Overexpression of a Multiprotein Bridging Factor 1 Gene \textit{DgMBF1} Improves the Salinity Tolerance of Chrysanthemum

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Abstract: Soil salinity represents a major constraint in the growth of chrysanthemum. Therefore, improving salinity tolerance of chrysanthemum has become an important research direction in tolerance breeding. Multiprotein bridging factor 1 (MBF1) is an evolutionarily highly conserved transcriptional co-activator in archaea and eukaryotes and has been reported to play important roles to respond to abiotic stresses. Here, a MBF1 gene induced by salt stress was isolated and functionally characterized from \textit{Dendranthema grandiflorum} and name as \textit{DgMBF1}. Overexpression of \textit{DgMBF1} in chrysanthemum increased the tolerance of plants to high salt stress compared to wild type (WT). It also showed fewer accumulations of hydrogen peroxide (H$_2$O$_2$), superoxide anion (O$_2^-$), higher activities of antioxidant enzymes, more content of proline and soluble sugar (SS) and more favorable K$^+$/Na$^+$ ratio than those of WT under salt stress. In addition, the expression level of genes related to antioxidant biosynthesis, proline biosynthesis, glyco-metabolism and K$^+$/Na$^+$ homeostasis was statistically significant higher in the \textit{DgMBF1}-overexpressed lines than that in WT. These results demonstrated that \textit{DgMBF1} is a positive regulator in response to salt stress and could serve as a new candidate gene for salt-tolerant plant breeding.

Keywords: multiprotein bridging factor 1; \textit{DgMBF1}; transgenic chrysanthemum; salt stress tolerance; gene expression

1. Introduction

Land salinization is a worldwide ecological issue. As one of the limiting factors, it severely hampers the growth of plants and the production of crops [1,2]. It has been determined that plants can effectively respond to environmental stress by identifying a series of complex biological signals and activating its transduction mechanism [3]. Generally, two phases of stress are revealed after the occurrence of salt stress: a rapid osmotic stress and a slower ionic stress [4]. Accordingly, there are three major physiological adaptive mechanisms of salt tolerance: osmotic stress tolerance, maintenance of ions homeostasis, and compartmentalization of Na$^+$ to reduce cytosolic Na$^+$ concentrations [5]. In addition, reactive oxygen species (ROS) plays a key role in the response to salt stress. ROS is a kind of toxic molecules that causes oxidative damage to proteins, DNA and lipids [6]. Under normal conditions, ROS was produced at a low level in organelles such as chloroplasts, mitochondria and peroxisomes. Under salt stress, the production rate of ROS is dramatically elevated. The balance between production and elimination of ROS determines its accumulation greatly [7].
At present, billions of hectares of various saline-alkali land exist globally, accounting for one-tenth of the world’s arable land. Twenty-three percent of cultivated land is salt-affected. [8] Chrysanthemums are produced mainly by facility cultivation. Due to the continuous increase of irrigation times in the facility, the transpiration of chrysanthemum speeds. However, the rainwater cannot take the salt away in time. Deep in the soil layer, the salt remains completed [9]. The continuous planting makes the salt accumulate year by year in soil. It poses a serious impact on the production and quality of chrysanthemums. Therefore, our question is how to cultivate and breed chrysanthemums under strong salt tolerance became a botanical concern. Genetic engineering methods can be used to transfer salt-tolerant genes into plants, which significantly improve the salt tolerance in transgenic plants.

Transcriptional regulatory proteins were involved in a variety of biological processes, especially in the expression of genomic information [10–12]. Among these proteins, transcriptional co-activators interacted with transcription factors, regulatory elements and the basal transcription machinery to complete the eukaryotic gene expression [13]. MBF1 proteins functioned as a non-DNA binding transcriptional co-activator. By bridging TATA box binding proteins, specific transcription factors were activated. The transcription of its target genes was enhanced to bind to target promoters in eukaryotes and to participate in multiple growth and developmental processes [14,15].

MBF1s elevated the regulation of multiple biological processes in yeast and animals [16,17]. MBF1s were also known to participate in a variety of abiotic and biotic stresses responses in plants [18]. There were three different homologs that could encode MBF1 in Arabidopsis thaliana. Their existence complemented the deletion of MBF1 in yeast [19]. The expression of AtMBF1a and AtMBF1b were regulated by plant development. The overexpression of AtMBF1a improved salt tolerance, fungal resistance and glucose insensitivity in transgenic Arabidopsis plants [20]. AtMBF1c was highly induced by high salt, dehydration, heat, H2O2, methyl viologen and pathogen infection [20–22]. Overexpression of CaMBF1 decreased high salt and cold stresses tolerance in Arabidopsis [23]. The expression level of StMBF1 in potato stems was enhanced by heat shock and H2O2 treatments [24]. Simultaneous treatment with high temperature and drought could induce the expression of MBF1 in tobacco [21].

Chrysanthemum is a world famous kind of cut flower and is susceptible to salt stress [25]. An et al. [26] demonstrated that overexpression of CcSOS1 improved the salt tolerance capacity in chrysanthemum. DgNAC1-overexpressed chrysanthemum held a higher survival rate under drought and salt stresses [27,28]; Overexpression of DgWRKY2, DgWRKY4 and DgWRKY5 genes enhanced salt tolerance in chrysanthemum [12,29,30]. To better understand the role MBF1 played in response to salt stress in chrysanthemum, we isolated a MBF1 gene from chrysanthemum and called it DgMBF1. Overexpression of DgMBF1 in chrysanthemum enhanced salt tolerance in plants, indicating that the DgMBF1 could be served as a new positive regulator of plants under salt stress.

2. Results

2.1. DgMBF1 Clone and Sequence Analysis

A salt-responsive multiprotein bridging factor gene identified from chrysanthemum was named as DgMBF1. The full-length cDNA of DgMBF1 was determined through polymerase chain reaction (PCR) and inserted into pCAMBIA 2300 controlled by the cauliflower mosaic virus (CaMV) 35S promoter. The obtained vector was transformed into the leaf disc of chrysanthemum by Agrobacterium tumefaciens. The expression level of DgMBF1 was measured through quantitative real-time PCR (qRT-PCR). Two fully overexpressed (OE) lines (OE-3, OE-34) was selected for subsequent experiments independently.

The full-length DgMBF1 gene was 733 bp, in which a 438 bp open reading frame (ORF) consisted. The ORF could encode 153 amino acids. Sequence alignments by DNAMAN showed that the DgMBF1 protein contained an MBF1 domain at the N-terminal region and a helix-turn-helix (HTH) domain at the C-terminal region (Figure 1a). DgMBF1 shared 92% identity with AaMBF1c (Artemisia annua, PWA60867.1), 83% with HaMBF1c (Helianthus annuus, XP_022027306.1) and LaMBF1c (Lactuca sativa, XP_023735987.1), 76% with LnMBF1c (Lpomoea nil, XP_019191938.1), 73% with AtMBF1c (Arabidopsis
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*thaliana*, AEE76905.1), and 71% identity with *MtMBF1* (*Medicago truncatula*, AES76734.2) (Figure 1b). Phylogenetic analysis indicated that *DgMBF1* was significantly more homologous to MBF1c than to MBF1a and MBF1b. Therefore, *DgMBF1* was classified as a member of the plant group II MBF1c protein.

**Figure 1.** Sequence analysis of *DgMBF1*. (a) Multiple alignments of predicted amino acid sequences of *DgMBF1* with other plant MBF1 proteins. The shade of colors is used to distinguish the degree of consistency. Dark blue: completely consistent; pink: ≥ 75%; light blue: ≥ 50%. Red frame is used to circle their domains. (b) Phylogenetic analysis of *DgMBF1* protein sequence with other plant MBF1 proteins. *DgMBF1* is highlighted with a red frame. MBF1 proteins used in this analysis were as follows: AaMBF1c (*Artemisia annua*, PWA60867.1), HaMBF1c (*Helianthus annuus*, XP_022027306.1), LaMBF1c (*Lactuca sativa*, XP_023735987.1), AtMBF1c (*Arabidopsis thaliana*, AEE76905.1), LnMBF1c (*Lpomoea nil*, XP_019191938.1), SoMBF1c (*Spinacia oleracea*, XP_021838271.1), PaMBF1c (*Polytrichastrum alpinum*, AJG41867.1), MtMBF1 (*Medicago truncatula*, AES76734.2), OsMBF1c-like (*Oryza sativa*, XP_015641831.1), VvMBF1 (*Vitis vinifera*, XP_002280992.1), GmMBF1a (*Glycine max*, XP_003527342.1), TaMBF1 (*Triticum aestivum*, ACO36694.1), ZmMBF1 (*Zea mays*, ACG33346.1), StMBF1 (*Solanum tuberosum*, AF232062.1), CaMBF1 (*Capsicum annuum*, JX402927.1), NtMBF1 (*Nicotiana tabacum*, BAB88859.1), AtMBF1a (*Arabidopsis thaliana*, AF370280), AtMBF1b (*Arabidopsis thaliana*, AF326909.1), LeMBF1 (*L. esculentum*, AF096246).
2.2. Expression Analysis of DgMBF1

The expression profile of DgMBF1 in different tissues was detected by qRT-PCR. The result suggested that the relative expression of DgMBF1 was highest in leaves, followed by stems, and lowest in roots, and DgMBF1 was expressed in the flowers (Figure 2a). Expression patterns of DgMBF1 gene in leaves under salt stress were also detected by qRT-PCR. Under salinity, DgMBF1 transcript increased continuously until 24 h and remained at a higher level compared to untreated control (Figure 2b). The result indicated that DgMBF1 was involved in salt tolerance.

![Figure 2](image-url)

**Figure 2.** Quantitative real-time PCR analysis of DgMBF1 expression in different tissues and in response to salt treatment. (a) Expression patterns of DgMBF1 in leaves, stems and roots. (b) Salt treatment. Data represent means and standard errors of three replicates. Different letters above the columns indicate significant differences ($p < 0.05$) on the basis of Duncan’s multiple range test.

2.3. Observation of Callus and Phenotype

The growth of small buds on the infested leaf disc in chrysanthemum leaves is shown in Figure 3a. Positive plants were screened through DNA detection. Under normal conditions, there was no significant difference in phenotype between WT and transgenic chrysanthemum plants (Figure 3b).

![Figure 3](image-url)

**Figure 3.** Callus transformation process and phenotypic observation. (a) Transgenic chrysanthemum callus transformation process. (b) Phenotypic observation of WT and transgenic chrysanthemums.
2.4. Overexpression of DgMBF1 in Chrysanthemum Enhanced the Salt Tolerance

To further verify the function of DgMBF1, we generated chrysanthemum transgenic plants. Two independent OE lines (OE-3, OE-34) were obtained and the transcript abundance of DgMBF1 in leaves were determined by qRT-PCR (Figure 4a). The result indicated that DgMBF1 transcript expression in two OE lines was significantly (p < 0.05) higher than that in WT. Under normal conditions, no obvious differences in phenotypes were observed between OE lines and WT lifelong (Figure 4d). Under salt treatment, leaves of WT plants turned yellow and even wilted with the gradual increase of salinity. While OE lines remained green and exhibited significantly (p < 0.05) higher survival rate than that of WT (Figure 4c, d). Moreover, the survival rates of OE-3 and OE-34 were 78.65% and 80.92%, respectively. In contrast, the WT was only 32.13% (Figure 4b).

Figure 4. Overexpression of DgMBF1 in transgenic chrysanthemum resulted in enhanced tolerance to salt stress. (a) Transcript levels of DgMBF1 in WT and OE lines. (b) The survival rates of OE lines and WT after two weeks’ recovery. (c) Phenotypic comparison of OE lines and WT under salt stress. (d) OE lines and WT grown under normal (control) and salt stress conditions, followed by a recovery. Data represent means and standard errors of three replicates. The different letters above the columns indicate significant differences (p < 0.05) according to Duncan’s multiple range test.

2.5. Influence of Salt Stress on the Growth and Development of Chrysanthemum

To study the salt impact on the growth and development of chrysanthemum, the root length and fresh weight were measured. Salinity inhibited the growth of chrysanthemum; all lines exhibited the atrophy of root and the reduction of fresh weight. While the reduction rate of OE lines was far (p < 0.05) smaller compared with WT (Figure 5).
Figure 5. Assay of root length, fresh weight in OE lines and WT under salt stress. (a) Relative root length. (b) Relative fresh weight. Root length and fresh weight are relative to that of WT for 0 days. The different letters above the columns indicate significant differences (p < 0.05) according to Duncan’s multiple range test.

2.6. Overexpression of DgMBF1 Enhanced Oxidation Tolerance under Salt Stress

The accumulation of H$_2$O$_2$ and O$_2^-$ in leaves was determined by diaminobenzidine (DAB) and nitro blue tetrazolium (NBT) staining in both WT and OE lines to measure the oxidation of chrysanthemum directly. The OE lines accumulated less H$_2$O$_2$ and O$_2^-$ than the WT with NaCl solutions treatment, for less brown and blue spots were observed (Figure 6c,d). Quantitative analysis also indicated that H$_2$O$_2$ and O$_2^-$ contents in leaves increased under salt stress in both WT and OE lines. OE lines (p < 0.05) accumulated less H$_2$O$_2$ and O$_2^-$ than WT. Under 15 days of salt stress, the content of H$_2$O$_2$ in OE lines increased to 2.11- and 2.36-fold, while the WT increased to 2.93-fold. The content of O$_2^-$ in OE lines increased by 31.08% and 23.52% than day 0, less than that in WT (Figure 6a,b). In addition, antioxidant enzymes including superoxide Anion (APX), peroxidase (POD), superoxide (SOD) and catalase (CAT) in leaves also exhibited higher activities in OE lines than that in WT plants (Figure 7a,d). Moreover, we examined the relative expression of DgCuZnSOD, DgCAT and DgAPX, which related to ROS-scavenging system in leaves. The results revealed that the transcript accumulation of DgCuZnSOD, DgCAT and DgAPX were statistically (p < 0.05) higher in OE lines than that in WT under salt stress (Figure 7e,g).

Taken together, through regulating the expression of these antioxidant-related genes, overexpressed DgMBF1 respond to against ROS persecution. Less H$_2$O$_2$ and O$_2^-$ contents and higher antioxidant enzyme activities would eliminate stress-induced ROS. Thus, the salt tolerance in transgenic chrysanthemum would be elevated.
Figure 6. Analysis of ROS accumulation levels under salt stress. (a,b) Quantitative measurement of \( \text{H}_2\text{O}_2 \) and \( \text{O}_2^- \) contents. (c,d) Analysis of \( \text{H}_2\text{O}_2 \) and \( \text{O}_2^- \) contents by NBT staining and DAB staining. Data represent means and standard errors of three replicates. The different letters above the columns indicate significant differences (\( p < 0.05 \)) according to Duncan’s multiple range test.
2.7. Overexpression of DgMBF1 Promoted the Accumulation of Osmotic Substances under Salt Stress

Proline and SS contents in leaves were measured to determine the regulation of osmotic mechanism in chrysanthemum under salt stress. The OE lines and WT exhibited little differences in the contents of proline and SS under normal conditions. However, the contents of these two osmotic regulatory factors both increased in OE lines and WT under salinity. The proline and SS highly accumulated in the OE lines (p < 0.05). By day 10, proline and SS contents in transgenic chrysanthemum reached a maximum; and by day 15, proline content was proximately 1.53- and 1.85-fold greater than that in WT. SS content was about 1.6- and 1.53-fold greater than that in WT. (Figure 8a,b). The relative expression of genes in leaves, related to proline biosynthesis and glycol-metabolism, including DgP5CS, Dg6PGDH, and DgMDH were also significantly (p < 0.05) up-regulated in OE lines compared with that in WT plants under salt stress (Figure 8c,e).

These results showed that overexpression of DgMBF1 conferred a higher osmotic pressure on transgenic chrysanthemum to cope with the dehydration stress caused by salinity.
Figure 8. Overexpression of DgMBF1 promotes the accumulation of osmotic substances. (a,b) The proline and SS contents in WT and OE lines under salt stress. (c–e) Relative expression level of genes involved in metabolism of proline and soluble sugars in WT and OE lines under salt stress. Data represent means and standard errors of three replicates. The different letters above the columns indicate significant ($p < 0.05$) differences according to Duncan’s multiple range test.

2.8. Overexpression of DgMBF1 Enhanced the $K^+/Na^+$ Selectivity under Salt Stress

The contents of $K^+$ and $Na^+$ in leaves were measured, without obvious differences under normal conditions. Having been exposed to salinity, the $Na^+$ content of OE lines was significantly ($p < 0.05$) lower than that of WT, whereas the OE lines maintained a significantly ($p < 0.05$) higher level of $K^+$ content compared to WT plants. $Na^+$ content in OE lines increased approximately by 15.54- and 17.1-fold, respectively, while in WT, $Na^+$ content was 25.45-fold higher than day 0. $K^+$ content in OE lines revealed approximately 1.6- and 1.74-fold higher than day 0, respectively, while in WT decreased by 46%. In addition, the $K^+/Na^+$ ratio was significantly ($p < 0.05$) higher in OE lines than that in WT, which was about 4.95- and 4.46-fold greater compared with WT. (Figure 9a–c).

Moreover, some ion transporter genes, which served to regulate the $K^+/Na^+$ homeostasis, were found in chrysanthemum leaves, including the vacuolar $Na^+/H^+$ antiporter DgNHX, the plasma membrane $Na^+/H^+$ antiporter DgSOS, the potassium channel protein DgAKT and the high-affinity potassium ion transporter protein DgHAK. The expression levels of these genes were all statistically significantly ($p < 0.05$) up-regulated in OE lines than that in WT under salt stress (Figure 9d–g).

Overall, through regulating the expression of ion transporter genes, the OE lines excluded $Na^+$ and imported $K^+$ more effectively than did WT in response to salt stress.
DgMBF1 were acclaimed to be involved in stress response. Our results indicated that the functional di
ferences between the two branches have not been discovered yet. Both members
were induced by NaCl and might participate salt stress respondence. VvMBF1 was induced by

Phylogenetic analysis of DgMBF1 and MBF1s in other plant species indicated sharp differences
between MBF1c (plant group II) and MBF1a/b (MBF1a and MBF1b, plant group I) proteins. However,
the functional differences between the two branches have not been discovered yet. Both members
were acclaimed to be involved in stress response. Our results indicated that DgMBF1 belonged to the
group II (Figure 1b) [31,32]. The transcription of DgMBF1 in chrysanthemum leaves was dramatically
promoted under salinity (Figure 2b), indicating that DgMBF1 might participate salt stress respondence.

To further verify the function of DgMBF1, this gene was overexpressed in chrysanthemum. The
OE lines exhibited higher salt tolerance compared with WT. Salt stress inhibited the growth and
development of plants, reducing root length and decreasing fresh weight [30]. In our study, the
gradually increased salinity inhibited the root length and fresh weight of chrysanthemum. The growth
inhibition of OE lines was significantly (p < 0.05) lower than that of WT (Figure 5). These results
accorded with previous studies of MBF1 genes in other plant species. For instance, overexpression of

Figure 9. Overexpression of DgMBF1 enhances the K+/Na+ selectivity. (a) Relative Na+ content.
(b) Relative K+ contents. (c) K+/Na+ ratio. (d–g) Relative expression level of ion transporter genes in
WT and OE lines under salt stress. Na+ and K+ contents are relative to that of WT for 0 days. Data
represent means and standard errors of three replicates. The different letters above the columns indicate
significant (p < 0.05) differences according to Duncan’s multiple range test.

3. Discussion

Chrysanthemum belonged to the herbaceous perennial plants. Because of its high medicinal
and aesthetic value, the chrysanthemum-related industry thrived. However, salinity hampered the
production badly [19]. Therefore, transgenic technology would be applied to improve the salinity tolerance of
chrysanthemum. In this study, a multiprotein bridging factor 1 gene DgMBF1 from chrysanthemum
was isolated. The results showed that overexpression of DgMBF1 elevated the salt tolerance of
crysanthemum.

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inhibition of OE lines was significantly (p < 0.05) lower than that of WT (Figure 5). These results
accorded with previous studies of MBF1 genes in other plant species. For instance, overexpression of
*TaMBF1c* conferred heat tolerance in rice and yeast [33]. *VeMBF1* was induced by dehydration stress and ABA treatment, and overexpression of *VeMBF1* improved drought tolerance in *Arabidopsis* [34].

Excessive ROS could cause severe damage to plants. The antioxidant system of plants would reduce cell damage caused by ROS while maintain ROS balance [35,36]. The analyses showed that the contents of H$_2$O$_2$ and O$_2^-$ in OE lines were lower than that in WT, and the activities of ROS scavengers (SOD, POD, APX, CAT) were statistically significant (p < 0.05) higher than that of WT under salt stress (Figures 6 and 7a–d), which was consistent with the significant up-regulation of antioxidant related genes (*DgCuZnSOD, DgCAT* and *DgAPX*) (Figure 6e–g). Cu/ZnSOD was a metal enzyme that acted as a major superoxide scavenger. The overexpression of the *AliCu/ZnSOD* gene improved drought and salt tolerance in tobacco [37]. The highly activated CAT, as the plant’s cleaning agents, spared plants from the harm of reactive oxygen. The overexpression of *TsApx6* increased the survival rate of *Arabidopsis thaliana* under drought and high salinity stress [39].

The results above indicated that ROS accumulation was less in *DgMBF1*-overexpressed chrysanthemum than that in WT plants under salt stress. The overexpression of *DgMBF1* activated ROS-scavenging system, increasing the salinity tolerance of chrysanthemum.

Salt stress accounted for cytoplasmic water loss, which led to osmotic stress [40]. As two important intracellular osmotic regulatory factors: proline and SS functioned well in osmotic stress response [41]. In this study, the OE lines accumulated more contents of proline and SS than that of WT plants under salt stress (Figure 8a,b). Additionally, the expression of the related genes, such as *DgP5CS, Dg6PGDH* and *DgMDH*, which functioned in osmotic adjustment, were statistically significant (p < 0.05) up-regulated in OE lines (Figure 8c–e). This was in accord with the raised contents of proline and SS. P5CS was the rate-limiting enzyme, which functioned in botanical proline biosynthesis. Overexpression of P5CS genes could increase proline production so that confer salt tolerance in transgenic plants, such as tobacco, wheat, rice and potato [42–45].

These results indicated that by regulating the osmotic adjustment ability of transgenic chrysanthemum, *DgMBF1* could improve its salt tolerance.

By regulating K$^+$/Na$^+$ homeostasis, *DgMBF1* conferred salt tolerance in chrysanthemum. In terms of salt tolerance, the ability to maintain a relatively high cytoplasmic K$^+$/Na$^+$ ratio was considered a crucial determinant [46]. Ion transporter genes played a critical role in the transport of Na$^+$ in plant cells and vacuoles [47]. Salt tolerance levels were linked with the capacity to discharge Na$^+$ and to maintain a relative high K$^+$/Na$^+$ ratio in the cells [48]. The K$^+$/Na$^+$ ratio exhibited little differences in OE lines and WT plants under normal conditions, while the ratio in OE lines was significantly (p < 0.05) higher than that in WT under salt conditions (Figure 9c). These results were supported by significant differences (p < 0.05) in the expression levels of certain ion transporter genes (*DgNHX, DgSOS, DgAKT* and *DgHAK*) (Figure 9a,b,d–g). In *Arabidopsis thaliana*, ion homeostasis was largely mediated by the SOS signaling pathway [49]. NHX transporters involved in cytoplasmic detoxification by vacuolar Na$^+$ accumulation, osmotic adjustment by Na$^+$ or K$^+$ accumulation. It has been reported that *LeNHX* genes served as determinants of salt tolerance in tomato [50]. A plasma membrane Na$^+$/H$^+$ antiporter SOS1 influenced the export of Na$^+$. The overexpression of *AtSOS1* enhanced the salt tolerance [51]. For plants growing under salt stress, it was important to maintain a Na$^+$/K$^+$ ratio by favoring the accumulation of potassium over sodium [52]. AKT genes encoded the root K$^+$ uptake channels. The expression of a AKT1-type K$^+$ channel gene *PutAKT1* enhanced salt tolerance in *Arabidopsis* [53]. K$^+$ absorption and distribution in plants under salt conditions are mediated by potassium channels and transporters. The high-affinity K$^+$ transporter (HAK) family was among the major K$^+$ acquisition systems in plants [54]. Overexpression of *OsHAK5* increased the accumulation of K$^+$ and improved the salinity tolerance in tobacco [55]. *McHAKs* in *Mesembryanthemum crystallinum* were significantly induced by salt stress [56].
The obvious changes in the contents of K\(^+\) and Na\(^+\), and the expression levels of DgNHX, DgSOS, DgAKT and DgHAK in the chrysanthemum together indicated that DgMBF1 could enhance the salt tolerance of chrysanthemum by maintaining a higher K\(^+\)/Na\(^+\) ratio.

At the current stage, the salt-tolerant genetic engineering technology has made major breakthrough in chrysanthemum. A variety of transgenic chrysanthemums have been proven to have better tolerance to salt stress. Due to the transgenes belonging to different families, there was no comparison with salt tolerance among various transgenic lines. High-quality genes should be the target of further study. By comparing salt tolerance to screen out the best gene, the production and cultivation of chrysanthemums would be increased.

4. Materials and Methods

4.1. Plant Materials and Stress Treatment

The chrysanthemum cv. ‘Jinba’ were potted in a 1:1 mixture of peat and perlite, and grown in an incubator with a 16h photoperiod (200 \(\mu\)mol·m\(^{-2}\)·s\(^{-1}\) illumination), 23 ± 2 °C, 70% relative humidity. Seedlings at the 6–7 leaf stage were treated with 200 mM NaCl solutions to create salt stress. Seedlings were sampled at 0, 1, 3, 6, 12 and 24 h time points from all the treatments and stored at −80 °C for RNA extraction.

4.2. DgMBF1 Clone and Sequence Analyses

Total RNA was extracted from chrysanthemum leaves by the TRIzol reagent (Mylab, Beijing, China), the first-strand cDNA was synthesized with the PrimeScript™RT reagent kit (Takara, Beijing, China). A complete DgMBF1 ORF which prepared for the inclusion of an XbaI and a SacI cloning sites was amplified from this cDNA template using the primer pair (DgMBF1-F/R) designed from the putative 5’ and 3’ untranslated region. The PCR products were purified by agarose gel and cloned into pMD18-T vector (Takara, Beijing, China) for sequencing.

4.3. Sequence Alignment and Phylogenetic Analysis

The BLAST online tool was used to search homologous protein sequences in the NCBI protein database. Sequences with over 90% coverage were selected. Sequence alignment of DgMBF1 was performed using ClustalX. The phylogenetic tree was constructed through MEGA5.

4.4. Expression Vector Constructs and Chrysanthemum Transformations

Initially, the complete ORF of DgMBF1 was cleaved by restriction with XbaI and SacI and inserted into XbaI and SacI digested the plant expression vector pCAMBIA2300 to produce pCAMBIA2300-DgMBF1. This pCAMBIA2300-DgMBF1 vector was driven by the CaMV 35S promoter and transmuted into LB4404 strain of Agrobacterium tumefaciens and following the protocol documented by An et al. [57].

Middle and upper leaves of the chrysanthemum “Jinba” were cut into a square centimeter pieces and cultured on Murashige and skoog (MS) medium. These pieces were used as transformation receptors, pre-cultured for 3 days, the single colonies of agrobacterium LB4404 of pCAMBIA2300-DgMBF1 were transformed into Agrobacterium liquid (cultured at 28 °C until the OD\(_{600}\) was 0.5) infecting leaf discs for 10 min, co-cultured for 3 days, delay-cultured for 3 days, the specific method according to Cui et al. [58]. Transformants were initially selected through DNA detection, chrysanthemum genomic DNA was isolated from medium-sized leaves using a DNA extraction kit (Takara, Beijing, China), primers used for DNA detection are listed in Table 1.
Table 1. Primers used in this study.

| Primers used for cloning of DgMBF1 | Forward Primers | Reverse Primers |
|-------------------------------------|-----------------|-----------------|
| DgMBF1 DNA detection                | TCCAGACCTCAACTCTA | ACAAATCGACACAAATAAAAA |
| EF1α                                | GAGTCAAAGATTCAAATAGGAC | ACAAAATCGACACAAATAAAAA |

| Primers used for qRT-PCR            | Forward Primers | Reverse Primers |
|-------------------------------------|-----------------|-----------------|
| DgMBF1                              | TGCCGCAAGAGGACAGG | TGGGACAGACGATCAG |
| EF1α                                | TTGTGATCTCTGACCTGG | GGCTTTCTGCAGACGAG |
| DgCuZnSOD                           | CCATTGCTGACAGGATAC | GACCTCTGCTCCAGACGAG |
| DgCAT                               | TACGACATCAGACAGG | GAGCTTTCTGCAGACGAG |
| DgAPX                               | GTCGTCGCTCCTTGGGGAT | GGCTTTCTGCAGACGAG |
| DgP5CS                              | TTTCTGCAGACAGG | GACCTCTGCTCCAGACGAG |
| DgMDH                               | CGAGACTTCGTCTCCCG | GCTTTCTGCAGACGAG |
| DgDHX                               | TGCTGCTAGATGATA | GCTTTCTGCAGACGAG |
| DgAKT                               | AGCTGCAACTCACGAG | GCTTTCTGCAGACGAG |
| DgDHX                               | TGCAGAGACTGAGCAGG | GCTTTCTGCAGACGAG |

4.5. Gene Expression Levels Analysis

The expression level of DgMBF1 was detected by qRT-PCR using the SsoFast EvaGreen supermix (Bio-Rad, Hercules, CA, USA) and Bio-Rad CFX96TM detection system. The settings for qRT-PCR were as follows, 95 °C 30 s, 95 °C 15 s, 60 °C 30 s, circulating for 40 times. The analyzing approach of $2^{-\Delta\Delta Ct}$ was applied in the experiment, using EF1α as the reference gene [59]. The primers used in qRT-PCR are listed in Table 1.

4.6. Determination of Salt Tolerance

Six transgenic strains were eventually obtained, and two independent OE lines (OE-3, OE-34) were included. The OE lines exhibited relatively high expression. These samples were reproduced through asexual propagation for salt treatment. The root, stem, leaf and flower tissues of chrysanthemum under normal condition were collected for tissue-specific expression analysis of DgMBF1 gene.

Totally, each line contained 120 seedlings. To avoid salt shock, the method according to Chen et al. [60] was used to irrigate the chrysanthemum seedlings with an increasing concentration of NaCl solutions. Chrysanthemum seedlings at 6–7 leaves stage were irrigated with NaCl solutions: 100 mM on day 1–5, 200 mM on day 6–10, 400 mM on day 11–15, and sampled at 0, 5, 10 and 15 days.

One third of these seedlings were sampled for staining experiments, root length and fresh weight measurements. The roots were rinsed clean. The whole seedling was put into a container of known weight. The fresh weight was measured with an analytical balance. The ascending fourth and fifth leaves were cut and stained immediately. The length of 30 cm was drawn and the shallow water layer was set on the table. Stretching the wet root with the mark, the length of root was calculated. The whole root length was the sum of each sample.

One third of seedlings were applied to statistical survival. When the salt stress treatment process was completed, the roots were rinsed with deionized water and transplanted in a new culture medium. The survival rate was counted after two-week recovery. The ascending fourth and fifth leaves collected on last one third of seedlings were used for qRT-PCR and physiological experiments.

4.7. Determination of Physiological Indexes of Chrysanthemum under Salt Stress

$\text{H}_2\text{O}_2$, proline and SS contents, APX, POD, SOD and CAT activities were measured with the Nanjing Jiancheng test kit, the content of $\text{O}_2^-$ was measured with the Suzhou Keming test kit. Both procedures were performed strictly following the instructions. In situ accumulation of $\text{H}_2\text{O}_2$ and $\text{O}_2^-$ were measured through histochemical staining, using DAB and NBT, respectively, following Zhao et al [28]. The Na$^+$ and K$^+$ concentrations were measured following An et al. [26].
4.8. Expression of Stress-Related Genes

Total RNA of OE lines and WT plants were isolated and converted to cDNA as described above. The expression of stress-related genes in chrysanthemum was measured by qRT-PCR. *DgCuZnSOD, DgCAT, DgAPX, DgP5CS, Dg6PGDH, DgMDH, DgNHX, DgSOS, DgAKT* and *DgHAK* were monitored, and *EF1α* as a reference. The primers used in qRT-PCR are listed in Table 1.

4.9. Statistical Analysis

All experiments were conducted in three biological repetitions. All data were analyzed using SPSS version 20.0 program, the significance discriminate analysis (*p* < 0.05) was performed according to Duncan’s multiple range test.

5. Conclusions

To conclude, our study identified a multiprotein bridging factor 1 *DgMBF1* as a salt-tolerant positive regulatory gene. *DgMBF1* was induced by salinity. The overexpressed *DgMBF1* enhanced the salt tolerance in transgenic chrysanthemum through maintaining a higher K⁺/Na⁺ ratio, greater activities of antioxidant enzymes, higher accumulation of osmotic regulatory factors. Therefore, *DgMBF1* could be served as a candidate gene for salt-tolerant plant breeding. The next step was to conduct an in-depth study of the downstream target genes of *DgMBF1* to understand its deeper molecular mechanisms in salt stress response.

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Abbreviations

| Abbreviation | Definition |
|--------------|------------|
| APX          | superoxide Anion |
| CaMV         | cauliflower mosaic virus |
| CAT          | catalase |
| DAB          | diaminobenzidine |
| HTH          | helix-turn-helix |
| H₂O₂         | hydrogen peroxide |
| MBF1         | multiprotein bridging factor 1 |
| NBT          | nitro blue tetrazolium |
| O₂⁻          | superoxide anion |
| OE           | overexpressed |
| ORF          | open reading frame |
| PCR          | polymerase chain reaction |
| POD          | peroxidase |
| qRT-PCR      | quantitative real-time PCR |
| ROS          | reactive oxygen species |
| SOD          | superoxide |
| SS           | soluble sugar |
| WT           | wild type |
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