Journal of Stress Physiology & Biochemistry, Vol. 9 No. 3 2013, pp. 220-231 ISSN 1997-0838 Original Text Copyright © 2013 by Nageshbabu, Usha, Jyothi, Sharadamma, Rai, Devaraj

## **ORIGINAL ARTICLE**

# Expression of miRNAs confers enhanced tolerance to drought and salt stress in Finger millet (*Eleusine coracona*)

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Received March 19, 2013

Plants respond to the environmental cues in various ways, recent knowledge of RNA interference in conferring stress tolerance had become a new hope of developing tolerant varieties. Here we attempt to unfold the molecular mechanism of stress tolerance through miRNA profiling and expression analysis in Finger millet *(Eleusine coracona)* under salt and drought stress conditions. The expression analysis of 12 stress specific conserved miRNAs was studied using semi-quantitative real time PCR and Northern blot assay. Our studies revealed that, although most of the miRNAs responded to the stresses, the expression of particular miRNA differed with the nature of stress and the tissue. The expression analysis was correlated with the existing data of their target genes. Abiotic stress up-regulated miRNAs are expected to target negative regulators of stress responses or positive regulators of processes that are inhibited by stresses. On the other hand, stress down-regulated miRNAs may repress the expression of positive regulators and/or stress up-regulated genes. Thus the current study of miRNAs and their targets under abiotic stress conditions displays miRNAs may be good candidates to attribute the stress tolerance in plants by transgenic technology.

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MicroRNAs (miRNAs) are an important class of endogenous small silencing RNAs in regulating the expression of a wide range of target genes that are involved in many important biological processes in both plant and animals. These small (generally 21-22 nt) RNA molecules, originated from primary "hairpin" transcripts, can induce translational suppression or direct mRNA cleavage. Similar to regular mRNAs, the expression of miRNAs is highly regulated. Their expression pattern could provide critical clues in understanding miRNA functions. However, a detailed expression pattern for specific miRNA is difficult to obtain because of their short length, presence of similar miRNA families and intermediate precursor sequences. Many miRNAs show unique tissue-specific, development-related, and stress-induced expression. The miRNAs that are found to be regulated in response to a given abiotic stress could be of three types (1) known miRNAs that are known to be regulated by the stress, (2) known miRNAs that are newly discovered and to be regulated by the stress and (3) new miRNAs that are regulated by the stress. MicroRNAs of the first type are good positive controls and also could reveal unknown aspects of the regulation, such as tissue or organ specificity. Many of the miRNAs discovered till date have confirmed their involvement in regulatory networks associated with plant stress response. Of which 12 were found to be involved in salt and drought stress. These key players include miR156, miR157, miR159, miR166, miR167, miR169, miR171, miR172, miR393, miR395, miR396, miR398 families (Xie et al., 2010; Matts et al., 2010). Finger millet (Eleusine coracona) is the primary food source for millions of people in tropical regions. Till date many of the investigators focused on biochemical and physiological responses under environmental constrains. However, no work has been carried out in unwinding the role of miRNAs from the millet in regulating abiotic stress responses. Here we attempted to divulge these stress induced epigenetic regulators involved in restoring the cellular homeostasis.

#### MATERIALS AND METHODS

#### Plant material and stress treatments

Finger millet *(Eleusine coracona L* MR-1.) seedlings were grown in a green house at 28 °C; 13 h light until 8 day old and were then randomly divided into three groups. Two groups were grown in half-strength modified Hoagland nutrient solution containing two salt concentrations: 200mM NaCl and 400mM NaCl for 48 h and the other group was used for drought (with-holding water for 3 weeks) stresses respectively. Separate control lines were maintained for each stress. The shoots and roots were harvested separately from both control and stress samples frozen immediately in liquid nitrogen until RNA isolation.

#### RNA isolation and real time PCR analysis

Total RNA was isolated from different tissues of control and stressed seedlings using miRNA Isolation kit (Ambion) according to manufacturer's instructions. The quality of miRNA was analysed by 15% Urea PAGE, using 14, 24, and 39 nt oligos as size standards.Twelve miRNAs were selected for this study, which included miR156, miR157, miR159, miR162, miR167, miR169, miR172, miR395, miR396, miR397, miR398, and miR399. A majority of these miRNAs have been reported to play a role under stress conditions in model plant species. Semi quantitative real time -PCR was used to characterize the expression of 12 miRNAs in both shoot and root under salt and drought stress conditions. First, the RT reaction was carried out using TaqMan micro RNA reverse transcription kit (Applied Biosystems) the obtained cDNA was used for PCR amplification using family specific miRNA primers. The primers used in these amplification reactions were listed in supplementary table 1.

#### **RNA gel blot analysis**

For miRNAs quantification, northern blot hybridization was conducted using high sensitive miRNA Northern blot assay kit (Signosis, USA). 30 µg total RNA of each sample was electrophoresed on 15% polyacrylamide gel and transferred to membrane. Biotin-labeled Antisense RNA (Invitrogen) was used as hybridization probes and U6 RNA as a loading control. Supplementary table-2 lists the biotin labeled probes used for hybridization reaction.

#### RESULTS

# Salt stress altered the expression pattern of miRNAs in Finger millet

To get an insight into possible tissue-dependent roles of miRNAs in Finger millet, the expression patterns of miRNAs in different tissues including root and shoot were examined using semi quantitative real time PCR and Northern blot assay. Among the 12 miRNAs which have been considered for the study, only eight miRNAs revealed different expression patterns with respect to the tissue and nature of stress. The miRNAs responded varied between root and shoot. The expression of miR159 was up-regulated by two - fold in shoots, where as it remain unexpressed in roots. Similarly the down regulation of miR399 was identified only in shoots. Our results revealed that miRNAs responded differentially with the concentration of salt. Although the effect of salt concentration over miRNA expression was negligible with 0.1 to 0.5 fold altered expression in roots (Figure 1A), there was two-three fold variation in case of shoots. Most of the miRNAs were down regulated in shoots except miR156 and miR159. Two fold up-regulation of miR156 was observed, which was dose dependent at the same time miR159 showed 1.5

fold down -regulation at 400mM salt concentration compared to 200mM. The repression of miR157, miR167, miR169, miR398 and miR399 was also found to be dose dependent with 0.5- 1.0 fold down-regulation with increasing concentration. But no expression of miR398 was found at 200mM salt (Figure 1B). The Northern blot assay was performed with these differentially expressed miRNAs. The results confirmed the response of miRNAs towards the stress was tissue specific (Figure 2).

# Response of known miRNAs to drought stress in Finger millet

Although Finger millet is a drought tolerant species, the prolonged exposure of seedlings to the stress at the early vegetative stage is found to affect the crop yield significantly. The underlying mechanism of drought tolerance may be attributed to the altered expression of miRNAs. Expression studies of these epigenetic regulators with respect to drought stress were performed by semiquantitative RT-PCR and validated by Northern blot assay. Among the 12 miRNAs considered for study, 8 miRNAs responded to drought stress. miR171 and miR172 which were unnoticed in salt stress were found to express under water deficient conditions and remain to be stress specific. In roots, miR156 was up-regulated by 0.5 fold and miR171 by 0.3 fold. miR482 was expressed only in drought stressed roots as a secondary miRNA expressed in response to up-regulation of miR171. However, miR166, miR167 were down regulated by 10 fold and miR395 by 9 folds respectively (Figure 3A). In shoots, the response of miRNAs was quiet different. There was no expression of miR156, miR395 and miR171, where as the expression was found with miR397 and miR398 which were down regulated by 10 fold. The miRNAs which were up-regulated include miR159, miR169, miR172, and miR396 and their fold change was in the range of 1.5 to 3.0. miR159 had shown 3.0 fold up-regulation while miR396 was up-regulated by 0.5 fold (Figure 3B). miR396 had shown 0.5 fold in both the tissues and was not either tissue specific or stress specific. The Northern blot assay was carried out with differentially expressed miRNAs which established the tissue specific expression of miRNAs (Figure 4).

Majority of the targets shared by the abiotic

stress specific miRNAs were transcription factors such as MYB-Protein, AHAP2, NF-Y $\alpha$  2, CCAAT binding protein and other zinc finger proteins involved in transcription machinery. There were also target genes encoding proteins involved metabolic pathways such as phosphatidate cytidyltransferase, protein kinases, membrane bound channels, protein conjugating systems and membrane transporters. A description of the target genes were listed in supplementary (Table 3).



Figure 1. Relative expression levels of conserved miRNAs under salt stressed (root and shoot) Finger millet seedlings.



**Figure 2.** Northern blotting confirming differential expression of miRNAs. Total RNA 30µg from each of three conditions (control and salt) was loaded and hybridized with family specific miRNAs probe. U6 was used as control.



**Figure 3.** Northern blotting confirming differential expression of miRNAs. Total RNA 30µg from each of three conditions (control and salt) was loaded and hybridized with family specific miRNAs probe. U6 was used as control.



**Figure 4.** Northern blotting confirming differential expression of miRNAs. Total RNA 30µg from each of three conditions (control and drought) was loaded and hybridized with family specific miRNAs probe. U6 was used as control.

#### DISCUSSION

Although the miRNAs of several plant species have been studied recently, neither miRNA identification nor an analysis of differential miRNA expression in response to salt and drought stress has been performed in Finger millet. One reason why Finger millet has gained attention as a dedicated food crop is that it can grow on suboptimal land under salt and drought conditions. Our results indicated that salt stress had a significant effect on the germination rate and growth of Finger millet in almost all tested abiotic stresses. Interestingly, drought stress had no obvious effect on the germination rate; the significant effect of drought stress on Finger millet growth was observed at prolonged water deficient conditions at the early vegetative state. This result suggests that Finger millet has evolved a more effective mechanism to cope with drought stress as opposed to salt stress. Therefore, it is interesting to further investigate the change in gene expression, especially the gene expression regulators, under such stress conditions.

Micro RNAs are an extensive class of newly discovered gene regulators. They have been reported to play important roles under abiotic stress in model plant species. Using gRT-PCR, we studied the expression change of 12 conserved miRNAs in 8 days old Finger millet seedlings exposed to salt and drought stress. Of the 12 miRNAs, 11 have been demonstrated to be involved in salt or drought stress in previous study, eight of these in both the Arabidopsis thaliana and rice. Our results indicate that both salt and drought stresses altered the expression pattern of miRNAs in a dosedependent manner. The expression of an abiotic stress mediated miRNA depends upon the nature of the target gene. MicroRNA targeting the positive regulators of gene encoding proteins that are involved in processes inhibited by stress responses and negative regulators of stress responses were up-regulated and those targeting the positive regulators of stress response and stress upregulated genes were repressed.

miR156 known to play an important role in multiple biological processes by targeting Squamosal Promoter binding protein-Like (SPL) genes, miR156 has been demonstrated to temporally regulate shoot development (Park *et al.*, 2011), control the development timing from juvenile to adult transition together with miR172 (Xing *et al.*, 2009; Gou *et al.*, 2011). Overexpression of miR156 in *Arabidopsis*, rice, and maize led to a prolonged vegetative phase together with the production of significantly higher number of total leaves, which resulted in enhanced biomass accumulation (Liu et al., 2008; Zhang et al., 2009), miR156 was demonstrated by microarray-based analysis to response to salt stress but not to drought stress in Arabidopsis; miR156 was induced by 1.6-fold by salinity stress (Zhang et al., 2009). In rice, miR156 was found to respond to drought stress and was down-regulated by 2.1-fold by drought stress (Zhao et al., 2009). Studies have been conducted on maize under salt stress conditions and showed miR156 was not responded (Ding et al., 2011). However, our results demonstrate that expression of miR156 was upregulated in roots of drought stressed seedlings and shoots of salt stressed samples in contrast to the results found with Arabidopsis, rice and maize. The expression of miR169 differs with species and the nature of stress applied. In Arabidopsis, the miR169 expression is down regulated resulting in the accumulation of NF-YA5 transcription factor which is crucial for the expression of several DRE (dehydration- responsive elements), while in rice, the expression is favored by cold and drought stress. Medicago truncatula showed no expression in miR169 towards drought stress (Wang et al., 2005; Zhao et al., 2010). Reactive oxygen species (ROS) were found to accumulate under drought and salt stress as the oxidative stress occurs concurrently. The accumulated ROS results in membrane damage and lipid peroxidation. The expression of miR398 is down regulated during these induced stresses which lead to increased activities of superoxide dismutase the target genes of miR398. miR171 targets the gene of another family of GRAS transcription factors which is found to be involved in nodule morphogenesis and floral development (Wang et al., 2011). Targets of miR482 were primarily Resistance gene receptor kinases with two out of three targets are defense related

kinases and induced bacterial infections. These miRNAs were reported to target NBR-LRR gene their over expression leads to nodulation with increasing lateral root mass (Hue et al., 2010; Eckardt et al., 2012). The expression of different members of the MIR172 family and, consequently, their corresponding mature miRNAs, depends on growth stages and tissue types. In addition, the expression level of miR172 is also affected by day length and temperature (Kwon et al., 2011; Li et al., 2011). In both dicot and monocots, the expression level of miR172 increases as plants grow and after flowering it accumulates in leaves and floral buds (Aukerman and Sakai, 2003; Chuck et al., 2007; Zhu et al., 2009) miR172 sequence that regulates the mRNA abundance and/or translation of the plant specific transcription factor gene APETALA2 (AP2) (Park et al., 2002; Chen, 2004) and a small group of AP2-like genes in Arabidopsis (Aukerman and Sakai, 2003), which contain sequences complementary to miR172. AP2 is a floral organ identity gene while the five AP2-like genes mainly act as flowering repressor (Poethig et al., 2009).

In summary, our results provide insight into the Finger millet miRNAs, highlighting the regulatory network triggered by salt and drought stress. Furthermore, there exists a cross talk between salt and drought stress induced miRNA expression. The plant exhibits varied mechanisms to alter the miRNA expression tissue specifically thereby enhancing the tolerance towards stress.

#### ACKNOWLEDGEMENT

This work was supported from Department of Science and Technology, Government of India, New Delhi (Ref No; SR/FT/LS-10/2012).

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Su	pp	lementar	y Table 1:	List of	primers used	l for rea	l time -	<ul> <li>PCR</li> </ul>
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miRNA	miRNA Sequence	Primer names	Primer Sequences
miR 156	UGACAGAAGAGAGUGAGCAC	miR156-7RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC GTGCTC
		miR156FP	GCGGCGGTGACAGAAGAGAGT
miR 157	UUGACAGAAGAUAGAGAGCAC	miR156-7RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC GTGCTC ACTGGATACGACGTGCTC
		miR157FP	GCGGCGGTTGACAGAAGATAGA
miR 159	UUUGGAUUGAAGGGAGCUCUA	miR159RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC TAGAGC
		miR159FP	GCGGCGGTTTGGATTGAAGGG
miR 162	UCGAUAAACCUCUGCAUCCAG	miR162RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC CTGGAT
		miR162FP	GCGGCGGTCGATAAACCTCTG
miR 167	UGAAGCUGCCAGCAUGAUCUA	miR167RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC TAGATC
		miR167FP	GCGGCGGTGAAGCTGCCAGCA
miR 169	CAGCCAAGGAUGACUUGCCGA	miR169RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC TCGGCA
		miR169FP	GCGGCGGCAGCCAAGGATGAC
miR 172	AGAAUCUUGAUGAUGCUGCAU	miR172RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC ATGCAG
		miR172FP	GCGGCGGAGAATCTTGATGAT
miR 395	CUGAAGUGUUUGGGGGAACUC	miR395RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC GAGTTC
		miR395FP	GCGGCGGCTGAAGTGTTTGGG
miR 396	UUCCACAGCUUUCUUGAACUG	miR396RT	GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC CAGTTC
miR 397	UCAUUGAGUGCAGCGUUGAUG	miR396FP	GCGGCGGTTCCACAGCTTTCT GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC
		miR397RT	CATCAA
miR 398	UGUGUUCUCAGGUCACCCCUU	miR397FP	GCGGCGGTCATTGAGTGCAGC GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC
		miR398RT	AAGGGG
miR 399	UGCCAAAGGAGAUUUGCCCUG	miR398FP miR399RT	GCGGCGGTGTGTTCTCAGGTC GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGAC
		miR399FP	CAGGGC GCGGCGGTGCCAAAGGAGATT

Supplementary Table 2: List of biotinylated probes used for Northern Blotting

miRNA	Sequence (5'-3')	Probe sequence (5'-3') (Biotinylated at 5' end)
miR156	UGACAGAAGAGAGUGAGCAC	ACUGUCUUCUCUCACUCGUG
miR159	UUUGGAUUGAAGGGAGCUCUA	AAACCUAACUUCCCUCGAGAU
miR167	UGAAGCUGCCAGCAUGAUCUA	ACUUCGACGCUCGUACUAGAU
miR169	CAGCCAAGGAUGACUUGCCGA	GUCGGUUCCUACUGAACGGCU
miR396	UUCCACAGCUUUCUUGAACUG	AAGGUGUCGAAAGAACUUGAC
miR398	UGUGUUCUCAGGUCACCCCUU	ACACAAGAGUCCAGUGGGGAA

#### Cumplementer, Table 1. List of primars used for

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# Supplementary Table 3: Targets of predicted miRNAs

miRNA family	Targeted genes	Target description
miR172	TC405657	Hypothesis protein
	TC392019	Transcription factor AHAP2
	BE659941	Floral homeotic protein APETALA 2
	TC366837	APETAL2-like protein
	TC407080	Transcription factor AHAP2
	TC378006	Hypothesis protein
	TC383306	PHAP2B protein
	TC404733	Hypothesis protein
	TC352579	Hypothesis protein
	TC383335	PHAP2B protein
miR159		MYB - Protein
miR167	GD753695	Hypothesis protein
	TC371467	Phosphatidate cytidylyltransferase
	TC379788	Hypothesis protein
	TC371879	Hypothesis protein
	BE805600	Auxin response factor 8
	BM732289	Hypothesis protein
	DB979348	Hypothesis protein
	TC389689	Hypothesis protein
miR169	TC364843	Hypothesis protein
	TC353076	Nuclear transcription factor Y subunit A-3
	TC379261	Os02g0776400 protein
	TC401273	CCAAT-box transcription factor complex WHAP12
	TC355136	Hypothesis protein
	TC383014	CCAAT-binding transcription factor
	TC366077	Hypothesis protein
	CO985073	Mitogen-activated protein kinase 10
miR393	DB989850	Auxin-responsive factor TIR1-like protein
	TC416229	Auxin-responsive factor TIR1-like protein
	TC366828	Transport inhibitor response 1
	TC365328	Transport inhibitor response 1
	TC362546	Transport inhibitor response 1
	TC398603	Hypothesis protein
	TC362758	Hypothesis protein
	FG999850	Hypothesis protein
miR395	BG789910	ATP sulfurylase
	TC359920	ATP sulfurylase
	TC358067	ATP sulfurylase
	TC360687	ATP sulfurylase
	EV282501	Hypothesis protein
	GE008734	Hypothesis protein
	TC369301	Hypothesis protein
	TC349703	Plastidial lipoyltransferase 2
	TC348882	Plastidial lipoyltransferase 2

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	CF922366	Phospholipase C
	TC358694	Low affinity sulfate transporter 3
	TC358694	Low affinity sulfate transporter 3
	TC405478	Hypothesis protein
	TC411365	Cation diffusion facilitator 9
	AW759383	Glutamate dehydrogenase 2
	GR836780	Hypothesis protein
	TC369607	Zinc finger, CCCH-type; Sugar transporter superfamily
	TC357118	Hypothesis protein
	TC397169	Zinc finger, CCCH-type; Sugar transporter superfamily
	TC365139	Ketol-acid reductoisomerase, chloroplast precursor
	GD787823	Dihydroflavonol-4-reductase DFR1
miR396	TC365248	Hypothesis protein
	FK003100	Hypothesis protein
	TC373306	Cytochrome P450 monooxygenase CYP72A65
	TC366659	Hypothesis protein
	TC393538	Hypothesis protein
	TC379767	Hypothesis protein
miR398	TC393753 At1g08050	Elovl2 protein Cytchrome C Oxidase, zinc finger (C3HC4-type ring finger) protein
	At3g15640	CDS, Cytchrome C Oxidase, zinc finger (C3HC4-type ring finger) protein
	At3g15640	Cytchrome C Oxidase, zinc finger (C3HC4-type ring finger) protein
	At1g08050	Cytchrome C Oxidase, zinc finger (C3HC4-type ring finger) protein
	TC417910	Superoxide dismutase [Cu-Zn]
miR399	At2g33770	Ubiquitin conjugating enzyme (UCE); vesicle-associated membrane protein
	At4g00170 At2g33770	Ubiquitin conjugating enzyme (UCE); vesicle-associated membrane protein Putative UCE2