

Genome-Wide Analysis of Putative *ERF* and *DREB* GENE Families in Indica Rice (*O. sativa* L. subsp. *Indica*)

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Abstract—Drought is a major constraint to rice production and its stability in rain-fed and poorly irrigated environments. Identifying genomic regions influencing the yield and its response to water deficits will aid in our understanding of the genetics of drought tolerance and development of more drought tolerant cultivars. Besides drought, the other major impediment to increased crop production is salt stress. In this context, identification of drought and salt-responsive genes assumes significance.

In this paper we carried out genome-wide analyses to explore putative genes encoding ethylene responsive factor (*ERF*) and dehydration-responsive element binding proteins (*DREB*) in the genome of indica rice. Reference nucleotides of well established molecular function, representing each of the protein families investigated, were chosen as query sequences for searches in the indica rice genome database. Clones having genomic sequences similar to the related genes were taken and converted to amino acid sequences. Putative sequences were subjected to PROSITE and Pfam databases and 31 signature sequences related to *ERF* family and 30 sequences related to *DREB* were obtained. Proteins showing more than 30% identity were taken and phylogenetic trees were generated for each family. The results of this study provide basic genomic information about new *ERF* and *DREB* gene families in indica rice.

Index Terms—*Oryza sativa*, genome wide analysis, *DREB*, *ERF*

I. INTRODUCTION

Rice is the staple food in India accounting for more than half of the calories consumed. It is also a major source of livelihood for farmers especially in villages with rice-based production systems providing the main income and employment for more than 50 million households in India. Rice production in developing country like India continuously faces the challenge of keeping pace with an annual population increase, while the area of fertile wetland (lowland) available for rice farming is steadily decreasing due to urbanization and industrialization. To satisfy the demand for increased production of rice in the next decades, the only alternative is to expand rice cultivation to marginal dry-land (upland) areas, where rice production is severely hampered by dehydration stress due to drought and salt stress.

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Intensive research has been undertaken in the past few decades to identify drought and salt-responsive mechanisms in plants, both from a biological and genetic perspective. Transcription factors play a significant role in regulation of abiotic-stress responsive gene expression [1]. Transcription factors of the ethylene responsive factor (*ERF*) family contain a single AP2/*ERF* domain, and are sometimes further divided into two major subfamilies, the CBF (C-Repeat/*DRE*-Binding Factor)/*DREB* (Dehydration Responsive Element Binding protein) subfamily and the *ERF* subfamily [1]. Genes in the CBF/*DREB* subfamily play a crucial role in the response of plants to abiotic stresses by recognizing the dehydration-responsive element (*DRE*) with a core motif of A/GCCGAC [2], [3].

Manifestation of several abiotic stresses such as drought, and salinity occurs via alteration of water status of plant cells and are result in cellular dehydration. Water movement across cellular membranes is regulated largely by a family of water channel proteins called aquaporins. Aquaporins contribute to drought/salt tolerance by facilitating water transport between different organs and at the cellular level by maintaining water homeostasis [4], [5]. The availability of complete genome sequence of indica rice enables genome wide analyses of gene families of abiotic-stress responsive gene expression. Several of putative proteins have not yet been assigned to any families or subfamilies in indica rice. To gain further information, we have carried out an analysis of the genomic sequences related to rice *ERF* and *DREB* gene families by searching the indica variety genome in public databanks. The purpose of this study was identification of new *ERF* and *DREB* genes that could play a major role in drought/salt tolerance in rice and their phylogenetic analyses.

II. MATERIALS AND METHODS

The reference sequences of well-established molecular function, representing drought and salt tolerant gene families chosen as query sequences for searches in the indica rice (*O. sativa*) genome databases were EU910896 (*Solanum lycopersicum* ethylene responsive factor *JERF3*) and EF583447 (*Glycine max* Dehydration Responsive Element Binding *DREB5*). Taking specific members of these families as query sequences, searches were made using the TBLASTN tool [16] against GenBank database non-redundant (NR), with search specifications for *O. sativa* L. subsp. *indica* cultivar. The BLAST server used was that of the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/BLAST/>). As selection criteria of BLAST hits for genomic sequences, a cut off e-value of e^{-10} was previously set. The genomic sequences

found were used to predict putative genes contained within them. A mixed procedure was adopted combining *ab initio* gene prediction algorithms of genomic sequence alignments with similar sequences from expressed genes. The prediction algorithms used was GenScan (<http://genes.mit.edu/GENSCAN.html>). Such expressed sequences were found by BLAST searches against EST and NR databases of GenBank, using the genomic sequence as query. The algorithm of choice for the multiple alignments of protein sequences was Clustal W2 [17]. All the proteins with greater than 30% identity, with at least one of the reference proteins used in the searches, were regarded as functionally similar (homologous) to the reference proteins [18] - [21]. Those sequences that did not conform to this criterion were discarded.

Prediction of homology and signature sequences for the drought and salinity genes were carried out with PROSITE (<http://www.ebi.ac.uk/InterProScan>) [22] and Pfam [23] databases. Sequences were included into families based on homology and presence of signature sequences. For topology prediction, HMMTOP [24] was used. Protein alignments obtained with ClustalW2 [17] were used as starting points for phylogenetic analysis.

III. RESULTS AND DISCUSSION

Understanding the mechanisms of plants tolerance to environmental stress has the potential to provide new tools and strategies to improve the tolerance to environmental stress. Until recently, functional gene dissection of plants was largely carried out by molecular characterization of individual genes from various species. With the availability of complete genome sequences, bioinformatic analyses have assisted unraveling information stored in these genomes and facilitated studies of gene evolution.

The availability of data of complete genome sequencing of indica rice has made it possible to search for new, putative drought/salt-stress responsive genes. To search for *ERF* and *DREB* family genes in indica rice, BLAST searches of indica rice genome databases was performed using *Solanum lycopersicum* ethylene responsive factor *JERF3* and *Glycine max* Dehydration Responsive Element Binding *DREB5* as query sequences. After searching the databanks with TBLASTN, clones having genomic sequences to the related family were taken and converted to amino acid sequences. In each family, similar sequences were removed and the sequences were subjected to PROSITE and Pfam databases to see the presence of signature sequences for the corresponding families.

Thirty one full length cDNA clones in the genome of indica rice were identified as encoding ERF domains after these sequences were subjected to PROSITE (Table 1). All the 31 putative ERF proteins exhibited more than 30% identity with *Solanum lycopersicum* ethylene responsive factor *JERF3* and were taken for the construction of phylogenetic tree. In an earlier study, [15] identified 139 ERF family genes in japonica rice (*Oryza sativa* L. subsp. *japonica*) using a comprehensive computational analysis. It has been demonstrated that the AP2/ERF proteins have important functions in the transcriptional regulation of a variety of biological processes related to growth and

development, as well as various responses to environmental stimuli. Wang and Wu have demonstrated that the transcriptional regulation of *JERF3* modulated the increased tolerance to drought, salt, and freezing in tobacco during germination and seedling development [13], [14]. The transcriptional regulation was two-fold: activation of the expression of osmotic stress genes and photosynthetic carbon assimilation/metabolism and oxidative genes establishing the involvement of *JERF3* in a ROS-mediated regulatory module in transcriptional networks that govern plant response to stress.

TABLE I: ERF GENE FAMILIES FOUND IN INDICA RICE GENOME ORTHOLOGOUS TO SOLANUM LYCOPERSICUM JERF3 AND THEIR GENBANK ACCESSION NUMBERS. (* INDICATES PRESENCE OF FULL LENGTH CDNA)

Sequence name	GenBank accession number	Full length cDNA	% identity with query
OsJERF3.1	CT834226.1	*	40%
OsJERF3.2	CT832042.1	*	41%
OsJERF3.3	CT830564.1	*	68%
OsJERF3.4	CT832041.1	*	71%
OsJERF3.5	CT831646.1	*	66%
OsJERF3.6	CT837654.1	*	58%
OsJERF3.7	CT829047.1	*	62%
OsJERF3.8	CT832402.1	*	83%
OsJERF3.9	CT837611.1	*	61%
OsJERF3.10	CT828880.1	*	44%
OsJERF3.11	CT833424.1	*	58%
OsJERF3.12	CT833260.1	*	75%
OsJERF3.13	CT833261.1	*	75%
OsJERF3.14	CT831330.1	*	67%
OsJERF3.15	CT837590.1	*	69%
OsJERF3.16	CT832482.1	*	65%
OsJERF3.17	CT832723.1	*	70%
OsJERF3.18	CT829445.1	*	55%
OsJERF3.19	CT832234.1	*	42%
OsJERF3.20	CT834242.1	*	58%
OsJERF3.21	CT831165.1	*	66%
OsJERF3.22	CT835379.1	*	37%
OsJERF3.23	CT829864.1	*	64%
OsJERF3.24	CT835592.1	*	54%
OsJERF3.25	CT831975.1	*	36%
OsJERF3.26	CT828886.1	*	48%
OsJERF3.27	CT837872.1	*	55%
OsJERF3.28	CT836522.1	*	32%
OsJERF3.29	CT828346.1	*	46%
OsJERF3.30	CT834499.1	*	56%
OsJERF3.31	CT829643.1	*	48%

Thirty full length cDNA clones in the genome of indica rice were identified as encoding DREB domains after these sequences were subjected to PROSITE (Table 2). All the 30 putative DREB proteins exhibited more than 30% identity with *Glycine max* Dehydration Responsive Element Binding *DREB5* and were taken for the construction of phylogenetic tree. Increased expression of DREB genes have been implicated in increased tolerance to drought in Arabidopsis [1] [8], soybean [9], maize [11] and rice [10]. Each type of rice DREB protein is defined on the basis of demonstrating high homology with the corresponding DREB-type proteins in *Arabidopsis* in the AP2 domain [10]. Five *OsDREB* genes have been reported earlier by Dubouzet in japonica rice [10]; four of them being *DREB1*-related (*OsDREB1A-1D*) and the fifth one *DREB2*-related (*OsDREB2A*).

TABLE 2: DREB GENE FAMILIES FOUND IN INDICA RICE GENOME ORTHOLOGOUS TO GLYCINE MAX DEHYDRATION RESPONSIVE ELEMENT BINDING DREB5 AND THEIR GENBANK ACCESSION NUMBERS. (* INDICATES PRESENCE OF FULL LENGTH CDNA)

Sequence name	GenBank accession number	Full length cDNA	% identity with query
OsDREB5.1	CT829047.1	*	68%
OsDREB5.2	CT830564.1	*	80%
OsDREB5.3	CT831646.1	*	83%
OsDREB5.4	CT837611.1	*	74%
OsDREB5.5	CT834246.1	*	63%
OsDREB5.6	CT837654.1	*	62%
OsDREB5.7	CT833424.1	*	59%
OsDREB5.8	CT832042.1	*	61%
OsDREB5.9	CT832041.1	*	62%
OsDREB5.10	CT828880.1	*	62%
OsDREB5.11	CT832234.1	*	63%
OsDREB5.12	CT829445.1	*	62%
OsDREB5.13	CT832723.1	*	59%
OsDREB5.14	CT831330.1	*	65%
OsDREB5.15	CT832482.1	*	57%
OsDREB5.16	CT837590.1	*	65%
OsDREB5.17	CT832402.1	*	61%
OsDREB5.18	CT834242.1	*	62%
OsDREB5.19	CT833261.1	*	59%
OsDREB5.20	CT835379.1	*	61%
OsDREB5.21	CT833260.1	*	60%
OsDREB5.22	CT831165.1	*	60%
OsDREB5.23	CT829864.1	*	59%
OsDREB5.24	CT831975.1	*	39%
OsDREB5.25	CT835592.1	*	53%
OsDREB5.26	CT837872.1	*	51%
OsDREB5.27	CT836522.1	*	32%
OsDREB5.28	CT829643.1	*	41%
OsDREB5.29	CT288886.1	*	57%
OsDREB5.30	CT834499.1	*	56%

The detailed structural and phylogenetic analyses in this study provide insights into the presence and organization of putative *ERF* and *DREB* gene families in indica rice. The outcome of the present study provides basic genomic information for the gene families involved in response to abiotic stresses and will pave the way for elucidating the precise role of these genes in plant growth and development in the future. The unraveling of roles of individual members of these gene families in abiotic stresses tolerance will require a concerted effort by adoption of diverse approaches, including molecular genetic analysis when proofreading spelling and grammar:

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