# An *in silico* strategy identified the target gene candidates regulated by dehydration responsive element binding proteins (DREBs) in *Arabidopsis* genome

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**Abstract** Identification of downstream target genes of stress-relating transcription factors (TFs) is desirable in understanding cellular responses to various environmental stimuli. However, this has long been a difficult work for both experimental and computational practices. In this research, we presented a novel computational strategy which combined the analysis of the transcription factor binding site (TFBS) contexts and machine learning approach. Using this strategy, we conducted a genomewide investigation into novel direct target genes of dehydration responsive element binding proteins (DREBs), the members of AP2-EREBPs transcription factor super family which is reported to be responsive to various abiotic stresses in Arabidopsis. The genome-wide searching yielded in total 474 target gene candidates. With reference to the microarray data for abiotic stresses-inducible gene expression profile, 268 target gene candidates out of the total 474 genes predicted, were induced during the 24-h exposure to abiotic stresses. This takes about 57% of total

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Biotechnology Research Centre, Jilin Academy of Agricultural Sciences (JAAS), Changchun 130033, People's Republic of China predicted targets. Furthermore, GO annotations revealed that these target genes are likely involved in protein amino acid phosphorylation, protein binding and Endomembrane sorting system. The results suggested that the predicted target gene candidates were adequate to meet the essential biological principle of stress-resistance in plants.

**Keywords** Target genes prediction · Transcription factor binding site · DREBs · SVM · *Arabidopsis* 

#### **Abbreviations**

TFs Transcription factors

TFBS Transcription factor binding site

DBD DNA binding domain

DREBs Dehydration responsive element binding

proteins

DFSs DRE frame sequences nDFSs Non-DRE frame sequences

MGs Master genes

SVM Support vector machine

# Introduction

The identification of downstream target genes of specific transcription factors (TFs) is necessary in understanding cellular responses to environmental stimuli. Most existing structures of gene regulatory network are highly complicated as it involves cooperative interactions and feedback regulations. The discovery of the direct targets of TFs is a fundamental step to elucidate the construction of regulatory networks. Genes regulated by a given transcription factor is partially determined by the DNA binding domain (DBD) of a protein (Pabo and Sauer 1992). The DBD in TFs binds to





a specific DNA motif at the regulatory region of the target genes.

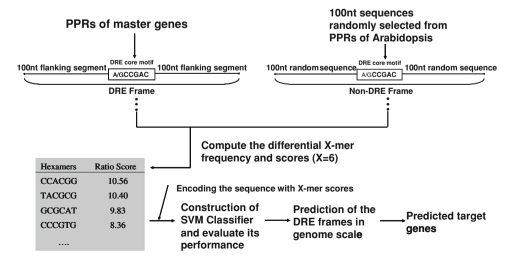
Availability of genome sequences made it possible to discover the target genes of a specific transcription factor by looking for the locations of the specific recognition motifs in genome. In practice, however, the task is still difficult due to the complication of plant genomes (Holstege and Clevers 2006). The development of microarray technology in measurement of mRNA expression profile promoted the identification of transcription factor target genes in genome (Young 2000). This approach allows the discovery of those genes that exhibit significant changes in mRNA levels upon inactivation or activation of a transcription factor (Qian et al. 2003, 2005; Maruyama et al. 2004). Despite this advancement, microarray profiling data might not be always sufficient in accurately identifying the direct target genes of TFs. For those organisms whose genomes are completed, an advanced computational strategy would be desirable to explore in genome-wide the binding motifs of the target genes for a given transcription factor.

During the last decade many computational methods have been developed to identify the target genes of TFs successfully (GuhaThakurta and Stormo 2001; Dieterich et al. 2003; Qian et al. 2003; Bigelow et al. 2004; Chan and Kibler 2005; Jolly et al. 2005; Zhang et al. 2005; Horsman et al. 2006; Redestig et al. 2007). Among the methods, the positional weight matrix (PWM) was the technique most widely used in describing the transcription factors binding sites (TFBS) and scanning the TFBS in the genome scale. However, owing to the looseness of the TFBS's conservation (Vavouri and Elgar 2005), these strategies were not capable of effectively identifying TFBS in genome scale. For this reason, the approach, including the PWM and the analysis of TFBS contexts, were developed to overcome the shortage (Kel et al. 2001; Rebeiz et al. 2002; Qian et al. 2005). The fundamental nature of the aforementioned approaches was in fact to develop appropriate algorithms that will describe the properties of the TFBSs and their contexts.

Dehydration responsive element binding proteins (DREBs) are important TFs that induce the expression of a series of abiotic stress-related genes and impart stress endurance to plants (Yamaguchi-Shinozaki and Shinozaki 1994; Agarwal et al. 2006). They belong to the ethylene responsive element binding factors (AP2-EREBPs) super family of 124 members (so-called ERF proteins), and among which 57 proteins are in the DREB subfamily (Riechmann et al. 2000). The ERF proteins share a conserved DNA binding domain (ERF domain) of 58-60 amino acids that, reportedly, binds to two typical cis-acting elements, that is, the GCC-box, and the C-repeat CRT/ dehydration responsive element (DRE) motif and involves in the expression of cold and dehydration responsive genes (Ohme-Takagi and Shinshi 1995; Hao et al. 1998; Fujimoto et al. 2000; Brown et al. 2003; Song et al. 2005). It is important to identify the target genes of DREBs in Arabidopsis since the DREBs play a vital role in various types of biotic and abiotic stress responses. Maruyama et al. identified the downstream genes of the DREB1A/CBF3 using two microarray systems (Maruyama et al. 2004). Fowler and Thomashow, Taji et al. also reported the downstream genes of DREBs proteins (Fowler and Thomashow 2002; Taji et al. 2002). Nevertheless, the overall target genes of DREBs are yet to be discovered.

In this paper, we reported a novel computational strategy to determine the DREB transcription factor binding sites in *Arabidopsis* genome by combination of the context analysis for the TFBS and machine learning approach. As illustrated in the flowchart (Fig. 1), the differences between the DRE frame sequences (DFSs, see section "Materials and methods") and non-DRE frame sequences (nDFSs, see section "Materials and methods") were given focus. A machine learning approach, specifically the support vector machine (SVM) based classifier, was developed to

Fig. 1 A flowchart showing the strategy for genome-wide identification of DREBs-targeted genes







categorize DRE-containing sequences into DFSs and nDFSs. Our results suggested that this algorithm was effective in the discovery of the DREB binding sites in the promoter region of the target genes, so as to infer the target genes of DREBs in *Arabidopsis*. Furthermore, we predicted 474 candidate genes as the direct targets of DREBs. With Reference to the AtGenExpress microarray data, we achieved the 268 direct targets of DREBs that was inducible by abiotic stress stimuli such as cold, salinity and drought during a 24 h observation.

#### Materials and methods

The flowchart of working method reported in this study was illustrated in Fig. 1. The data sets used were listed in Table 1. Three definitions operationally used in this study were made as:

Master genes (MGs)—The DREB-targeted genes that were experimentally identified contained the detectable DRE motifs in their putative promoter regions (PPRs). DRE frame sequences (DFSs)—DNA fragments of 206 bp, which were retrieved from the PPRs of MGs, contained a DRE motif (A/GCCGAC) at their center region.

Non-DRE frame sequences (nDFSs)—DNA fragments of 206 bp, which were collected randomly from the PPRs of *Arabidopsis* genome, with a DRE motif inserted artificially at their center region.

Collection of Arabidopsis putative promoter regions

Arabidopsis putative promoter regions (PPRs), the sequences of 1000 bp upstream of the start codon, were retrieved based on the TAIR (www.arabidopsis.org) genome annotations. All PPRs were scanned to explore the DRE motifs containing the A/GCCGAC consensus.

Collection of MGs

We searched for target genes of DREBs in public databases, including PubMed searching. A total of 32 genes

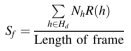
(Seki et al. 2001; Fowler and Thomashow 2002; Taji et al. 2002; Sakuma et al. 2006) were extracted which were further confirmed to be the target genes of DREBs by experimental approaches such as microarray analyses and real time PCR. The detailed information for these genes was listed in Table 2.

Analysis of the differential hexamers in the context of DRE sites

We collected 48 DFSs from the PPRs of MGs as the positive dataset and 1000 nDFSs as the negative dataset. HexDiff algorithm (Chan and Kibler 2005) was implemented to compute the hexamers frequency and ratio scores. Accordingly, the frequency of hexamers for positive  $F_p(h)$  and negative  $F_n(h)$  datasets on both strands were counted and used to calculate the ratio, R(h), as:

$$R(h) = \frac{F_p(h)}{F_n(h)}$$

The hexamers of larger ratio scores than the threshold were placed into  $H_d$ . In total we obtained 83 hexamers whose ratio scores  $R_h$  were greater than the threshold of 3.0. As a result of this selection process,  $H_d$  contained hexamers presented over in DRE frame rather than in non-DRE frame sequences. In this case, once the hexamers for  $H_d$  were chosen, the number of each hexamer  $N_h$  in a frame was counted and used to calculate the frame score  $S_f$  as:



The hexamer score  $S_h$  in a frame was calculated as:

$$S_h = N_h R(h), h \in H_d$$

Encoding the DRE-containing sequences

The hexamers in  $H_d$  were selected as the features used to encode the 206 bp DRE-containing sequences. Each sequence was encoded into an 84-dimension vector with 83 hexamer scores  $S_h$  and one label of either +1 or -1 to distinguish the DFSs (+1) or nDFSs (-1).

Table 1 The datasets used in this research

Dataset	Number of sequences	Description
DRE frame	48	Retrieved from the PPRs of master genes
Non-DRE frame	1000	Retrieved randomly from the PPRs
Random dataset	1000	Random sequence with equal nucleotide frequency $p(A) = p(T) = p(G) = p(C) = 0.25$ (used to evaluate the basic false positive rate, basic-FPR)



Table 2 List of the reported target genes of DREBs

AGI	DRE motif	Motif position	Functional description	Publications
AT5G52310	CATGGACCGACTACTA	-267	Hydrophilic protein	1, 3
	ATCATACCGACATCAG	-217		
	ATACTACCGACATGAG	-160		
	ATCAAGCCGACACAGA	-123		
AT2G17840	AAGCGACCGACCA	-204	Chloroplast related	2, 3, 5, 6
AT2G23120	CTACAGCCGACATCAG	-121	Unknown protein	1, 2, 5
AT5G62350	TCTGGACCGACGTTTA	-554	Endomembrane system	5, 6
	TGCTTGCCGACCTCTA	-186		
	TTTGTACCGACTATAA	-34		
AT5G15960	TAGCTACCGACATAAG	-79	Late embryogenesis-abundant protein	4, 5
AT5G15970	AAGCTACCGACATAAG	-125	Late embryogenesis-abundant protein	2, 5
AT1G01470	ATTCCACCGACGTGCA	-371	Late embryogenesis-abundant protein	1, 5, 6
	CATCGACCGACTTCAT	-23		
AT1G20440	ACATGACCGACATCTA	-989	Dehydrin	1, 3, 4, 5, 6
	TCAAAGCCGACCATTC	-960		
	CATCTACCGACTTCAA	-155		
AT1G20450	TTTCTGCCGACGTGGC	-603	Dehydrin	3, 4, 5
	ACATGACCGACATCCA	-567		
AT2G42530	TGATGGCCGACCTCTT	-188	Late embryogenesis-abundant protein	1, 2, 5
AT2G42540	TGTTGGCCGACATACA	-83	Hydrophilic protein	5, 6
AT3G50970	ACTACACCGACGTCTT	-207	Dehydrin	5, 6
AT4G15910	ATCTCACCGACCTCTT	-574	Late embryogenesis-abundant protein	5, 6
AT2G21660	CGTAAACCGACTCTAA	-81	Glycine-rich RNA-binding protein	4, 5, 6
AT1G22730	CATTGACCGACAATTC	-479	Zinc finger DNA binding protein	5, 6
AT5G58670	GGACAGCCGACAGATC	-154	Late embryogenesis-abundant protein	5, 6
AT3G24190	GTGGCGCCGACGTAGC	-302	Chloroplast	1, 5, 6
	TCGTTGCCGACGTAAT	-257		
AT4G24960	CTCTCACCGACCGACG	-329	HVA22 family	2, 3, 6
AT5G17460	GCCACGCCGACATAGT	-318	Mitochondrion	1, 5, 6
	AACAGGCCGACATAAT	-132		
AT5G04340	AAGTAGCCGACTTAAT	-406	Zinc finger DNA	1, 3, 5, 6
	TCTTAGCCGACTTCCA	-244	Binding protein	
AT2G17840	AAGCGACCGACA	-204	Unknown protein	2, 3, 4, 5, 6
AT4G33070	TTTAGACCGACATAAA	-181	Chloroplast	3, 4, 5, 6
AT4G35300	ATTATGCCGACATTAA	-398	Sugar transport protein	5
AT4G14000	CACAGACCGACTTTAA	-976	Unknown protein	5
	TGGAAGCCGACTAAAA	-640		
	CGGAAGCCGACCAAAG	-553		
	CTCGTACCGACCGGTT	-44		
AT4G15910	ATCTCACCGACCTCTT	-574	Unknown protein	2, 5, 6
AT1G52690	CTCTCGCCGACCAAGA	-41	Late embryogenesis-abundant protein	5, 6
AT1G69870	ATATGACCGACAACAC	-334	Proton-dependent oligopeptide transport family protein	5, 6
AT1G22985	TCAAAACCGACTTGAT	-117	AP2 domain-containing DNA binding protein	4, 5, 6
AT5G52300	CGTGGACCGACTAAAA	-166	Hydrophilic protein	2, 5, 6
AT3G09390	ATTGAACCGACGTACG	-145	Metallothionein protein	5
AT1G01470	ATTCCACCGACGTGCA	-371	Late embryogenesis-abundant protein	5, 6
	CATCGACCGACTTCAT	-23		
AT2G23120	CTACAGCCGACATCAG	-121	Unknown protein	5, 6

Publications refer to the literatures that reported the genes as the regulatory targets of DREBs. 1. Fowler and Thomashow (2002); 2. Seki et al. (2002); 3. Taji et al. (2002); 4. Maruyama et al. (2004); 5. Sakuma et al. (2006); 6. Yamaguchi et al. (2006)



## Support vector machines (SVMs)

The SVMs were standard supervised machine learning algorithms based on recent developments in statistical learning theory (Burges 1998; Vapnik 1998; Schölkopf et al. 1999; Schölkopf and Smola 2002). SVMs were binary classification tools that provided nonlinear function approximations by nonlinearly mapping the input vectors into feature spaces and using linear methods for regression or classification in feature space. Thus SVMs, and more generally kernel methods, combined the advantages of linear and nonlinear methods. For more details of SVMs, please refer Burges (1998), Vapnik (1998), Schölkopf and Smola (2002). Furthermore, SVMs have been used in many fields of bioinformatics such as splice site prediction, protein structural relationship prediction and microarray analysis and so on (Yu et al. 2003; Phan et al. 2005; Baten et al. 2006; Ogul and Mumcuoglu 2006; Towsey et al. 2006; Wee et al. 2006; Huang et al. 2007). In this study, we used SVM to discriminate DFSs from nDFSs. The Perl package of LIBSVM [http://www.csie.ntu.edu.tw/~cjlin/ libsym], an implementation of SVM by Chih-Chung Chang et al., was used to construct our classifier.

## Prediction of the target genes of DREBs

Once the construction of SVM classifier was completed, the SVM classifier was then used to scan the PPRs in order to get the DFSs in Arabidopsis genome. Using Regular Expression technique, we first scanned the PPR of genes in genome-scale to discover the DREB consensus (A/GCC-GAC). After that, to determine whether those consensuses with the 200 bp flanking sequences on both sides were DRE frames or not, they were classified through the optimized SVM classifier. In this case, the gene containing at least one DFS in its PPR was considered to be the target gene candidate of DREBs. Thereafter, the gene ontology (GO) enrichments of the candidate genes were then investigated using GO annotation system (Ashburner et al. 2000). The GO term file for Arabidopsis was downloaded from TAIR (filename: gene\_association.tair.gz) and GO-TermFinder (version 0.81) (Boyle et al. 2004) Perl module was used to get the term distribution of genes and the significance of the each groups.

## Collection of gene expression data

The AtGenExpress gene expression datasets, GSE5620 (for control), GSE5621 (cold stress), GSE5623 (salinity stress) and GSE5624 (drought stress), were downloaded from the gene expression omnibus (GEO) of the National Center for

Biotechnology Information (NCBI). Those genes were considered as inducible genes if their expression level fluctuated over 2-fold (either up or down) after cold, drought or salinity stress treatments.

#### Results

Being a group of important TFs responsible for abiotic stresses in plants (Chen et al. 2003; Kasukabe et al. 2004; Li et al. 2005; Agarwal et al. 2006, 2007; Wang et al. 2006; Xiong and Fei 2006), some DREBs were studied in the last decade and only 32 direct target genes of DREBs were reported (Fowler and Thomashow 2002; Seki et al. 2002; Taji et al. 2002; Maruyama et al. 2004; Sakuma et al. 2006; Yamaguchi et al. 2006). The DFSs in the PPRs of master genes were collected using home-made Perl scripts. The data was presented in Table 2. It was apparent that the collected DRE motifs of the MGs located within the 1000 bp upstream of the translation start codon (TSC). Thus, sequences of 1000 bp upstream of TSC were selected as the PPRs for this study. Given that not much information about promoters of non-DRE binding proteins target genes was available, as a control, the nDFSs were randomly extracted from the PPRs with DRE consensus (A/GCCGAC) added artificially (Fig. 1). In addition, a random sequences dataset was generated and was used to evaluate the false positive rate (Table 1).

# Discovery of over-represented hexamers in the DFSs

To identify the over-represented hexamers distributed among DFSs, the hexamers frequency ratios R(h) between DFSs and nDFSs were calculated in each dataset. Because there were only 48 DFSs found in positive dataset, which was indeed far less than the number of the nDFSs in the negative dataset, the sampling with replacement was then carried out to increase the positive dataset from 48 DRE frames to 500 DRE frames, which was about half the scale of the negative dataset. Appling the threshold varying from 2.0 to 5.0 with an interval of 0.5, we were able to get seven hexamer collections. Furthermore, we accessed the performance of SVM classifiers trained preliminarily on datasets encoding with each hexamer collections. Such was done to find out the satisfactory hexamer collections that were used as sequence feature. The results showed that the threshold at 3.0 gave rise to an optimal SVM classifier training in terms of accuracy and sensitivity (See Electronic supplementary material). Consequently, we collected 83 hexamers (See Electronic supplementary material) with



ratio scores >3.0 that were considered to be the distinctive motifs for further analysis.

Optimization of SVM parameters and performance evaluation

In general, the SVM involves two classes of parameters: the capacity parameter C and kernel type. C is a regularization parameter that controls the tradeoff between maximizing the margin and minimizing the training error. In the radial basis function (RBF) kernel used in our study,  $\gamma$  is an important parameter to dominate the generalization ability of SVM by regulating the amplitude of the kernel function. Accordingly, two parameters, C and  $\gamma$ , should be optimized. Thus, the parameter optimization was performed using a grid search approach within a limited range.

Using the optimized value of C and  $\gamma$ , the prediction model was thereafter constructed based on the training set with the RBF kernel. Moreover, to verify whether the hexamer features used in our study were able to effectively discriminate the DRE frames from the non-DRE frames and to avoid the dependence of the nDFSs dataset, five datasets were then constructed. Each dataset contained the 48 DFSs and 100 nDFSs that were randomly selected from the non-DRE frame dataset. The performances of SVM classifiers training each dataset were accessed by 5-fold cross-validation, and the data obtained from such procedure were presented in Table 3. It was shown that our classifier achieved the average accuracy of 85.6% with the average basic false positive rate of 9.12%. The parameters were likewise taken (C,  $\gamma = 64$ , 16) to scan the genome, as they were in the best prediction accuracy (87.2%). We also constructed the prediction models using three other kernel functions, i.e. polynomial, sigmoid, and linear functions, on the same data sets. However, the prediction accuracy of the aforementioned kernel functions achieved was not as satisfactory as that of the RBF kernel (data not shown).

Hunting the target gene candidates of CRT/DRE binding factors in genome scale

The regular expression protocol was used at first to explore the putative DFSs in the PPRs of genes in genome scale. After each sequence was encoded as vectors with 83 hexamers scores, the SVM classifier was then used to categorize whether the sequence was a DFS or not. Apparently, each gene occupied at least one predicted DFS among its PPRs was considered to be the target gene candidate of DREBs. From such procedure, a total of 474 target gene candidates, including 26 MGs, were obtained. The average number of targets per DREB transcription factor was about 8. The locus identifiers of target genes and DRE frame scores were listed in (See Electronic supplementary material). Mapping the positions of each predicted target gene to chromosomes indicated that the target genes candidates were well-distributed among the all five chromosomes in Arabidopsis.

Target genes responsive to abiotic stress

In this study, the expression profiles of those target gene candidates were investigated after 30-min exposure to the stresses such as cold, drought and salinity. Under such condition, a number of candidate genes induced were obtained and were presented in the Venn diagram (Fig. 2). A total of 160 genes (See Electronic supplementary material) were considered to be the direct targets of DREB TFs responsive to the abiotic stresses. Among them, 27, 91 and 106 genes were respectively responsive to drought, salinity and cold stress, respectively. In addition, there were ten genes were identified to be responsive to all the three types of abiotic stresses, indicating that they act as the cross-talkers in all likelihood among the drought, salinity and cold responsive pathways.

The positional distributions of the DRE motif occurred in each promoter of stress responsive genes were investigated. It was shown that the 160 target gene candidates

Table 3 Performances of SVMs trained on different datasets

Dataset	(C, γ)	Accuracy (%)	TP	FN	TN	FP	Sn (%)	Sp (%)	Basic-FPR <sup>a</sup> (%)
1	8, 128	85.2	32	16	94	6	66.7	94.0	8.44
2	64, 16	87.2	33	15	96	4	68.8	96.0	13.74
3	2, 512	85.9	33	15	94	6	68.8	94.0	14.32
4	8192, 0.0625	85.2	32	16	94	6	66.7	94.0	5.18
5	256, 4	84.5	30	18	95	5	62.5	95.0	9.76
Average		$85.6 \pm 1.02$					$66.7 \pm 2.6$	$94.6 \pm 0.9$	$9.12 \pm 3.08$

Accuracy = (TP + TN)/(TP + FN + TN + FP)

Sn = TP/(TP + FN), Sp = TN/(TN + FP)

<sup>&</sup>lt;sup>a</sup> Basic-FPR was evaluated by the random dataset



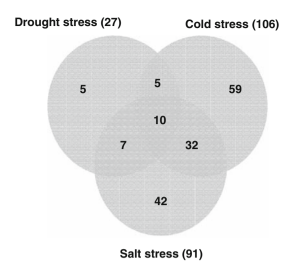


Fig. 2 Venn diagram showing the target genes induced after 30 min exposure to cold, drought or salinity

harbored the most DRE motifs within the 300 bp upstream of the start codon (Table 4). By analyzing of the gene expression profiles for 24 h, other 108 candidates were detected to be responsive to abiotic stresses (See Electronic supplementary material). In total, 268 target gene candidates (about 57% of predicted candidate genes) exhibited to be responsive to abiotic stresses.

Gene Ontology enrichment analysis of candidate targets of DREBs

Through the gene ontology enrichment analysis, it was found out that DREB target gene candidates were diverse, in terms of their functionality, molecular process and cell component categories. Approximately 5% of the candidates (38) were categorized as responsive to stress, and other 4.7% candidates (33) were functionally assigned as responsive to abiotic or biotic stimulus. Figure 3 illustrated that candidate target genes were among other categories significantly represented in "GO: 0012505 (endomembrane system)", "GO: 0006355 (regulation of transcription)", "GO: 0006468 (protein amino acid phosphorylation) and "GO: 0005515 (protein binding)". In addition, the target gene candidates induced by abiotic stresses showed the similar functional distributions pattern as the total predicted candidates (see Fig. 3).

# Discussion

In this study, attempts were made to discover computationally the DREBs-specific target genes through exploration of the DREBs binding sites in genome scale. Owing to the looseness of the DRE motif, it is insufficient to infer the targets of DREBs by simply scanning the DRE consensus in the PPRs of genes. A number of studies have already revealed that *cis*-acting elements distributed among the context of conserved TFBS also influenced the recognition by TFs (Wasserman and Fickett 1998; Kel et al. 2001; Krivan and Wasserman 2001; Zhang et al. 2005). It can be presumed that, in the PPRs of DREBs target genes, there were not only the DRE motifs but also other cisacting elements were flanking to the DRE motifs. Those cis-acting elements were thus used as supplementary information to promote the inference of the target genes of DREBs. The HexDiff algorithm, which was previously reported to predict novel cis-regulatory modules (CRMs) (Chan and Kibler 2005), was employed to get the differential cis-acting elements between the DFSs and the nDFSs. Yamamoto et al. and Gertz et al. identified the promoter constitution and regulatory regions using similar algorithms (Gertz et al. 2006; Yamamoto et al. 2007). The requirements of HexDiff algorithm suggested that it worked effectively in a well-defined system where the TFBS frames and non-TFBS frames were already clarified. As with many bioinformatic analyses, there is restriction based on the initial dataset on which predictions are based. In this study, we used 48 DFSs as positive data, which were relatively less than the negative data. It is expected that the accuracy of our algorithm would be improved when more DFSs are discovered and used for training in the future.

In machine learning, the imbalance problem (Japkowicz 2000) has to be considered when there existed great disparity between the size of the positive and negative training sets. In general, there are two kinds of method which were commonly introduced to overcome such imbalance problem, the methods in which the class represented by a small dataset gets over-sampled so as to match the size of the other class and the methods in which the class represented by the large dataset can be down-sized so as to match the size of the other class. In this study, the imbalance problem was encountered since there were more negative samples (nDFSs, 1000) than positive samples (DFSs, 48). In the process of discovering of over-represented hexamers, we augmented the size of positive dataset by the sampling with replacement to match the size of the negative dataset. While in the process of SVM training, we used 100 negative samples by downsizing the negative dataset that will match the positive dataset (48 samples).

A further attempt to understand the meaning of the hexamers was done by searching for hexamers in the databases of JASPAR (Sandelin et al. 2004) and TRANSFAC (Matys et al. 2003). Based on the gathered information, some hexamers were shown to be the conserved sequences of known *cis*-acting elements, such as the abscisic acid responsive element which was reported to be the coupling *cis*-acting element with the DRE motif in the stress





Table 4 The position distribution and sequence logo of DRE motifs in the master genes (MGs) and stress responsive genes

	Position distribution of DRE motifs	DRE motif frequency logo
Master genes (38)	15 0 -1000 -900 -800 -700 -600 -500 -400 -300 -200 -100 0 Position	ATTAGA COLACATAAA CERTEGO COLACATAAA LERGING COLACATAAA
Cold responsive genes (106)	25 20 15 10 5 -1000 -900 -800 -700 -600 -500 -400 -300 -200 -100 0 Position	ATTTGACCACAPATA FEEEE COMPLESSES
Drought responsive genes (27)	10 5 -1000 -900 -800 -700 -600 -500 -400 -300 -200 -100 0 Position	ATTTGACCACTAA Jeggese
Salt responsive genes (91)	20 15 10 10 5 10 10 10 10 10 10 10 10 10 10 10 10 10	ATTTAACCCACTATAA , eaeeeeeeeeeeeeeeeeeeee

The DRE motif frequency logos were generated by the WEBLOGO (Crooks et al. 2004) online server (http://weblogo.berkeley.edu/). The overall height of each stack indicated the sequence conservation at that position, whereas the height of symbols within the stack presented the relative frequency of the corresponding nucleic acid at that position

response (Zhang et al. 2005) and the Myb responsive element which was reported to be effective to increase the tolerance of the cold, drought and salinity stress (Dai et al. 2007). This information was valuable in analyzing the cooperative gene regulation of TFs.

The frame scores of DFSs and nDFSs showed that 43% of nDFSs scores were under 0.05, while only 5% accounted for DFSs scores (Fig. 4). It was difficult to give out a cut-off value to discriminate between DFSs and nDFSs as that was used in the reference (Chan and Kibler 2005). For this



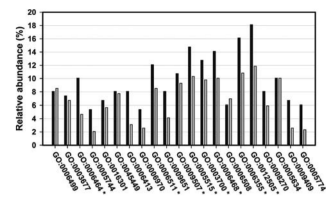
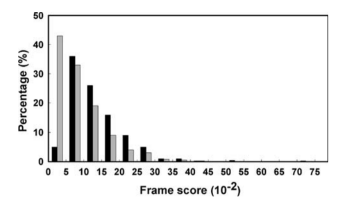


Fig. 3 Gene Ontology enrichments of target gene candidates. Black bars represented the relative frequency of GO classes in the set of total 474 target genes identified; grey bars indicated the relative frequency of GO classes for the set of stress responsive target genes. GO classes covering <2% of the genes were omitted for clarity and unknown classes were not shown. Significant overrepresented classes for the total 474 target genes (black bars) were indicated (\*, P < 0.01). (GO:0006499, N-terminal protein myristoylation; GO:0003677, DNA binding; GO:0006464, protein modification process; GO:0005744, mitochondrial inner membrane presequence translocase complex; GO:0016301, kinase activity; GO:0045449, regulation of transcription; GO:0006413, translational initiation; GO:0006970, response to osmotic stress; GO:0006511, ubiquitindependent protein catabolic process; GO:0009651, response to salt stress; GO:0005515, protein binding; GO:0003700, transcription factor activity; GO:0006468, protein amino acid phosphorylation; GO:0006508, proteolysis; GO:0006355, regulation of transcription, DNA-dependent; GO:0012505, endomembrane system; GO:0008270, zinc ion binding; GO:0005634, nucleus; GO:0009409, response to cold; GO:0005774, vacuolar membrane)

reason, we introduced the machine learning technique. To describe the characteristics of the TFBS context, the applications of linear classification model (Kel et al. 2001) and logistic regression model (Krivan and Wasserman 2001) were previously reported. The SVM was for the first time introduced in our study to discriminate the characters of the TFBS context. The results suggested that our computational approach was satisfactory and the resulting accuracy (more than 85%) was superior to that reported by Krivan and Wasserman (2001).

In this study, we mined 474 genes as the direct target genes of DREBs from the pool of genes with the DRE motifs in its PPRs. Among the 474 target genes, 56 genes (12%) were annotated as the stress responsive genes by GO annotations. Additionally, the functional distribution of the predicted 474 target genes was diverse. The top four categories involved were the endomembrane system, the regulation of transcription, protein binding and protein amino acid phosphorylation. Endomembrane system was reported to be a proteinase-sorting system involving in assistance of the plant cells under various stress conditions (Matsushima et al. 2002; Chitteti and Peng 2007; Wang et al. 2007; Prak et al. 2008). Protein phosphorylation was another mechanism which was largely reported to be



**Fig. 4** Histogram showing the distribution of the frame scores in the DFSs and nDFSs datasets. Black bars indicated the frame score distribution of the DFSs dataset; grey bars indicated the frame score distribution of the nDFSs dataset

responsible for the stress-relative proteins activation or deactivation (Kiegerl et al. 2000; D'Angelo et al. 2006; Boudsocq et al. 2007; Chae et al. 2007; Schweighofer and Meskiene 2008). Moreover, the phosphoproteome analyses of rice and *Arabidopsis* revealed the important roles of the phosphorated proteins responsive to different stresses (Khan et al. 2005; Chitteti and Peng 2007). These results suggested that the target gene candidates of DREBs predicted through the strategy proposed in this study were adequate in meeting the essential biological principle of stress-resistance in plants.

Exposing to the stresses for 30 minutes, 27, 91 and 106 genes out of the 474 candidate genes were induced respectively by drought, salinity and cold. Extending the stress exposure time to 24 h, further 108 genes out of the remaining genes in *Arabidopsis* were induced. The remaining genes may be the stress-inducible genes that appeared after the 24-h exposure to abiotic stresses, though the possibility of the false positives occurring in this case was not ruled out in this study. Nevertheless, our strategy was able to produce an immediate prediction with the existing TFBS knowledge and the microarray experiments that would be useful as a first step in the discovery of TF target genes.

## Conclusions

In this paper, we reported a novel computational strategy. Using the method proposed in this study, the analyses of TFBS contexts and SVM classification led to the identification of 474 target gene candidates of DREBs, a group of TFs playing a vital role in abiotic stresses responsive regulatory network in plants. With reference to the relevant DNA microarray data, the results showed that 160 candidate genes were induced in 30 min and other 108 candidate



genes were induced during 24-h exposure to abiotic stresses. The results obtained in this study provided the primary information that warranted further experimental investigation regarding the anti-stress regulatory network of DREBs in plants.

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