10 is induced by the same treatments [44,46]. MdZAT5 transgenic apple calli and Arabidopsis were confirmed to perform biological functions. In this study, MdZAT5 was involved in anthocyanin synthesis and salt stress response in apple.

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Figure 9. Ectopic expression of MdZAT5 enhances ROS buildup under salt treatment. DAB staining for H$_2$O$_2$ (A) and NBT staining for O$_2^-$ (B) in Col-0 and MdZAT5-OE Arabidopsis leaves after 14 days of salt treatment. H$_2$O$_2$ content (C) and O$_2^-$ generation rates (D) in Col-0 and MdZAT5-OE Arabidopsis after 14 days of salt treatment. Different lowercase letters represent a significant difference ($p < 0.05$). Data are the mean ± SD of three independent replicates.
reported the functions of zinc finger proteins in the regulation of anthocyanin synthesis; thus, more work is needed in this area to perfect the content.

Salt stress is one of the most important limiting factors in plant growth, development, and yield [50,51]. Therefore, it is necessary to study salt-stress-related genes and their functions to improve crops. Previous studies have reported that many C2H2-type zinc finger proteins are involved in the regulation of salt stress as transcriptional activators or inhibitors [52]. In Arabidopsis, SALT-INDUCED ZINC FINGER PROTEIN1 (AtSIZ1) positively regulates salt tolerance by maintaining osmotic balance and ion homeostasis [53]. RING/FYVE/PHD ZFP (AtRZFP) enhances salt and osmotic tolerance by scavenging ROS, maintaining Na\(^{+}\) and K\(^{+}\) homeostasis [54]. AtZFP3 can enhance the salt resistance of Arabidopsis, and its expression level is inhibited under salt stress [55]. OsZFP213 interacts with OsMAPK3 to improve salt tolerance by scavenging reactive oxygen [56]. In other species, there are also reports of C2H2-type zinc finger proteins involved in salt stress, such as wheat, soybean, tomato, and sweet potato [35,57–59].

In this study, ProMdZAT5::GUS transgenic Arabidopsis and calli also further confirmed that MdZAT5 was related to salt stress (Figure 5). Under salt stress, transgenic MdZAT5 apple calli and Arabidopsis showed weaker growth than wild-type plants, indicating that MdZAT5 plays a negative regulatory role in plant response and adaptation to salt stress (Figures 7 and 8). We measured the content of MDA in apple calli and Arabidopsis, and found that the MDA content of transgenic plants was higher than in those of WT plants (Figures 7C and 8E). The content of MDA can reflect the degree of stress damage to plants [60]. The accumulation of ROS was related to the content of MDA [61]. Deeper levels of DAB/NBT staining, higher H\(_2\)O\(_2\) content levels, and higher O\(_2^{-}\) generation rates under salt treatment indicate that more ROS accumulation occurred in transgenic plants (Figure 9). In addition, maintaining ion balance is an important method for plants to resist salt stress [62]. NHX1, a Na\(^{+}\)/H\(^{+}\) antiporter located on the vacuolar membrane, plays an important role in maintaining ion homeostasis in plant cells [63]. ABI1 is a salt-stress-related gene [64]. The AtNHX1 and AtABI1 expression levels in the overexpressed MdZAT5 Arabidopsis were significantly lower than those in Col-0 (Figure 8F,G). The above results show that overexpression of MdZAT5 enhances the sensitivity of plants to salt stress by reducing the expression level of NHX1 and ABI1.

In this study, we identified a novel MdZAT5 transcription factor that directly or indirectly activates the expression of anthocyanin synthesis-related genes to increase anthocyanin accumulation or reduce the expression of salt-stress-related genes to improve the sensitivity of salt stress. This study provides new insights for future research on anthocyanin accumulation and resistance to salinization and provides new candidate genes for improving apple quality and abiotic stress. However, the potential mechanism of MdZAT5 in regulating anthocyanin accumulation and salt stress is not clear. We next intend to further improve its regulatory mechanism by verifying the direct downstream protein or gene targets of MdZAT5.

4. Materials and Methods
4.1. Plant Materials and Growth Conditions

The apple (Malus × domestica ‘Royal Gala’) shoot cultures were stored at 25 °C on MS solid medium containing 0.5 mg/L 6-benzylaminopurine (6-BA) and 0.5 mg/L naphthyl acetic acid (NAA) for a 16/8 h light/dark photoperiod and subcultured at 30-day intervals. To obtain self-rooted plantlets, the 3-week-old shoot cultures were transferred to a root-inducing MS solid medium containing 0.2 mg/L indoleacetic acids (IAA). For tissue expression analysis, the roots, stems, leaves, flowers, and fruits were collected 80 days after flowering from 7-year-old ‘Gala’ apple tree (Taian, China). Four-week-old self-rooted apple seedlings were treated with NaCl (150 mM), PEG 6000 (10%), temperature (4 °C), and ABA (100 µM), as described in [45].

Apple calli from the ‘Orin’ cultivar were grown on MS solid medium of 1.5 mg/L 2, 4-dichlorophenoxyacetic acid (2, 4-D), and 0.4 mg/L 6-BA for 18 days in the dark at
24 °C. Then, for the stress treatment, calli were cultured in a medium with high light and NaCl (100 mM) for 18 days. The seeds of ecotype Columbia (Col-0) and transgenic Arabidopsis (MdZAT5-OE1, MdZAT5-OE2, MdZAT5-OE3) were sterilized and vernalized for 3 days at 4 °C. Then, Arabidopsis seedlings were sown on MS solid medium for 3 days under a photoperiod of 16/8 h light/dark and transferred to solid medium with high light (~300 µmol m⁻² s⁻¹) and NaCl (150 mM) for 7 and 14 days, respectively. Twenty-one-day-old seedlings of Arabidopsis were supplied with 150 mM NaCl for 14 days in soil.

4.2. Bioinformatics Analysis of the MdZAT5 Gene

The basic information of the MdZAT5 sequence came from the NCBI database (https://www.ncbi.nlm.nih.gov/, accessed on 3 November 2020). The secondary and tertiary structure prediction of MdZAT5 adopted SOPMA (https://npsa-prabi.ibcp.fr/cgi-bin/npsa_automat.pl?page=npsa_sopma.html, accessed on 15 March 2021) and Phyre2 (http://www.sbg.bio.ic.ac.uk/~phyre2/html/page.cgi?id=index, accessed on 15 March 2021), respectively [65].

4.3. Phylogenetic Analysis and Multiple Sequence Alignment of ZAT5 Proteins

The adjacency algorithm of the online software MEGA_X was used to construct ZAT5 evolutionary neighbor-joining trees of different plants (the step test was set to 1000 times, substitution method was the Poisson model) [66]. A multiple sequence alignment of ZAT5 proteins from 10 different plants was performed using Clustal Omega. We found several highly conserved domains, which were visualized by the online software Jalview.

4.4. Analysis of the MdZAT5 Promoter

The cis-element in the MdZAT5 promoter (2000 bp upstream of the transcription initiation site) was analyzed with the online software PlantCARE (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/, accessed on 18 March 2021) [67].

4.5. RNA Extraction and qRT-PCR Assays

Total RNAs of plant materials, including apple and Arabidopsis, were isolated using the RNA Plant Plus Reagent Kit (Tiangen Biotech, Beijing, China). Reverse transcription was conducted for single-stranded DNA synthesis using the PrimScript™ First Strand cDNA Synthesis Kit (TaKaRa, Dalian, China), per the manufacturer’s protocol. qRT-PCR was performed on the extracted RNA using an ABI7500, in which 18S (apple) and AtACTIN (Arabidopsis) were used as internal control. Then, relative gene expression analysis was conducted using the cycle threshold (Ct) $2^{-ΔΔCt}$ method. Quantitative primers are listed in Supplementary Table S1.

4.6. Construction of the Expression Vectors and Genetic Transformation

To construct the overexpression vector of MdZAT5, we cloned the full-length coding sequence of MdZAT5 into the plant expression vector pRI101 [40]. The MdZAT5 (2000 bp promoter fragment from start codon) was cloned into a p1300-GN vector to construct ProMdZAT5::GUS, and the vector drives Gus (β-glucuronidase) reporter gene [64]. The MdZAT5 overexpression vector and ProMdZAT5::GUS constructs were transformed into Arabidopsis by the flower dipping method to obtain transgenic plants [68]. The calli of transgenic apple were obtained by the Agrobacterium-mediated method [69]. Transgenic Arabidopsis and calli were identified with kanamycin.

4.7. GUS Histochemical Staining

The 18-day-old transgenic apple calli and 7-day-old transgenic Arabidopsis seedlings of ProMdZAT5::GUS were cultured on solid medium with NaCl (150 mM), PEG 6000 (10%), temperature (4 °C), ABA (100 µM), and high light (~300 µmol m⁻² s⁻¹) for 24 h. Apple calli or Arabidopsis seedlings were subjected to staining using GUS staining buffer (containing
0.5 mM ferrocyanide, 0.1% Triton X-100, 0.1 mM EDTA, 0.5 mM ferricyanide, and 1 mM X-Gluc) and then decolorized with absolute ethanol for 12 h [70].

4.8. Measurements of Anthocyanin, Relative Electronic Conductivity, and MDA

The anthocyanin of plants was extracted by the methanol hydrochloric acid method [71]. The plant materials were placed in anthocyanin extract (95% ethanol: 1.5 M HCl = 85:15, \( v/v \)) in the dark at room temperature for 24 °C. The absorbance value of the extracted samples was measured at 530, 620, and 650 nm with a UV–Vis spectrophotometer. The anthocyanin content was calculated according to the previous method [72].

The relative electronic conductivity content of apple calli was measured by this method and a DDS-12 conductometer (Hangzhou Wanda Instrument Factory, Hangzhou, China) [73]. The content of MDA in apple calli and Arabidopsis was determined by the thiobarbituric acid (TBA)-based method [74].

4.9. Measurement of ROS

The \( \text{H}_2\text{O}_2 \) content and \( \text{O}_2^- \) production rate were determined using a kit (Keming, Suzhou, China). \( \text{H}_2\text{O}_2 \) level was detected by diaminobenzidine (DAB) histochemical staining, and \( \text{O}_2^- \) level was detected by nitro blue tetrazolium (NBT) staining in accordance with the methods described in [75].

4.10. Statistical Analysis

Each experiment was repeated at least three times (biological repetitions). Each biological repetition was performed at least three times (technical repetitions). All Data processing system (DPS) was used to analyze the significance of the data [76].

5. Conclusions

In short, overexpression of \( \text{MdZAT5} \) promotes the expressions of anthocyanin biosynthesis-related genes to actively regulate anthocyanin synthesis and increase sensitivity to salt stress by regulating the expression of \( \text{NHX1} \) and \( \text{ABI1} \). Our study provides new insight into \( \text{MdZAT5} \)-mediated anthocyanin synthesis and salt resistance and is helpful to further clarify the mechanism of anthocyanin synthesis and salt stress.

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