Full Paper

Prediction of bipartite transcriptional regulatory elements using transcriptome data of Arabidopsis

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Abstract

In our previous study, a methodology was established to predict transcriptional regulatory elements in promoter sequences using transcriptome data based on a frequency comparison of octamers. Some transcription factors, including the NAC family, cannot be covered by this method because their binding sequences have non-specific spacers in the middle of the two binding sites. In order to remove this blind spot in promoter prediction, we have extended our analysis by including bipartite octamers that are composed of ‘4 bases—a spacer with a flexible length—4 bases’. 8,044 pre-selected bipartite octamers, which had an overrepresentation of specific spacer lengths in promoter sequences and sequences related to core elements removed, were subjected to frequency comparison analysis. Prediction of ER stress-responsive elements in the BiP/BiPL promoter and an ANAC017 target sequence resulted in precise detection of true positives, judged by functional analyses of a reported article and our own in vitro protein–DNA binding assays. These results demonstrate that incorporation of bipartite octamers with continuous ones improves promoter prediction significantly.

Key words: promoter prediction, transcriptome, plant genome

1. Introduction

The availability of large-scale gene expression data by microarray and RNASeq analyses provides the possibility of predicting transcriptional regulatory elements in promoters using these data. For such predictions, methods for extraction of consensus sequences from a set of promoter sequences have been used (e.g. Gibbs Sampler1 and MEME2). As the accuracy of these established methods is not sufficient for experimental validation, we have developed a novel, more accurate method based on a simple frequency comparison.3 The new method shows considerably higher accuracy and sensitivity than conventional methods, judged by re-prediction of experimentally identified regulatory elements.3 In addition, our recent
studies demonstrated that it is also useful for predicting novel regulatory elements, and it has paved the way for prediction-oriented promoter analysis.\textsuperscript{4,6}

Our prediction method compares the frequency of every octamer between focused and global promoter sets. This octamer-based methodology has worked well for recognition sets of many types of transcription factors including bZIP, AP2/DREB, zinc finger, CAMTA, and also PCF families. However, it is theoretically not good for the detection of spacer-containing bipartite motifs that are recognized by protein complexes including dimers. As a consequence, these promoter elements are considered a blind spot of our octamer-based prediction method.

There are several databases for detection of transcription factor-binding sites in the promoter region. Among them, JASPAR\textsuperscript{7} is the most popular public database that covers higher plants. Detection of promoter elements is based on DNA motifs with each transcription factor proteins, and in the case of Arabidopsis which is the most intensively collected data for higher plants in datasets, motifs for 48 factors are available out of \(\sim 1,900\) transcription factors in the genome.\textsuperscript{8} The 48 factors include bipartite motifs for several MADS family proteins and an AP2 family protein, ANT. Obviously, the coverage is low for continuous and also bipartite motifs.

In this report, we have developed a supplemental method for promoter prediction that compensates for this blind spot by incorporating bipartite octamers (4 + 4 bases) to predictions that contain a spacer sequence in the middle. Our results using the pre-selected bipartite octamers showed improved prediction of the ER-stress responsive element (ERSE) as an element for tunicamycin-activated ER stress, and of ANAC017 target sites in promoters of the downstream genes of ANAC017.

2. Materials and methods

2.1. Bioinformatics analysis

24,956 Arabidopsis promoter sequences of 1 kb long, starting from the most major transcription start site (TSS), were prepared previously.\textsuperscript{7} Counting of each bipartite octamer, statistical analyses, and LDSS (local distribution of short sequences) analysis\textsuperscript{10} were achieved with home-made Perl and C\textsuperscript{++} programs, and Excel (Microsoft Japan, Tokyo). Distribution profiles of bipartite octamers were subjected to smoothing with a 21 bin, which is a good width for regulatory sequences (REG),\textsuperscript{10} clustered with Cluster\textsuperscript{11} by the hierarchical method with correlation measurement, and visualized with TreeView\textsuperscript{11} as described previously.\textsuperscript{10} Prediction of transcriptional regulatory elements based on microarray data with the aid of a list of bipartite octamers was achieved essentially according to our previous report\textsuperscript{7} using modified software. Motifs of clustered sequences were expressed using WebLogo.\textsuperscript{12} For prediction of ERSEs, 162 genes with a fold-change of over 2.5 were selected from up-regulated genes (E-MEXP-3186 from ArrayExpress, https://www.ebi.ac.uk/arrayexpress/ (11 January 2017, date last accessed)).\textsuperscript{13} as a positive promoter group. Microarray data of ANAC017 mutants were also retrieved from ArrayExpress (E-GEOD-41136).\textsuperscript{14}

2.2. DNA–protein binding analysis

Complementary DNA for ANAC017 was prepared by reverse transcriptase-polymerase chain reaction (RT-PCR) from total RNA of Arabidopsis seedlings using a conventional method as described elsewhere.\textsuperscript{15} The coding sequence of the Flag tag and a T7 promoter were added to the ANAC017 CDS excluding the transmembrane domain (1–523 aa\textsuperscript{14}) by PCR. The prepared PCR product was subjected to in vitro transcription by T7 RNA polymerase and subsequent in vitro translation by a wheat germ system as described previously.\textsuperscript{6} Binding assays of the Flag-tagged ANAC017 protein and biotinylated oligo-DNA probes were achieved using AlphaScreen (Perkin Elmer Japan, Tokyo) as described.\textsuperscript{6} Sequences of a biotinylated DNA probe and non-biotinylated competitors are shown in Fig. 6.

3. Results and discussion

3.1. Pre-selection of potential regulatory sequences

In our previous study, promoter prediction using microarray data based on enumerative octamer analysis was developed.\textsuperscript{3} The degree of overrepresentation for each octamer among a set of promoter sequences showing some transcriptional response over total promoters in a genome is calculated as the ‘Relative Appearance Ratio (RAR)’ . When predicting a specific promoter, an octamer is taken from the 5’ end of the promoter, the corresponding RAR value for the octamer is put back to the corresponding promoter position. Repeating these steps sliding 1 bp each time towards the 3’ direction provides a promoter scan with the RAR.\textsuperscript{3–5} Peak positions in the scanned results are predicted regulatory sites. We extended this procedure using a new type of bipartite octamers composed of ‘4 bases + a spacer + 4 bases’. In this article, we will refer to these octamers as ‘bipartite octamers (4 + 4)’.

As a pilot analysis, we surveyed 12 lengths of the spacer from 1 to 12 bases long. Analysis of all the possible bipartite octamers with a 12-spacer length gives 12 RAR values for each promoter point. While trials of this prediction method for cold- and high light-stress responsive elements detected novel promoter sequences that could not be detected by the previous octamer analysis without spacers, they also gave us highly complex results making their analysis difficult (data not shown). Therefore, we decided to pre-select potential regulatory sequences before carrying out frequency analysis. For this purpose, we set the priorities to high coverage over accuracy in the sequence selection, because an accurate, and thus small, selection at this point results in low sensitivity of prediction. Accuracy can be increased at the subsequent prediction steps.

During the analysis mentioned above, we noticed an overrepresentation of a specific spacer length over the others when the octamer (4 + 4) sequence is fixed. Fig. 1 shows examples of the relationship between the spacer length and the counts in the total 24,956 promoters of the Arabidopsis genome. In the case of ACGnTCGT (\(n = 0–12\)), a spacer of three appears \(\sim 4\) times more frequently than the other spacer lengths. The probability of this count profile, represented by the peak and total counts, under the assumption of random occurrence is very low (\(P = 2.2E-195\)), suggesting a strong selection pressure towards this biased profile. Interestingly, a count profile of its complementary sequence (ACGnCGTT) also shows the same characteristics where a spacer of three gives a single high peak.

Preferential appearance in the promoter sequence and conservation of characteristics between forward and reverse sequences may suggest that they are transcriptional regulatory sequences. Therefore, we decided to select bipartite octamers that have a preference for the spacer length among the promoter sequences.

With spacer lengths from 0 to 12, the number of possible bipartite octamers (4 + 4) is \(4^4 \times 13 = 65,536 \times 13\) (Table 1). First, one spacer length from 0 to 12 bp for each octamer with the highest count...
among the promoter sequences was selected, extracting 65,536 sequences from 65,536 \( \times \) 13. Second, sequences with spacers of 0 and 1 were removed from 65,536 sequences because they were thought to be detected by established analysis using octamers without a spacer. Then, the \( P \) value of the count profile with 12 spacer lengths for each bipartite octamer, as shown in Fig. 1, was considered, focusing on a peak height and the total count for 13 spacers, and 9,022 octamers with \( P < 1E-5 \) were selected.

These sequences were further subjected to LDSS analysis, where preference of appearance according to promoter position was evaluated. According to this analysis, 903 of them are revealed to be related to core promoter elements (Fig. 2A), including TATA box (Fig. 3A) and Y Patch (Fig. 3B), and 75 sequences with ‘Solid’ distribution profiles (Fig. 2B), which potentially include transposon-related sequences, will not be transcriptional regulatory elements. These 978 identified sequences (903 + 75) were removed for further analysis. A summary of the sequence extraction is shown in Table 1.

The remaining 8,044 sequences (Fig. 2C) were classified into three groups according to the LDSS profiles: the REG type (Clusters 1 and 2 in Figs 2C and 3C) that shows local distribution around -40 to -400 relative to the TSS and is a characteristic of a certain type of transcriptional regulatory element,\(^9\) the non-REG type whose distributions are scattered evenly with regard to promoter position (Clusters 5–9 in Figs 2C and 3D) and the intermediate REG-like type (Clusters 3 and 4 in Fig. 2C).

Figure 3C shows two distribution profiles of complementary sequences to each other, and both of them have very similar distribution profiles with peak positions around -100. These results demonstrate their distribution profile is direction-insensitive, and this feature is the same as the REG type of continuous octamers.\(^9\)

Considering the distribution profiles, sequences in clusters 1 and 2 of Fig. 2C are classified as the REG type.

Sequence lists of the removed core and solid groups, and also the 8,044 selected sequences containing REG and non-REG types can be found in Supplementary Tables S1–S3, respectively.

In order to pick up sequence groups from the extracted sequences, 1,053 selected REG-type sequences which are predicted to be regulatory sequences\(^9\) were mapped to 24,714 Arabidopsis promoter sequences with 1 kb length, and a matrix of presence/absence (1,053 \( \times \) 24,714) was subjected to 2D clustering analysis (Fig. 4). This analysis provided us the clearest classification of cis elements.\(^9\) As shown in Panel A, several clusters have been identified each of which shares bipartite octamers and promoters.

We subsequently selected 20 largest clusters for summarization of the bipartite octamers for each cluster as sequence motifs. The motifs prepared with WebLogo\(^1\)\(^2\) are shown in Panel B. Several tandem repeats (clusters 1, 6, 9, 12, and 13) and also palindromic sequence (clusters 8 and 11) are included. We suggest that they are recognized by a dimer or a trimer of the same protein family.

### 3.2. Inclusion of known transcriptional regulatory elements in selected bipartite octamers

The 8,044 selected sequences include REG-type groups. According to their LDSS profiles, they are strongly suggested to be transcriptional regulatory sequences.\(^9\) Their inclusion in the selected sequences is an indication of the ability of the procedure utilized to identify potential regulatory sequences.

In order to find other features that may be regulatory elements in the sequences selected, we surveyed known transcriptional regulatory elements within the sequences. One of the reported bipartite regulatory motifs is the endoplasmic reticulum stress-responsive element (ERSE, consensus sequence CCAAT-N_{9-12}CCACG) that is conserved between mammals and higher plants.\(^1\)\(^6\) The ERSE in mammals is known to be recognized by a general transcription factor (NF-Y for CCAAT and the stress responsive transcription factor ATF6 for CCACG, and spacing between the two elements is fixed to nine nucleotides.\(^1\)\(^7\) An ERSE is also found in Arabidopsis promoters and is known to have a conserved function.\(^1\)\(^8\)

When we searched the selected sequences for ERSEs, 28 related sequences were found (Table 2), indicating the inclusion of these elements in the extracted sequences. Then, microarray data of the transcriptional response to ER stress activated by tunicamycin treatment\(^1\)\(^9\) were used to calculate the RAR values of the selected sequences. Typically, RAR values of over 3.0 are considered ‘positive’ as transcriptional regulatory sequences.\(^3\) As shown in Table 2, the
RAR values of most of the ERSE-related sequences are much >3.0 (shown in bold in the table). These results mean that the bipartite ERSEs can be detected as ER stress (tunicamycin treatment)-responsive elements in our promoter prediction using the selected bipartite octamers (4+4). The application of such promoter prediction will be demonstrated later in this article.

Another reported bipartite motif of higher plants is the NAC-binding motif (CTTG-N5-CAAG) that is recognized by Arabidopsis ANAC013.20 This is also called the mitochondrial dysfunction motif (MDM) which includes the NAC-binding site and has several sequence variations.20 Our extracted bipartite sequences included 13 sequences related to the NAC-binding motif (Table 3). The presence of these sequences in the extracted sequences also supports the idea that our method does extract bipartite transcriptional regulatory sequences. One feature of this group is non-REG-type distribution (Table 3).

As the NAC gene family is composed of 94–106 genes in Arabidopsis,9 variations in the target sequence among this family

Figure 2. LDSS analysis of bipartite octamers. The distribution profiles of each octamer are shown. Each profile is shown as a colored horizontal line (rare = black, frequent = red), and the profiles of 903 (A), 75 (B), and 8,044 (C) octamers were subjected to hierarchical clustering. The cluster number of the ‘Rest’ group is indicated at the right end of the heat map (C). REG (1, 2), REG-like (3, 4), and non-REG (5–9).

Figure 3. Typical LDSS profiles of bipartite octamers. Typical distribution profiles are shown: TATA type (A), Y Patch-related Core type (B), REG type (C), and non-REG type (D). Profiles were subjected to smoothing with a bin of 21 bp, and the maximum value was adjusted to 1.0. The peak position of Panel A is -47, which means the first T of the TATA within the sequence, ‘ACTT ... TATA’, comes at -38. The two sequences in Panel C are complementary each other.
and differentiation of their function could be present. Therefore, the sequences in Table 3 may be recognized by different NAC proteins.

3.3. Applications of bipartite octamer analysis to promoter prediction

The selected 8,044 bipartite octamers were utilized for promoter prediction. An ER stress-responsive BiP3 promoter was selected for the prediction of tunicamycin-responsive elements. The results of this prediction were compared with those obtained using the established method using octamers without any spacers.

Figure 5A shows the results of promoter scans of BiP3 for tunicamycin-responsive elements. The vertical axis shows the RAR, and typically peaks >3.0 are selected as prediction sites. As shown, several peaks are >3.0, and two of them match experimentally identified ERSEs, ERSE-A, and ERSE-B.21 The RAR values of bipartite octamers are indicated with red X marks, and they show much higher RAR values than the continuous octamers at the positions of ERSEs, demonstrating the superiority of the bipartite octamers in detection of the ERSE.

Figure 5B and C shows sequences around ERSE-A (Panel B) and ERSE-B (Panel C), and the predictions using continuous octamers are shown in green letters. In both cases, the CCACG site of the ERSE, which is the recognition site (CGTGT, the underlining shows the match with the ERSE) of the ATF6 homolog of Arabidopsis, bZIP28,18 was detected by continuous octamer analysis, but the
Table 2. ERSE sequences in the selected bipartite octamers

| LDSS cluster | Sequence | Motif | RAR (Tunicamycin_up) |
|--------------|----------|-------|----------------------|
| 1_REG        | ACCA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.42 |
| 1_REG        | TCCA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 6.4  |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 3.52 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.65 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 6.42 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 5.64 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 5.84 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 7.76 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 6.83 |
| 1_REG        | CCAA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.95 |
| 2_REG        | AATA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 2.3  |
| 2_REG        | AATA      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 6.8  |
| 4_REG-like   | CACG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 8.53 |
| 2_REG        | GACG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.39 |
| 1_REG        | ACCT      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 3.93 |
| 2_REG        | ACCT      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.07 |
| 1_REG        | ACCT      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 8.69 |
| 1_REG        | ACCT      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 10.25|
| 2_REG        | CGTG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.37 |
| 1_REG        | CGTG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.28 |
| 1_REG        | CGTG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 6.64 |
| 1_REG        | CGTG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 11.96|
| 1_REG        | CGTG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 12.8 |
| 1_REG        | TGGT      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 10.88|
| 2_REG        | TGGT      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 5.69 |
| 3_REG-like   | AGTG      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 4.87 |
| 2_REG        | TGGC      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 5.74 |
| 4_REG-like   | TGTC      | ERSE: CCAAT-N<sub>9</sub>-CCACG | 5.09 |

LDSS cluster indicates the group number of LDSS clustering shown in Fig. 2C. Sequence matching with ERSE is indicated underlining. A dot in a sequence means any nucleotide. The RAR values of each octamer for tunicamycin up-regulation are also shown. RAR values > 3.0 are shown in bold.

Table 3. NAC target sequences in the selected bipartite octamers

| LDSS cluster | Sequence | Motif |
|--------------|----------|-------|
| 7_non-REG    | ACTT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 3_REG-like   | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 9_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 9_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 3_REG-like   | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 9_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 7_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 9_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 4_REG-like   | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 9_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |
| 9_non-REG    | CTCT      | NAC: CTTG-N<sub>9</sub>-CAAG |

LDSS cluster indicates the group number of LDSS clustering shown in Fig. 2C. Sequence matching with the NAC target motif is underlined. A dot in a sequence means any nucleotide.

NF-Y binding site (ATTTG, the underlining shows the match with ERSE) was not. These results indicate that the bZIP binding site of the ERSE has some specificity in ER-stress responsive promoters, but the NF-Y site does not have any specificity in the same promoters, resulting in the failure to detect the NF-Y site as an ERSE.

For comparison, the same gene list from the microarray data of tunicamycin response has been applied to Gibbs Sampler, MEME, CONSENSUS and MD Scan provided by Melina II. These four methods could not detect either ERSE-A or ERSE-B (data not shown).

Predictions using our bipartite octamers are shown in red letters in Fig. 5B and C. Both the bZIP and NF-Y binding sites were detected as putative ERSEs, indicating the advantage of using the bipartite octamer analysis to predict the bipartite ERSE. Comparison of the RAR signals between the two methods reveals that signals of the bipartite ERSE in the bipartite octamer analysis are much higher than those of the continuous octamer analysis (Panel A). This indicates higher sequence specificity of the detected ERSE sequences in the bipartite octamer method than in the continuous method.

These results, shown in Fig. 5, demonstrate that detection of ERSEs among tunicamycin-stimulated ER-stress responsive promoters has been considerably improved by introducing the bipartite octamer analysis.

The second application of our new method is the prediction of ANAC017 target sites using microarray data of Arabidopsis ANAC017 mutants. As ANAC017 is a mediator of H2O2 signaling, the effect of a mutation was compared with wild type after H2O2 treatments. As shown in Fig. 6A, a promoter region of ATERF71 was predicted as a target site of ANAC017 (underlined) according to the following results: the expression level of ATERF71 was reduced in the ANAC017 mutants, and a target site of
ANAC017 predicted by bipartite octamer analysis was mapped to the promoter region of *ATERF71*. In order to confirm the predicted target site of ANAC017, direct binding of ANAC017 protein to the DNA sequence was examined in vitro by the AlphaScreen method. A FLAG-tagged ANAC017 protein was prepared in vitro using wheat germ and subjected to binding assays with a biotinylated oligo DNA probe. As shown in Fig. 6, binding assays of the ANAC017 protein to the probe containing the predicted target site gave positive signals, indicating direct binding (no competitor, Panel B). Addition of a non-biotinylated competitor reduced the signal (competitor M0), but addition of mutated competitors of M1, M2, M3, and M6 failed to reduce the binding signals. The target sequence, revealed by these assays, that is necessary for the binding is shown with asterisks in Panel A. These results demonstrate that the bipartite octamer analysis developed here has predicted the exact target sequence of the ANAC017 protein based on microarray data of ANAC017 mutants.

3.4. Possibility of further extension

We have tried several patterns other than [4 + 4], but could not obtain good improvements so far. In an example, a zinc finger protein, STOP1, recognizes [3 + 3 + 2] in a target promoter, but prediction of the target site using its knockout data was better with [8] than with [3 + 3 + 2] for some unknown reason. Currently, only results of [4 + 4] are worth reporting. We have given up to increase division patterns. We now think that extension of octamer analysis is enough with addition of [4 + 4].

In summary, we have successfully developed a supplementary method for promoter prediction for spacer-containing transcriptional regulatory elements in Arabidopsis. This method should be used alongside the octamer analysis that was established year ago and is applicable to any other genomes that are thought to contain bipartite transcriptional regulatory elements.
Conflict of interest
None declared.

Supplementary data
Supplementary Tables S1–S3 are available at www.dnaresearch.oxfordjournals.org.

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