



Monitoring Response of a Few bZip Transcription Factors in Response to Osmotic Stress in Sunflower

Zeinab Esmailzadeh Mianlengh¹, Masood Soltani Najafabadi^{2*}, Abbas Saidi^{1*}, Hossein Askari¹

¹ Department of Plant Sciences and Biotechnology, Faculty of Life Sciences and Biotechnology, Shahid Beheshti University, G.C, Tehran, Iran.

² Genetic Research Department, and Iranian National Plant Gene Bank, Seed and Plant Improvement Institute (SPII), Agricultural Research, Education, and Extension Organization (AREEO), Karaj, Iran

*Corresponding authors: Masood Soltani Najafabadi, Genetic Research Department, and Iranian National Plant Gene Bank, Seed and Plant Improvement Institute (SPII), Agricultural Research, Education, and Extension Organization (AREEO), Karaj, Iran Tel/Fax: +98-26-32716923/+98-26-32704851 E-mail: M.soltani@spii.ir, and Abbas Saidi, Department of Plant Sciences and Biotechnology, Faculty of Life Sciences and Biotechnology, Shahid Beheshti University, G.C, Tehran, Iran. Tel/Fax: +98-21-22431964, E-mail: abbas.saidi@gmail.com

Received: 20 Dec. 2015; Revised: 20 Sep. 2017; Accepted: 14 Oct. 2017; Published online: 15 May 2018

Background: Sunflower (*Helianthus annuus* L.) is one of the important vegetable oil supplies in the world and in Iran, as well. It is classified as a drought semi-tolerant crop; however, its yield is adversely affected by drought stress. Understanding the initial events in sensing stress and the related physiologic and biochemical events thereafter, is crucial in designing drought stress breeding programs. Transcription factors are master molecules directly involved in the plant responses under drought stress, from signal perception and transduction to the regulation of physiologic processes.

Objective: The expression pattern of some bZip transcription factors in response to osmotic stress was investigated in sunflower.

Material and Methods: Employing real-time PCR to monitor, the response of 10 bZIP transcription factors was performed under different osmotic stress conditions including -0.3, 0.9, and 1.2 MPa. Whole seedling was sampled at 6, 12, and 24 h after the osmotic condition application.

Results: Exposure to osmotic potential of 0.9 MPa for 24 h caused a reduction in the fresh weight of the seedling. Among the evaluated genes, eight genes, *bz-497*, *bz-502*, *bz-485*, *bz-499*, *bz-492*, *bz-504*, *bz-505*, and *bz-509* appeared as the osmotic stress responsive transcription factor. Changes in the expression of the genes under 0.3 MPa was observed for four genes. Most of the osmotic responsive genes appeared to be up-regulated. Most of responsiveness in the gene expression was happened under 0.9 MPa of the osmotic stress which is corresponding to fresh weight reduction in the seedlings. Among the investigated genes, two genes was identified to have possible roles in sensitive response of sunflower against drought stress.

Conclusions: It was a focus to have systemic view on the complex response of the plant to abiotic stress, and avoidance of the single gene analysis. Also, the importance of molecular data in molecular breeding procedures toward achievement of the stress tolerant lines was highlighted.

Keywords: BZip, Osmotic stress, Patchy pattern, Sunflower, Transcription factor

1. Background

Plants are continuously exposed to the constraints which prevent them from reaching their maximum productivity. Various abiotic stresses, especially drought stress are the most important of these restrictive factors for the crops growth and productivity worldwide. The earth is

warming and dry areas are increasing (1), on the other hand, food requirement due to growing population has been a major concern for all nations (2), thus drought stress and developing drought tolerant crops has been considered by researchers to a significant extent. The mechanisms of the response to the drought stress can

be evaluated at many different levels, from the whole plant to the molecular level (3). The physiological and molecular aspects of the plant response in drought condition has been broadly been studied (4). Some of these responses include stomatal closure (5), repression of the cell growth and photosynthesis (3), activation of the respiration (6), and accumulation of osmolytes and proteins (7). It is well known that all these changes are due to the changes in the gene expression level (8-10). The extensive investigation on the molecular aspect of the plant responses toward drought and osmotic stress which were gained via high-throughput methods have revealed that two classes of the genes are responsive in this case (10, 11): i. Functional genes which include genes that code proteins which function directly in abiotic stress tolerance such as chaperones, late embryogenesis abundant (LEA) proteins, osmotin, antifreeze proteins, mRNA-binding proteins, key enzymes for osmolyte biosynthesis, water channel proteins, sugar and proline transporters, detoxification enzymes, and various proteases and ii. the regulatory genes (i.e., genes coding for the proteins involved in further regulation of the signal transduction and stress-responsive gene expression). The second category include various transcription factors (TFs), protein kinases, and protein phosphatases.

Transcription factors are master proteins which regulate downstream genes upon recognition of specific sites on their regulatory regions. These low abundant proteins, based on presence of their conserved DNA-binding domains, are classified into several families, including MYB, MYC, WRKY, NAC, bZip, etc (12, 13).

bZip family of TFs contains members which possess two structural features, a basic region which binds to DNA and a leucine zipper dimerization motif. Plant bZip proteins preferentially bind to DNA sequences with an ACGT core. Several evidences indicate the essential role of bZip TFs in various plant developmental processes and adaptation responses against abiotic (14-16) or biotic stresses such as pathogen defense (17), light, seed maturation, and flower development (17). Their overexpression has been shown to have significant effect on the increased osmotic stress (18). As most responses in the plants under various conditions are governed by TFs, monitoring expression behaviors of the TFs could give us a better understanding of the processes running inside plants, which lead to the adaptive responses (4). The best way of gaining this understanding is to have the whole view on the system which is accessible through TF profiling (19). Although the bZIP TF family has been studied well among different plant species

(20-22), a comprehensive profiling for expression of *bZIP* gene family in response to the osmotic stress is still lacking in several important crop plants, including sunflower. Having information over this can be very helpful for future breeding programs through genetic engineering. The best technology to perform TF profiling was proved to be real-time PCR (23).

Sunflower (*Helianthus annuus* L.), is an important oil crop that produces a major part of the vegetable oil worldwide. Sunflower seed oil has high nutritional property and is composed of unsaturated fatty acids (90%) (24). This annual crop is semi-sensitive to drought, however reduction in sunflower yield due to drought stress has been reported worldwide (25, 26). Despite the importance of the crop in supplying edible oil, several factors such as large genome size (3.5 Megabase (27)) and a probably duplicated and complex, (28, 29) lead to limited information available at molecular level for the plant.

2. Objective

As there was not data available on the functional behavior of bZipTFs under osmotic stress in sunflower and the need for a systemic assessment on its function, in this paper, we tried to take a survey on this subject. To this end, we tried to find data on bZipTFs and measure the responses of the TFs to the different osmotic potential during time courses. The results have indicated the presence of a patchy pattern for expression of the evaluated TFs. This might provide another prove to the multi-level of regulation in response to suboptimal condition.

3. Materials and Methods

3.1. Plant Materials and Stress Treatments

Seeds of sunflower line AF81-112, a drought sensitive line (30), were supplied from Seed and Plant Improvement Institute, Iran. The seeds were sterilized before and after de-hulling using 50% sodium hypochlorite for 10 min followed by 70% ethanol for 5 min, and germinated on water agar medium for two days under 16 h of light, 24-25°C temperature. After two days, when seedlings were emerged, seedlings were transferred to liquid B5 medium and kept over a shaker-incubator with 60 rpm for five days. Osmotic stress was then applied by adding specific amount of polyethylene glycol6000 (PEG6000) to the B5 medium to provide osmotic potentials of 0, 0.3, 0.9 and 1.2 MPa. The amount of PEG to be added to gain the above osmotic potential was calculated using the equation of Michel and Kaufmann (31) and the osmotic potential

of the media was measured and confirmed using a thermocouple osmometer (Wescor Vapor Pressure Osmometer Model 5100C, Wescor Inc., Logan, Utah, USA) according to the manufacturer's instruction. After six, 12, and 24 h of stress application, the whole seedlings (including root and shoot) were washed with sterilized water, weighed, and deep frozen in the liquid nitrogen for RNA extraction. For each treatment (i.e., combination of the osmotic potentials and time courses) seven seedlings were considered. The experiment was repeated two more times. As the experiment was conducted in a completely randomized design with three replications. The fresh weight of samples was subjected to ANOVA and the mean comparisons was done using Duncan's multiple range test.

3.2. RNA Extraction and cDNA Synthesis

The seedlings for each treatment were pooled and grinded to a fine powder in the liquid nitrogen. The total RNA was extracted from the samples using RNX-plus Kit (CinnaGen) according to the manufacturer's protocol. The integrity and quantity of the extracted RNA was evaluated by loading on 1% agarose gel and spectrophotometer, respectively. The residuals of the genomic DNA were removed by treating RNA with RNase-free *DNaseI* (Fermentas, Germany) according to the manufacturer's instructions. The absence of contamination with the genomic DNA was subsequently confirmed by real-time PCR using primer pairs for tubulin and elongation factor 1 (Table 1).

Reverse transcriptase (RT) reactions were performed on 3 µg of the total RNA with oligo-dT primer and CinnaGen reverse transcriptase kit, according to the manufacturer's instructions. The efficiency and concentration of the synthesized cDNA was evaluated using Real-Time PCR amplification of the two reference genes encoding tubulin and elongation factor 1 (Table 1).

3.3. Real-Time PCR Analysis

PCR reactions were conducted using a StepOne™ Real-Time PCR system (Applied Biosystems), employing SYBR Green to monitor dsDNA synthesis. Reactions with a final volume of 10 µL contained 1 µL of template (cDNA or total RNA), 200 nM of each gene or sequence-specific primer (4 µL of the mixed 0.5 mM forward and reverse primers, respectively), and 5 µL of the SYBR Green master mix (Takara, Japan). The following standard thermal profile, as recommended by the manufacturer, was used for all Real-Time-PCR reactions: 50 °C for 2 min, 95 °C for 10 min; 40 cycles of the 95 °C for 15 s, and 60 °C for 1 min.

After 40 cycles, the specificity of the amplifications was checked by heating from 60 °C to 95 °C with a ramp speed of 1.9 °C min⁻¹, resulting in melting curves. The threshold cycle, C_T , which is the cycle number at which SYBR Green fluorescence in a PCR reaction reaches to an arbitrarily defined threshold value during the exponential phase of DNA amplification, were used for expression analysis. For each treatment, arithmetic mean of C_T for *Tubulin* and *Elongation Factor 1 (EF1)* genes (C_{TM}) was used for normalization of all C_T for genes (C_{Tg}) so that $\Delta CT = C_{Tg} - C_{TM}$. For comparative expression analysis, $2^{-\Delta\Delta CT}$ method (32) was used, where $\Delta\Delta CT$ is subtraction of ΔCT for stress condition from ΔCT for control condition. The $2^{-\Delta\Delta CT}$ was considered as fold change (FC).

The pattern of gene expression was obtained on Log₂FC of all genes using Multiple Experiment Viewer (MeV) software version 4.1.10. In this analysis Hierarchical Clustering (HCL) was performed on both treatments and genes using average linkage method and Manhattan distance.

3.4. Bioinformatics Analysis

The partial genomic and proteins sequences of TFs family bZip were iterated from sunflower TFs site (http://planttfdb_v1.cbi.pku.edu.cn:9010/web/index.php?sp=ha). As the sequences were not completed, we first tried to find ORF sequences which their translation were more closer to the partial protein sequences presented in the site. To confirm that the sequences belong to bZip family, the protein sequences were then blasted (33) against plant database (NCBI, <http://www.ncbi.nlm.nih.gov/>). In order to design specific primers for each member of the bZip family, the bZip domain of the proteins were identified using tools in Prositeexpasy site (<http://prosite.expasy.org/>). The primers for real-time PCR were designed based on the non-conserved sequence of the genes using Oligo7 software (online accessibility). The criteria for primer designing were: predicted melting temperatures (Tm) of 58-62 °C, primer lengths of 19-25 nucleotides, guanine-cytosine (CG) percent of 45-55%, and PCR amplicon lengths of 60-150 base pairs (bp). The primer sequences and their characteristics are presented in Table 1. Primers for the internal control genes including *Tubulin* and *EF1* were selected according to Fernandez *et al* (2011, (34)), the sequences of which are included in the Table 1.

4. Results

4.1. Stress Experiment

The increase in the osmotic potential of the media was

resulted in the decrease in the sunflower seedling fresh weights. However, the decreases were not so steep, but rather a gradual behavior was observed (**Fig. 1**). For each osmotic potential level, effect of duration of the osmotic potential imposition was not significant. This may indicate that when the seedlings sense a specific ambient osmotic potential, they may modify their internal osmotic statuses and/or trigger a chronic osmotic potential adjustment system which prevent desiccation during time.

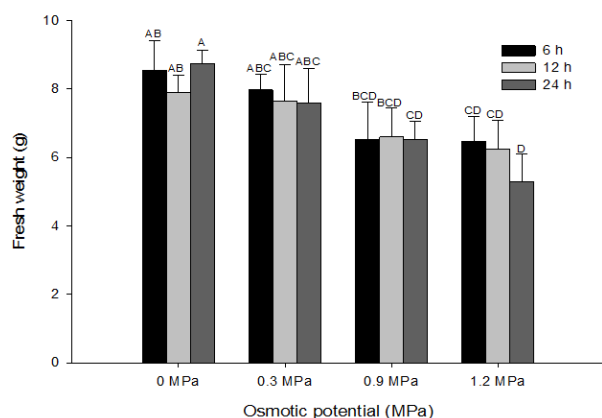


Figure 1. Responses of fresh weight of sunflower inbred line AF81-112 to increases in osmotic potential of the growth media. The seedlings were grown under each osmotic potential for 6, 12, and 24 h. The experiment was repeated three times and average fresh mean over the three biological replications is indicated. Columns having at least one letter in common are not significantly different ($\alpha \leq 0.5$). Data are represented as means \pm SD.

Under optimum condition (0 MPa), although non-significant, the effect of time on increasing fresh weight (an indication of the growth) was observed (fresh weight at 24 h vs 12 h). Increasing osmotic potential of the medium up to 0.3 MPa did not affect the fresh weights.

The first significant effect of the low osmotic potential on fresh weight as compared to the optimum condition was observed after a prolonged (24 h) application of 0.9 MPa osmotic potential. The same effect was observed when seedlings were grown for 24 h on 1.2 MPa media. In fact in this case, the reduction in fresh weight was significant compared to 0.3 MPa media. In both cases, the effect of osmotic potential of the media on fresh weight was accelerated by time. This may indicate a probable accumulation effect of damaging factors under hypo-osmotic potentials. This cumulative effect was not detectable under 0.3 MPa potential. Thus, in this experiment, 24 h incubation on osmotic potential of 0.9 MPa was the start point for seedlings to show their influences by the stress.

4.2. Bioinformatics Survey

In the sunflower transcription factor database, gene sequences for 33 accessions have been deposited as bZip transcription factors. Running ORF finding procedure on the gene sequences showed that four genes had no ORF greater than 50 bp in length. Running protein BLAST for the rest of 29 genes, we found that there is less than 25% homology for 19 genes. As a result, for out of 33 accessions deposited as bZip transcription factor, expression analysis was conducted for 10 genes (**Table 1**).

4.3. Expression Analysis

Melting curve analysis revealed that the primer pairs for each gene were able to amplify a single and therefore specific amplicon (Data are not shown). Also, no track of primer dimer was observed for the primers. Thus, expression analysis using the designed primers was considered to be valid.

In expression analysis, we defined genes to be osmotic stress responsive if their expression fold change (FC) was greater than 2 or less than 0.5 relative to the control condition. The two limits were considered for up- and down-regulated genes, respectively, and the genes with expression values in between as non-responsive genes to the osmotic stress. In each osmotic condition, the response of genes were detected at early (6 h), mid (12 h), and late (24 h) incubation times.

The hierarchical cluster analysis showed the presence of different waves in gene expression among the 10 genes and samples (**Fig. 2**). Based on the results, gene *bz-504*, *bz-505*, *bz-492*, and *bz-485* were appeared as genes with distinct expression patterns over the samples. While genes *bz-503*, *bz-499*, and *bz-487* which show nearly the same expression behaviours, were grouped in the same cluster, while, *bz-502*, *bz-509*, and *bz-497* genes which showed a different expression pattern appeared as a distinct group. Interestingly, a unique expression pattern of all genes was observed in 0.9 MPa for 12 h. In this analysis, genes with negative and positive Log_2FC were considered as down-regulated- and up-regulated ones. This analysis which is based on the average value of the gene expression gives us an overview on the scattered pattern of the gene expression among the different levels of the stress and time points, and thus, details of information on the behaviour of the genes is not reachable. This could be true, because most of TF gene expressions are suffered from the noises in the expression (35) and so the wide confidence intervals for their expression values make interpretations on the average values as hard and not precise.

The genes of *bz-503*, *bz-487*, and *bz-499* were

Table 1. Primers used for real-time PCR analysis.

Code	Gene name*	Primer	5' to 3'	TM (°C)	GC%	Amplicon size (bp)
bz-485:	PTHa00485.1	Forward	AACAACAGCCGCCTCCATAC	56.2	55	64
		Reverse	CCATAACCACCAGCAACACC	54.8	55	
bz-487	PTHa00487.1	Forward	GAAATCTCTGGACGAGAAGAAACG	45.8	57.2	80
		Reverse	TATATCTGCCGCCTGATG	55	57	
bz-492	PTHa00492.1	Forward	GAGCAGTATGGATGGGGAAGTG	54.6	57	80
		Reverse	GCCGACAGACATAAAGTCAAGT	54.6	56.2	
bz-497	PTHa00497.1	Forward	CTAAATATGGGGATGGACTTGTGG	45.8	57.3	63
		Reverse	CGGTCGCATTTTCATGGTTC	50	57.4	
bz-499	PTHa00499.1	Forward	CGAGCAATGGTTACTCGATAGC	47.8	56.8	85
		Reverse	GGTGGGTTACAAGTATGCGTATGG	50	58.2	
bz-502	PTHa00502.1	Forward	GCAAGAAACGCAGTGGATAGG	52.4	56.1	103
		Reverse	ACCAGGAACCGTAACCGATG	55	55.9	
bz-503	PTHa00503.1	Forward	AAGATCACAACCGCCGAATC	51	56.1	147
		Reverse	AGACATCAGCCTTAGCAGCATC	50	55	
bz-504	PTHa00504.1	Forward	CGGAAAGAAAAATGTGGATGTG	40.9	55.8	94
		Reverse	ATGTTGCTGCGGTATCGTTG	50	55.8	
bz-505	PTHa00505.1	Forward	GGGATACCGAAGACAAAAGC	55	54.8	70
		Reverse	TGGCATCAGAGACCACAAGC	55	55.7	
bz-509	PTHa00509.1	Forward	GATACATCGTCGGTTTCGCCTT	50	58.9	64
		Reverse	CGTTCTTCTACCCCTTACAATCC	50	58.2	
-	<i>Tubulin</i>	Forward	CGAGAGCAACATGAATGATCTG	45.5	45.5	93
		Reverse	CCTCTTCTCTCTTCTTCTCA	52.2	56.8	
-	<i>EF1</i>	Forward	AGGCGAGGTATGATGAAATTGTCA	41.7	58.4	200
		Reverse	GTCTCTTGGGCTCATTGATTTGGT	45.8	59.2	

*. The accessions are presented in http://plantfdb_v1.cbi.pku.edu.cn:9010/web/index.php?sp=ha.

appeared to be as the osmotic stress non-responsive or noisy genes (**Fig 3.**, also the pannel inside). It means that under different osmotic stress levels and time points, either their FC fall in the non-stress responsive region (FC between 0.5 and 2) or noise in gene expression, which is shown as the high standard deviation on gene expression, is observed.

There was no detectable expression for *bz-485* in the control condition and also osmotic stress in all three levels in 6 h (**Fig 4.**, and the pannel inside). Thus for, the gene is *de novo* induced upon osmotic stress imposition longer than 6 h. In the 0.3 MPa, induction of the gene was observed only at the late time (24 h). Under osmotic condition of 0.9 MPa and in time points

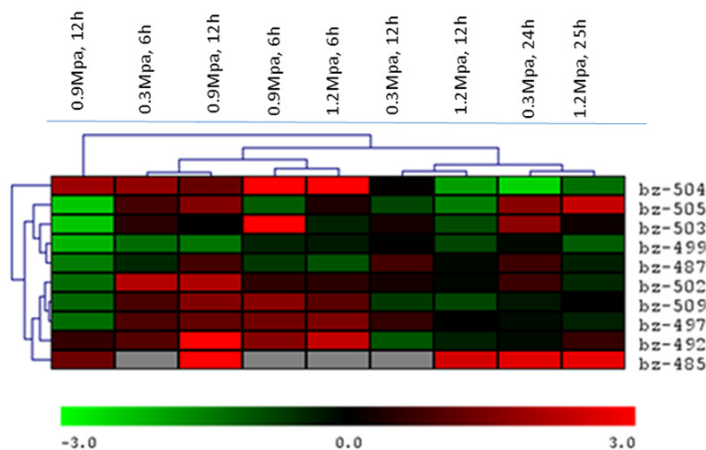


Figure 2. Hierarchical cluster analysis for the relative expression of the 10 genes belonging to the bZip TF of the sunflower seedling. The mean values for the three independent biological replications were subjected to average linkage method and Manhattan distance hierarchical clustering. Sunflower seedlings were subjected to 0.3, 0.9 and 1.2 MPa for 6T 12, and 24 h. The color saturation reflects the magnitude of the averaged \log_2FC . Expression values higher and lower than those of the control are shown in red and green, respectively. The vertical dendrogram (left) indicates the relationship among the TF genes regarding their expression patterns. The horizontal dendrogram (top) indicates the relationship among the treatments with respect to the expression of the bZipTFs. The color scale (bottom) indicates the color assigned to each \log_2FC .

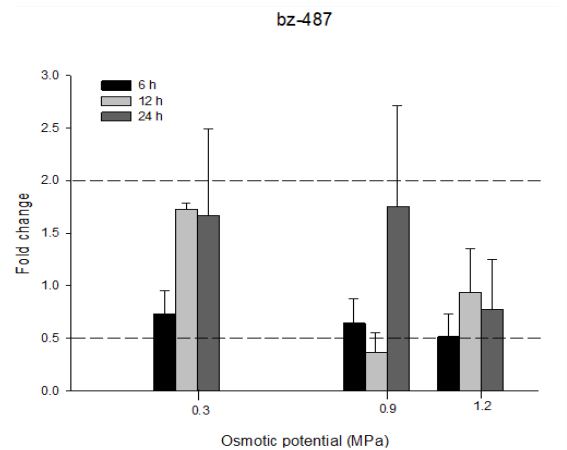
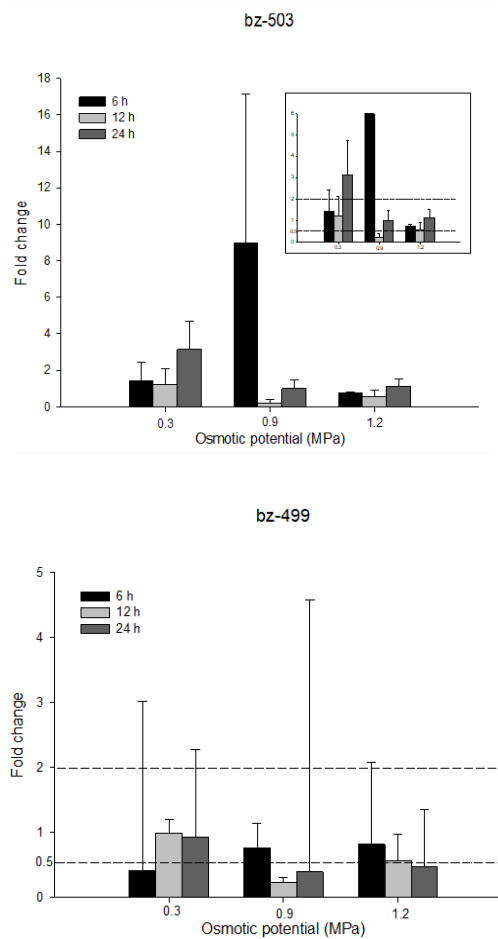
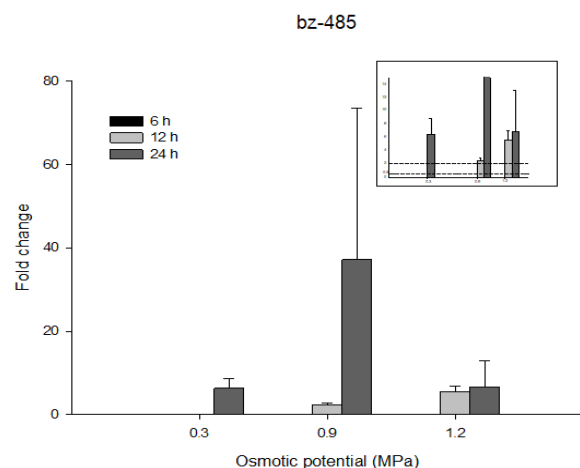


Figure 3. Relative expression of three bZip transcription factors, *bz-503*, *bz-487*, and *bz-499* in response to different osmotic potentials (0.3, 0.9, and 1.2 MPa) at various time course of osmotic potential application. *bz-503* and *bz-487* did not responded to osmotic potential. Gene *bz-499* mostly showed noisy pattern, but down-regulated under 0.9 MPa in 12 h. The time courses included are represented as early (6 h), mid (12 h), and late (24 h) course. The vertical axis is the relative expression of each gene under the osmotic potentials relative to control condition. The dashed horizontal lines define the threshold for up-regulation (upper limit line) and down-regulation (lower limit line). To show the threshold lines for up- and down-regulation, and because of the scale of Y axis, for gene *bz-503*, an inset panel with increased scale is included. Data are represented as means \pm SD of three replicates.

of 12 and 24 h, the gene responded non-significantly and noisily to the stress, respectively. In a more severe osmotic stress, 12 h imposition of the stress resulted in a prominent over expression of the gene, and a noisy behaviour upon longer imposition. It seems that the response of the gene to the reduction of osmotic stress depends on the stress severity and a threshold of the stress signal level required for its induction.

Figure 4. Relative expression of one bZip transcription factor (*bz-485*) showing *de novo* induction upon osmotic potential in response to different osmotic potentials (0.3, 0.9, and 1.2 MPa) at various time course of osmotic potential application. Lack of data on some treatments have caused by no gene expression. The time courses included are represented as early (6 h), mid (12 h), and late (24 h) course. For the matter of scaling, inside panel is included. The vertical axis is the relative expression of each gene under the osmotic potentials relative to control condition. The dashed horizontal lines define the threshold for up-regulation (upper limit line) and down-regulation (lower limit line). To show the threshold lines for up- and down-regulation, and because of the scale of Y axis, an inset panel with increased scale is included. Data are represented as means \pm SD of three replicates.

The genes *bz-492* and *bz-509* showed approximately the same behaviour in response to the different osmotic condition as well as time courses (**Fig. 5**). A bell-shaped pattern in the gene expression was observed for the genes. This means gene expression from a non responding (for *bz-509* for the three time points under 0.3 MPa and *bz-492* for 6 h and 24 h) or down regulation (for *bz-492* in 0.3 MPa for 12 h) was raised



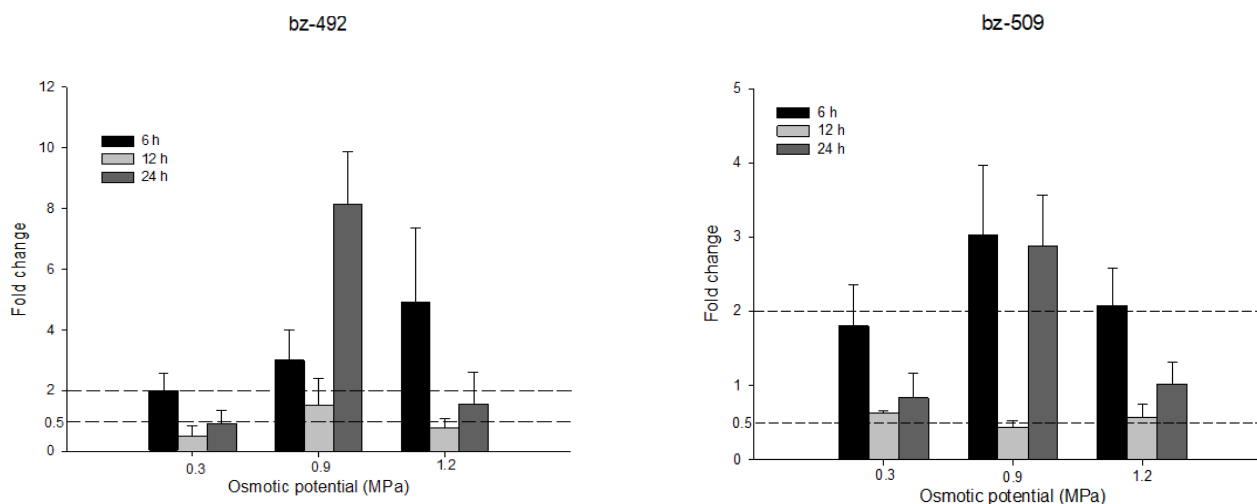


Figure 5. Relative expression of two bZip transcription factors, *bz-492* and *bz-509* showing a bell-shaped pattern upon osmotic potential in response to different osmotic potentials (0.3, 0.9, and 1.2 MPa) at various time course of osmotic potential application. The time courses included are represented as early (6 h), mid (12 h), and late (24 h) course. The vertical axis is the relative expression of each gene under the osmotic potentials relative to control condition. The dashed horizontal lines define the threshold for up-regulation (upper limit line) and down-regulation (lower limit line). Data are represented as means \pm SD of three replicates.

to up-regulation under 0.9 MPa in the 6 h and 24 h, but not-responding in 12 h. Their respected FC was reduced under 1.2 MPa osmotic potential to non-responding values.

We may propose an abundant and shortage effect model for the signals triggered by osmotic potential which affect on gene *bz-509*. Under this model, the amount of signal under 0.3 MPa, even in 24 h is not enough to trigger gene expression. Nevertheless, when severity of the stress reached to 0.9 MPa the amount of signal reached above a threshold and starts to trigger gene expression. The product of the gene (i.e., *bz-509* protein) may be high enough to prevent more gene expression, and through a negative feedback, could suppress its expression *per se*. When osmotic potential prolonged for more time (24 h), demands for the *bz-509* protein triggered its expression again. When no significant FC for *bz-506* under 1.2 MPa is observed, we may conclude that under severer osmotic stress, plant undertakes another signaling and regulatory pathway.

The *bz-505* TF was appeared to be the late responsive to the three osmotic potentials (Fig. 6). Except for a down-regulation under 0.9 MPa in 12 h, the gene was up-regulated in 24 h for all the osmotic potentials. The expression pattern for the gene under 0.9 MPa follows a non-linear behavior, that is, in early time point there is not a significant for this TC's relative expression followed by a down-regulation and an up-regulation in late time point, respectively. This behavior could be attributed to the cumulative stress signaling in which in

the early time point the amount of signal is not enough to be effective for changing gene expression, but under a prolonged stress condition for (i.e., more than 6 h) has negative effect on the gene expression. Nevertheless, upon prolonged stress, the suppressive effect might be withdrawn in a competition between suppressing and inductive effectors, and hence, causing an up-regulation event.

It could be concluded that the production of *bz-505* protein may serve as a counter-osmotic or sensor for osmotic severity which is activated upon exceeding of the respected signals above a biological-defined threshold.

bz-504 is an early responding gene, which is up-regulated in 6 h and then returned to the same level of expression of the controls under all examined osmotic potentials (Fig 6). Interestingly the up-regulation values showed an exponential behavior as osmotic potential was increased. In the other word, the relative rises in the *bz-504* transcript were observed to be small in 0.3 and 0.9 MPa followed by a huge increase in 1.2 MPa. Thus, we may conclude that the gene serves as an alarm for the early induction of osmotic stress to the biological system, triggers the responsive downstream mechanisms, and returns back to its background expression thereafter.

Bz-502 was up-regulated when seedlings were subjected to the 0.3 MPa for 6 h and returned to the background expression level thereafter (Fig. 7). Nevertheless, without responding to early and middle

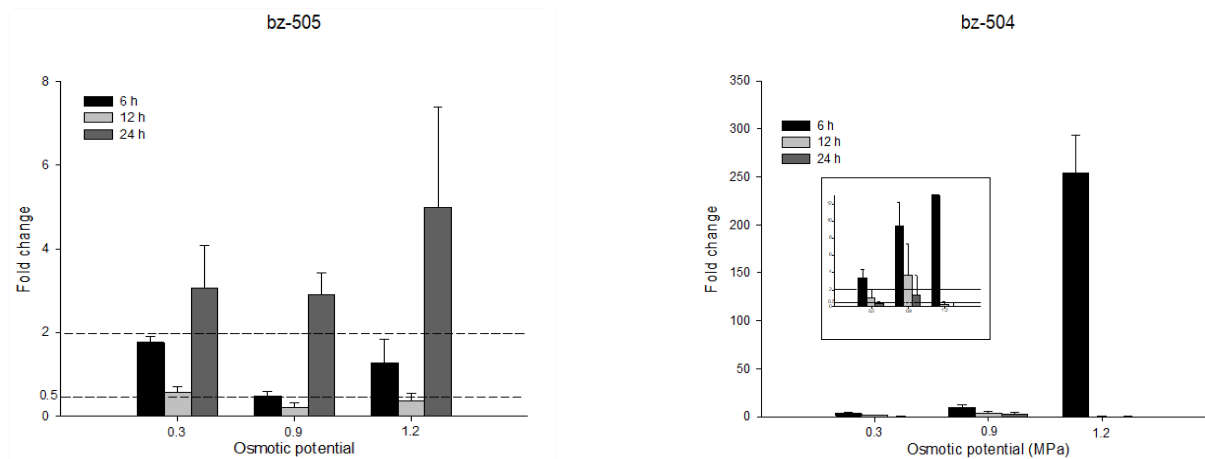


Figure 6. Relative expression of two bZip transcription factors, *bz-505* with late up-regulation and *bz-504* with early up-regulating expression patterns in response to different osmotic potentials (0.3, 0.9, and 1.2 MPa) at various time course of osmotic potential application. For the matter of scaling, inside panel in (b) is included. The time courses included are represented as early (6 h), mid (12 h), and late (24 h) course. The vertical axis is the relative expression of each gene under the osmotic potentials relative to control condition. The dashed horizontal lines define the threshold for up-regulation (upper limit line) and down-regulation (lower limit line). To show the threshold lines for up- and down-regulation, and because of the scale of Y axis, for gene *bz-504*, an inset panel with increased scale is included. Data are represented as means \pm SD of three replicates.

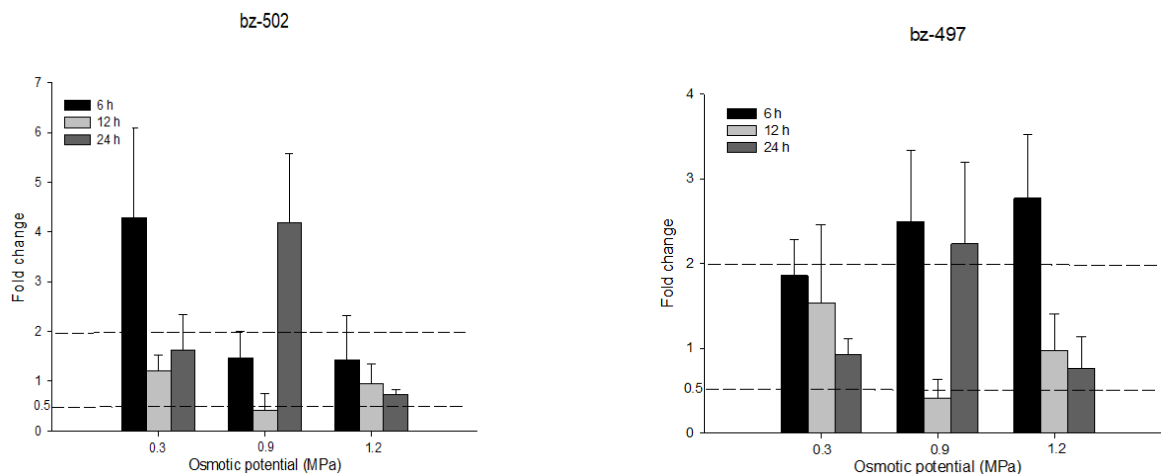


Figure 7. Relative expression of two bZip transcription factors, *bz-502* with stress responding behavior only in low stress level (0.3 MPa) and *bz-497* with stress responding behavior only in severe stress level (1.2 MPa) in response to different osmotic potentials (0.3, 0.9, and 1.2 MPa) at various time course of osmotic potential application. The time courses included are represented as early (6 h), mid (12 h), and late (24 h) course. The vertical axis is the relative expression of each gene under the osmotic potentials relative to control condition. The dashed horizontal lines define the threshold for up-regulation (upper limit line) and down-regulation (lower limit line). Data are represented as means \pm SD of three replicates.

time points under 0.9 MPa, second rise in the relative expression level was occurred at the late time point (24 h). As well, a more imposition to the osmotic potential (1.2 MPa) had no effect on its relative expression. In fact, this gene seems to be an early responding gene and has a low threshold for induction upon osmotic stress. These characters may represent position of this gene as a beginner in osmotic stress signal perception.

Upon signal perception, a series of signaling events are triggered, and thus no need for further expression of the gene.

The gene *bz-497*, however, showed an opposite response compared to that of *bz-502* (Fig. 7) which was no prominent relative expression, but rather a noisy feature under 0.3 and 0.9 MPa of the osmotic stress, and only an up-regulation response at early exposure

(i.e., 6 h) under 1.2 MPa.

Bz-499 has mostly shown a noisy pattern (Fig. 3), that is among, the gene appeared once as down-regulated and in another instance as an up-regulated gene. Nevertheless, the gene was down-regulated under imposition of 0.9 MPa osmotic stress for 12 h.

5. Discussion

The osmotic potential of the growth media has significant effects on the reduction of the fresh weight. In this study, sunflower seedling showed a level of tolerance (or buffering) to the increases of the medium osmolarity up to 0.9 MPa. Thus, 0.9 MPa could be considered as the threshold for overcoming the tolerance, or attenuating the buffering. So, the seedlings must employ different mechanism in order to cope against increases in the osmotic stress below 0.9 MPa. The early tolerance to the increases in the medium's osmotic potential has been reported in *Arabidopsis thaliana* (36). Therefore, it is a common response in plants to respond with a kind of buffering during the initial phase of stresses. During this phase, stress signal which has been already sensed by the plant is transduced and through activation of complex array of pathways adaptive responses will be initiated (37, 38). The growth reduction of sunflower seedlings under osmotic stress can be a result of both meristem growth retardation (39) and wasting metabolic energy for biosynthesis of osmoprotectants (40, 41), which are produced in high concentrations and upon osmotic stress (42, 43).

Arabidopsis and rice genomes contain 77 and 90 loci, respectively, coding for a large group of bZipTFs (<http://plntfdb.bio.uni-potsdam.de/v3.0/>), so, it is possible to perform TF profiling over the gene family on the plant under various conditions (44, 45). Although genome size of sunflower is estimated to be approximately 3.5 Gb (27), and not comparable to *Arabidopsis thaliana* (125 Mb) and *Oryza sativa* (389 Mb) (46, 47), only 33 bZip genes have been predicted in its genome (http://plantfdb_v1.cbi.pku.edu.cn:9010/web/index.php?sp=ha). As the gene sequences of these 33 genes are not fully presented in the aforementioned database, and number of sequences which met real-time PCR criteria for designing primers (23) were limited, we were able to monitor relative expression level of only 10 members of the gene family. Thus, we employed micro-transcription factor profiling term for cases which a subset member of a TF family was investigated. Nevertheless, updating the database could provide the possibility to monitor expression level of more genes from this TF family.

We were able to observe different patterns of gene

expressions among the genes under different osmotic potentials and time courses. Six TFs did not show any changes under the low stress level of 0.3 MPa. These observations are in agreement to the behavior of seedling fresh weight under this osmotic potential where no significant reduction in the fresh weight was noticed. As the effect of osmotic potential of 0.3 MPa on seedling growth is negligible, a phenomenon which we call it as seedling buffering, and because of up-regulation of a number of genes in this condition (e.g., *bz-505*, *bz-504*, *bz-502*, and *bz-485*), the genes may be involved in the protection of seedling under the mentioned osmotic potential. Indeed, buffering of seedling to 0.3 MPa might be a result of up-regulation of the mentioned genes in different time points. Certainly, in this buffering, many other TFs might be involved either belonging to this family or the other families, respectively.

Under 0.9 MPa condition a significant reduction in the fresh weight was observed for seedlings experiencing this potential for 24 h compared to the control condition, which was concomitant with down-regulation of gene *bz-499*. It is not unlikely that the gene has protective effects on biology of the seedling, and when plants is not able to keep its expression high enough to employ its protective role, fresh weight reduction is observed. Nevertheless, up-regulation of genes *bz-492*, *bz-509*, *bz-505*, and *bz-504* might be attributed to either their non-osmotic relevance or suppressing effect of the genes in the protective pathways.

The severe reduction in the fresh weight of seedlings that was observed in 1.2 MPa for 24 h may draw our attention to the negative role of *bz-504* on plant protection events that is when stress severity was increased from 0.9 MPa to 1.2 MPa the *bz-505* relative expression showed an increasing trend. Moreover, the role of *bz-504* in plant protecting systems is highlighted when its relative expression in early time of stress imposition for the three-osmotic potential is significantly high. The very high relative expression of the gene in the seedlings imposed for 6 h by 1.2 MPa osmotic stresses, and a deep reduction in the expression during subsequent time points may reveal that the gene belongs to the first line of defense of the seedling against osmotic stresses. This hypothesis gets closer to the reality when we realize that the used sunflower line in this experiment is classified as drought sensitive line.

During the time points, treating of the seedling with 24 h osmotic stress had significant effects on the seedling which was reflected in reducing the fresh weight. There is an interesting and clear companion between the reduction in the fresh weight and up-

regulation of *bz-505* in 24 h of stress impositions. Thus, suppressing effect of the gene in breaking of buffering against osmotic stress in sensitive line is proposed. Therefore, it is not unlikely that *bz-505* might play a more prominent role than other evaluated genes in this study in the adaptation and response of the seedling to the increases in the osmotic potential of the medium.

We made a small scale bZipTF profiling on a drought sensitive line. Although two of the ten evaluated genes did not show any responses to the osmotic stress, there also is not a solid evidence for the linkage of the eight other genes in response to the osmotic adaptation responses. When plants are exposed to the abiotic stresses, a plenty of genes appear to become subject of up- or down-regulation. However, not all the deregulated genes (genes which are effected by expression) have a relevant effect in the adaptation to the stresses (48, 49). However, this statement is not fully supported for TFs (10), which are low abundant master molecules with prominent effects on all aspect of biology (23). Indeed, their expression is strongly regulated in several levels (50), therefore, they are infrequently expressed as not-relevant genes in specific conditions. Although in this study a link was found between reduction in the fresh weight of the seedling under osmotic stress and the expression behavior of the genes *bz-504* and *bz-505*, for the other genes there was a lack of clear and straightforward pattern which could be drawn between phenotypic observations and the relative expression of the genes under investigation (patchy behavior of the gene expression (51), (Fig. 2). The reason for this patchy behavior can be explained by the two possibilities: i. small number of genes for analysis. It is possible that the genes under investigation are not the most important genes from this gene family. ii. the presence of a precisely regulated coordination between different components. Under this assumption, to gain a special response, several components such as various TFs are working in concert and these TFs may insert minor impacts on the whole response. BZip family of TFs contains protein members which their responses in several aspects of plants such as growth and development and stress signaling have been already proved (15, 16). So, we may consider different bZipTFs as dispersed pieces of a puzzle, which upon stress, they are brought together to make a whole response at molecular level. This finding one more time highlights the need for having systematics view on interpretation of biological data. This finding, additionally focuses on the requirement of having a systemic view on the complex response of the plant to the abiotic stress. By advancing in the genome information of the sunflower,

it is advised to bring more genes form the bZipTFs into account for elucidating more about the molecular aspect of the sunflower response to the osmotic stress. As an applied conclusion, monitoring of the molecular events in response of a drought sensitive sunflower line, rather than a tolerant one, to the osmotic stress can show us the weakness shunts of the lines under stress application. Having the knowledge will help molecular breeding procedures to fortify the weak points toward creation of stress tolerant cultivars in future.

Acknowledgments

We would like to mention and appreciate lab supports by Mr Esmail Rajaei from Shahid Beheshti University and Ms Neda Iravani, for their kind contribution through progress of the present work.

References

- Hanson PJ, Weltzin JF. Drought disturbance from climate change: response of United States forests. *Science Total Environ.* 2000;**262** (3):205-220. doi: 10.1016/S0048-9697(00)00523-4
- Ahuja I, de Vos RCH, Bones AM, Hall RD. Plant molecular stress responses face climate change. *Trend Plant Sci.* 2010;**15** (12):664–674. doi: 10.1016/j.tplants.2010.08.002
- Chaves MM, Flexas J, Pinheiro C. Photosynthesis under drought and salt stress: regulation mechanisms from whole plant to cell. *Ann Bot.* 2009;**103**:551-560. doi: 10.1093/aob/mcn125
- Chaves MM, Maroco JP, Pereira JS. Understanding plant responses to drought - from genes to the whole plant. *Func Plant Biol.* 2003;**30** (3):239-264. doi: 10.1071/FP02076
- Flexas J, Medrano H. Drought inhibition of photosynthesis in C3 Plants: stomatal and non-stomatal limitations revisited. *Ann Bot.* 2002;**89** (2):18318-9. doi: 10.1093/aob/mcf027
- Flexas J, Bota J, Galmés J, Medrano H, Ribas-Carbó M. Keeping a positive carbon balance under adverse conditions: responses of photosynthesis and respiration to water stress. *Physiol Plant.* 2006;**127** (3):343-352. doi: 10.1111/j.1399-3054.2006.00621.x
- Serraj R, Sinclair TR. Osmolyte accumulation: can it really help increase crop yield under drought conditions? *Plant Cell Environ.* 2002;**25**:333-341. doi: 10.1046/j.1365-3040.2002.00754.x
- Fujita Y, Fujita M, Shinozaki K, Yamaguchi-Shinozaki K. ABA-mediated transcriptional regulation in response to osmotic stress in plants. *J Plant Res.* 2011; 124(4):509-525. doi:10.1007/s10265-011-0412-3
- Nakashima K, Yamaguchi-Shinozaki K. Regulons involved in osmotic stress-responsive and cold stress-responsive gene expression in plants. *Physiol Plant.* 2005;**126** (1):62-71. doi: 10.1111/j.1399-3054.2005.00592.x
- Shinozaki K, Yamaguchi-Shinozaki K, Seki M. Regulatory network of gene expression in the drought and cold stress responses. *Curr Opin Plant Biol.* 2003;**6** (5):410-417. doi: 10.1016/S1369-5266(03)00092-X
- Rabbani MA, Maruyama K, Abe H, Khan MA, Katsura K, Ito Y, et al. Monitoring expression profiles of rice genes under cold, drought, and high-salinity stresses and abscisic acid application using cDNA microarray and RNA gel-blot analyses. *Plant*

- Physiol.* 2003;**133**:1755–1767. doi: 10.1104/pp.103.025742
12. Pérez-Rodríguez P, Riaño-Pachón DM, Corrêa LGG, Rensing SA, Kersten B, Mueller-Roeber B. PlnTFDB: updated content and new features of the plant transcription factor database. *Nucleic Acid Res.* 2009;**25**:1-6. doi: 10.1093/nar/gkp805
 13. Riechmann JL, Heard J, Martin G, Reuber L, Jiang C-Z, Keddie J, et al. Arabidopsis transcription factors: genome-wide comparative analysis among eukaryotes. *Science.* 2000;**15** (5499):2105-2110. doi: 10.1126/science.290.5499.2105
 14. Uno Y, Furihata T, Abe H, Yoshida R, Shinozaki K, Yamaguchi-Shinozaki K. Arabidopsis basic leucine zipper transcription factors involved in an abscisic acid-dependent signal transduction pathway under drought and high-salinity conditions. *Proc Nat Acad Sci U S A.* 2000;**97** (21):11632–11637. doi: 10.1073/pnas.190309197
 15. Kobayashi F, Maeta E, Terashima A, Kawaura K, Ogihara Y, Takumi S. Development of abiotic stress tolerance via bZIP-type transcription factor LIP19 in common wheat. *J Exp Bot.* 2008;**59** (4):891-905. doi: 10.1093/jxb/ern014
 16. Nijhawan A, Jain M, Tyagi AK, Khurana JP. Genomic survey and gene expression analysis of the basic leucine zipper transcription factor family in rice. *Plant Physiol.* 2008;**146**:333-350. doi: 10.1104/pp.107.112821
 17. Correˆa LGG, O-Pachó'n DMR, Schrago CG, Santos RVd, Mueller-Roeber B, Vincentz M. The role of bZIP transcription factors in green plant evolution: adaptive features emerging from four founder genes. *PLOS ONE.* 2008;**3** (8):e2944. doi: 10.1371/journal.pone.0002944
 18. Huang XS, Liu JH, Chen XJ. Overexpression of PtrABF gene, a bZIP transcription factor isolated from *Poncirus trifoliata*, enhances dehydration and drought tolerance in tobacco via scavenging ROS and modulating expression of stress-responsive genes. *BMC Plant Biol.* 2010;**10**:230. doi: 10.1186/1471-2229-10-230
 19. Chen W, Provart NJ, Glazebrook J, Katagiri F, Chang H-S, Eulgem T, et al. Expression profile matrix of arabidopsis transcription factor genes suggests their putative functions in response to environmental stresses. *Plant Cell.* 2002;**14**:559–574. doi: 10.1105/tpc.010410
 20. Baloglu MC, Eldem V, Hajyzadeh M, Unver T. Genome-wide analysis of the bZIP transcription factors in cucumber. *PLOS ONE.* 2014;**9** (4):e96014. doi: 10.1371/journal.pone.0096014
 21. Liu X, Chu Z. Genome-wide evolutionary characterization and analysis of bZIP transcription factors and their expression profiles in response to multiple abiotic stresses in *Brachypodium distachyon*. *BMC Genom.* 2015;**16**:227. doi: 10.1186/s12864-015-1457-9
 22. Rodriguez-Urbe L, O'Connell MA. A root-specific bZIP transcription factor is responsive to water deficit stress in tepary bean (*Phaseolus acutifolius*) and common bean (*P. vulgaris*). *J Exp Bot.* 2006;**57** (6):1391-1398. doi: 10.1093/jxb/erj118
 23. Czechowski T, Bari RP, Stitt M, Scheible WR, Udvardi MK. Real-time RT-PCR profiling of over 1400 Arabidopsis transcription factors: unprecedented sensitivity reveals novel root- and shoot-specific genes. *Plant J.* 2004;**38**(2):366-379. doi: 10.1111/j.1365-313X.2004.02051.x
 24. Jamieson GS, Baughman WF. The chemical composition of sunflower-seed oil. *J Am Chem Soc.* 1922;**44** (12):2952–7. doi: 10.1021/ja01433a036
 25. Nezamia A, Khazaiea HR, Rezazadehb ZB, Hosseini A. Effects of drought stress and defoliation on sunflower (*Helianthus annuus*) in controlled conditions. *DESERT.* 2008;**12**:99-104
 26. Haseeb M, Maqbool N. Influence of foliar applied nitrogen on reproductive growth of sunflower (*Helianthus annuus* L.) under water stress. *Agric Sci.* 2015;**6**:1413-1420. doi: 10.4236/as.2015.612137
 27. Staton SE, Bakken BH, Blackman BK, Chapman MA, Kane NC, Tang S, et al. The sunflower (*Helianthus annuus* L.) genome reflects a recent history of biased accumulation of transposable elements. *Plant J.* 2012;**72** (1):142-153. doi: 10.1111/j.1365-313X.2012.05072.x
 28. Giordani T, Cavallini A, Natali L. The repetitive component of the sunflower genome. *Curr Plant Biol.* 2014;**1**:45-54. doi: 10.1016/j.cpb.2014.05.001.9iuui800
 29. Pegadaraju V, Nipper R, Hulke B, Qi L, Schultz Q. De novo sequencing of sunflower genome for SNP discovery using RAD (Restriction site Associated DNA) approach. *BMC Genom.* 2013;**14**:556. doi: 10.1186/1471-2164-14-556
 30. Daneshian J. Response of sunflower lines to drought stress. Seed and Plant Improvement Research Institute; 2016. Personal communications.
 31. Michel BE, Kaufmann MR. The osmotic potential of polyethylene glycol 6000. *Plant Physiol.* 1973;**51** (5):914-916. doi: 10.1104/pp.51.5.914
 32. Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the 2^{-ΔΔCT} method. *Methods.* 2001;**25**:402-408. doi: 10.1006/meth.2001.1262
 33. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. *J Mol Biol.* 1990;**215**(3):403-410. doi: 10.1016/S0022-2836(05)80360-2
 34. Fernandez P, Rienzo JAD, Moschen S, Dosio GA, Aguirrezabal LA, Hopp HE, et al. Comparison of predictive methods and biological validation for qPCR reference genes in sunflower leaf senescence transcript analysis. *Plant Cell Rep.* 2011;**30** (1):63-73. doi: 10.1007/s00299-010-0944-3
 35. Bajić D, Poyatos JF. Balancing noise and plasticity in eukaryotic gene expression. *BMC Genom.* 2012;**13**:343. doi: 10.1186/1471-2164-13-343
 36. Weele CMVD, Spollen WG, Sharp RE, Baskin TI. Growth of Arabidopsis thaliana seedlings under water deficit studied by control of water potential in nutrient-agar media. *J Exp Bot.* 2000;**51**(350):1554-1562. doi: 10.1093/jexbot/51.350.1555
 37. Bartels D, Sunkar R. Drought and salt tolerance in plants. *Crit Rev Plant Sci.* 2005;**24**(1):23-58. doi: 10.1080/07352680590910410
 38. Ingram J, Bartels D. The molecular bases for dehydration tolerance in plants. *Ann Rev Plant Physiol Plant Mol Biol.* 1996;**47**:377-403. doi: 10.1146/annurev.arplant.47.1.37
 39. West G, Inzé D, Beemster GTS. Cell cycle modulation in the response of the primary root of Arabidopsis to salt stress. *Plant Physiol.* 2004;**135**(2):1050-1058. doi: 10.1104/pp.104.040022
 40. Yoshida Y, Kiyosue T, Nakashima K, Yamaguchi-Shinozaki K, Shinozaki K. Regulation of levels of proline as an osmolyte in plants under water stress. *Plant Cell Physiol.* 1997;**38**(10):1095-1102
 41. Valliyodan B, Nguyen HT. Understanding regulatory networks and engineering for enhanced drought tolerance in plants. *Curr Opin Plant Biol.* 2006;**9** (2):189-195. doi: 10.1016/j.pbi.2006.01.019
 42. Hare PD, Cress WA, Staden JV. Dissecting the roles of osmolyte accumulation during stress. *Plant Cell Environ.* 1998;**21**(6):535-553. doi: 10.1046/j.1365-3040.1998.00309.x
 43. Serraj R, Sinclair TR. Osmolyte accumulation: can it really help

- increase crop yield under drought conditions? *Plant Cell Environ.* 2002;**25**:333-3341. doi: 10.1046/j.1365-3040.2002.00754.x
44. Caldana C, Scheible W-R, Mueller-Roeber B, Ruzicic S. A quantitative RT-PCR platform for high-throughput expression profiling of 2500 rice transcription factors. *Plant Methods.* 2007;**3**:7. doi: 10.1186/1746-4811-3-7
45. Balazadeh S, O-Pacho'n DMR, Mueller-Roeber B. Transcription factors regulating leaf senescence in *Arabidopsis thaliana*. *Plant Biol.* 2008;**10**(Suppl. 1):63-75. doi: 10.1111/j.1438-8677.2008.00088.x
46. Matsumoto T, Wu J, Kanamori H, Katayose Y, Fujisawa M, Namiki N, et al. The map-based sequence of the rice genome. *Nature.* 2004;**436**:793-800. doi: 10.1038/nature03895
47. Kaul S, Koo HL, Jenkins J, Rizzo M, Rooney T, Tallon LJ, et al. Analysis of the genome sequence of the flowering plant *Arabidopsis thaliana*. *Nature.* 2000;**408**:796-815. doi: 10.1038/35048692
48. Fowler S, Thomashow MF. *Arabidopsis* transcriptome profiling indicates that multiple regulatory pathways are activated during cold acclimation in addition to the CBF cold response pathway. *Plant Cell.* 2002;**14** (8):1675-1690. doi: 10.1105/tpc.003483
49. Rabbani MA, Maruyama K, Abe H, Khan MA, Katsura K, Ito Y, et al. Monitoring expression profiles of rice genes under cold, drought, and high-salinity stresses and abscisic acid application using cDNA microarray and RNA gel-blot analyses. *Plant Physiol.* 2003;**133**(4):1755-1767. doi: 10.1104/pp.103.025742
50. Everett L, Hansen M, Hannehalli S. Regulating the regulators: modulators of transcription factor activity. *Methods Mol Biol.* 2010;**673**:297-312. doi: 10.1007/978-1-60761-854-6_19
51. Chen W, Provart NJ, Glazebrook J, Katagiri F, Chang HS, Eulgem T, et al. Expression profile matrix of *Arabidopsis* transcription factor genes suggests their putative functions in response to environmental stresses. *Plant Cell.* 2002;**14**(3):559-574. doi: 10.1105/tpc.010410