



Genome-wide identification and evolutionary analysis of the plant specific SBP-box transcription factor family

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ABSTRACT

We made genome-wide analyses to explore the evolutionary process of the SBP-box gene family. We identified 120 SBP-box genes from nine species representing the main green plant lineages: green alga, moss, lycophyte, gymnosperm and angiosperm. A maximum-likelihood phylogenetic tree was constructed using the protein sequences of the DNA-binding domain of SBP-box genes (SBP-domain). Our results revealed that all SBP-box genes of green alga clustered into a single clade (CR group), while all genes from land-plants fell into two distinct groups. Group I had a single copy in each species except for poplar while group II had several members in each species and can be divided into several subgroups. The SBP-domain encoded by all SBP-box genes possesses two zinc fingers. The C-terminal zinc finger of both group I and group II had the same C2HC motif while their N-terminal zinc finger showed different signatures, C4 in group I and C3H in group II. The patterns of exon–intron structure in *Arabidopsis* and rice SBP-box genes were consistent with the phylogenetic results. A target site of microRNA *miR156* was highly conserved among land-plant SBP-box genes. Our results suggested that the SBP-box gene family might have originated from a common ancestor of green plants, followed by duplication and divergence in each lineage including exon–intron loss processes.

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1. Introduction

Transcription factors (TFs) regulate and control gene expression in all living organisms. They are usually classified into different families and subfamilies based on the sequence of DNA-binding domains (Luscombe et al., 2000; Riechmann et al., 2005). *SQUAMOSA* is an *Antirrhinum majus* floral meristem identity MADS-box gene and the *SQUA* subfamily of MADS-box genes are critical in floral development (Huijser et al., 1992; Saedler et al., 2001; Fornara et al., 2004; Robles and Pelaz, 2005). SBP-box genes were first characterized as *SQUAMOSA* promoter binding proteins (SBPs) to regulate the expression of MADS-box genes in early flower development of *A. majus* (Klein et al., 1996). Since then, SBP-box genes have been identified in many plants including green alga, moss, silver birch, *A. majus*, *Arabidopsis* and maize. They play critical roles in regulating flower and fruit development as well as other physiological processes (Moreno et al., 1997; Eriksson et al., 2004; Lannenpaa et al., 2004; Arazi et al., 2005; Kropat et al., 2005). It has been reported that *Arabidopsis* *SPL3*, *SPL8*

and *SPL14* involves in flowering, sporogenesis, GA signaling and toxin resistance (Cardon et al., 1997; Unte et al., 2003; Stone et al., 2005; Zhang et al., 2006) while maize *tga1* and tomato *LeSPL-CNR* affect fruit development (Wang et al., 2005a; Manning et al., 2006). Recently, Xie et al. identified 19 SBP-box genes from the rice genome and revealed their predominant expression in several organs (Xie et al., 2006). Despite the importance of its function and the divergence of its gene structure, the origin and evolutionary process of the SBP-box gene family has not been reported.

SBP-box genes encode proteins sharing a conserved DNA-binding domain of 79 amino acid residues. It has been proved that the DNA-binding domain of SBP-box genes is necessary and sufficient to bind to a palindromic GTAC core motif (Klein et al., 1996; Cardon et al., 1997; Cardon et al., 1999; Lannenpaa et al., 2004; Birkenbihl et al., 2005). Studies on the NMR solution structure of the fragment of *Arabidopsis* *SPL4* and *SPL7* revealed that the DNA-binding domain of SBPs consisted of two separate zinc-binding sites. One zinc finger is C3H or C4 and the other is C2HC (Yamasaki et al., 2004). We are interested in the evolutionary process of these genes with two different zinc fingers.

Rhoades et al. predicted that 8 *Arabidopsis* SBP-box genes as potential targets of microRNA (miRNA) *miR156/157* (Rhoades et al., 2002). It has been reported that *miR156* is responsible for the temporal expression of *SPL3* during vegetative development (Schwab et al., 2005). *PpSBP3*, a moss SBP-box gene containing a miRNA target

Abbreviations: AA, Amino acid(s); CR, *Chlamydomonas reinhardtii*; EST, Expressed sequence tag; miRNA, microRNA; ML, Maximum likelihood; MP, Maximum-parsimony; NJ, Neighbor-joining; SBP, *SQUAMOSA* promoter binding protein; SPL, *SQUAMOSA* promoter binding protein like; TF, Transcription factor; UTR, Untranslated region.

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site, has been identified as a target of *miR156* (Arazi et al., 2005). Recently, Xie et al. identified 11 *OsmiR156* targets from rice SBP-box genes and revealed tissue-specific interactions between *OsmiR156* and *OsSBP* genes (Xie et al., 2006). However, it is unclear whether miRNA regulation is conserved in all land-plants and whether the SBP-box gene with a miRNA target site expand in each lineage.

Determining the phylogenetic relationships of the SBP-box gene family is an important step in elucidating the evolution and function divergence of this gene family. Phylogenetic analyses have been described for other plant TF families such as WRKY, MADS, GATA, AP2, DOF, etc. (Reyes et al., 2004; Wu et al., 2005; Zahn et al., 2005; Zhang and Wang, 2005; Moreno-Risueno et al., 2007; Shigyo et al., 2006; Shigyo et al., 2007). Cardon et al. (1999) identified 12 SBP-box genes from *Arabidopsis* and performed phylogenetic analysis for these *Arabidopsis* SBP-box genes together with other 12 SBP-box genes from *A. majus*, rice and maize. With more available plant genome sequences, comparison and phylogenetic analysis of SBP-box genes at genome scale are now possible. In this study, we made a genome-wide identification of SBP-box genes from 9 species representing the main plant lineages and performed phylogenetic analysis and classification to explore the evolution of SBP-box gene family. The feature of the exon–intron structure, the pattern of the conserved motifs, the role of the miRNA target, and the divergence of function are also discussed.

2. Materials and methods

2.1. Identification of SBP-box genes

We obtained the *Arabidopsis* SBP-box gene list from the DATF SBP-box gene family (Guo et al., 2005) (<http://datf.cbi.pku.edu.cn>) which was built based on the *Arabidopsis* TAIR6 genome release (<http://www.arabidopsis.org/>). *Oryza sativa* ssp. *japonica* genome data were downloaded from the TIGR rice genome annotation database release 4 (Yuan et al., 2005) (<http://rice.tigr.org/>). We performed HMMER (<http://hmmerr.wustl.edu/>) search using the Pfam profile PF03110 against *japonica* proteome sequences and refined the results manually to obtain the rice *japonica* SBP-box genes. We combined the BLAST search results generated from both TIGR maize gene index and TIGR maize database release 4.0 (<http://maize.tigr.org/>) with previously reported 8 members (Cardon et al., 1999) to identify maize SBP-box genes. We searched the draft genome sequences from PHYSCObase (<http://moss.nibb.ac.jp/cgi-bin/blast-assemble>) for moss (*Physcomitrella patens*) and lycophyte (*Selaginella moellendorffii*) SBP-box genes. We obtained poplar (*Populus trichocarpa*) SBP-box genes from DPTF (<http://dptf.cbi.pku.edu.cn>), which was built based on the JGI poplar genome release 1.1 (http://genome.jgi-psf.org/Poptr1_1/Poptr1_1.home.html). We downloaded green alga (*Chlamydomonas reinhardtii*) genome sequence from the US Joint Genome Institute (<http://genome.jgi-psf.org/>) and identified green alga SBP-box genes by HMMER search. Finally, we made BLAST search against the NCBI non-redundant and dbEST databases to obtain SBP-box genes in other species including pine and spruce. We used E value < $1e-5$ as the cutoff in HMMER and BLAST searches since the SBP-domain is very conserved and specific (<http://www.sanger.ac.uk/Software/Pfam/data/jtml/seed/PF03110.shtml>). Finally, we manually checked the search results to reduce hits with partial SBP-domain and other false positives.

2.2. Phylogenetic analysis

For the phylogenetic analysis, we considered only the amino acid sequence of the SBP-domain since no other regions could be aligned unambiguously for all the sequences available. We used ClustalW (v1.81) (Higgins et al., 1996) for the multiple sequence alignment of the SBP-domains with default settings and manually refined the alignment. We used PHYLIP (v3.6) ([\[ton.edu/phylip.html\]\(http://ton.edu/phylip.html\)\) to construct neighbor-joining \(NJ\) and maximum-parsimony \(MP\) phylogenetic trees with 1000 replicate bootstrap tests. For the NJ method, pairwise distances were calculated using PROTDIST under the JTT model \(Jones et al., 1992\). Site unweighted MP trees were generated using PROTPARS under ordinary parsimony and randomized input order. Finally, the 1000 trees were reduced to the majority-rule consensus tree using the program CONSENSE. The maximum-likelihood \(ML\) trees were obtained using the program MOLPHY \(v2.3\) \(<http://www.ism.ac.jp/ism/lib/softother.e.html>\). An NJ tree was obtained with the NJdist program based on the ML distance in the JTT model. The NJ tree was used as a start tree for a local rearrangement search. The likelihood of trees was calculated using the ProtML program under the JTT model, and the trees were sorted according to the Akaike information criterion \(AIC\) values. The local bootstrap probability of each branch was estimated using the resampling-of-estimated-log-likelihood \(RELL\) method. Two SBP-box genes from spruce and maize were excluded in the phylogenetic analysis because only the N-terminal 50 residues of the SBP-domain were available from the incomplete sequence data.](http://evolution.genetics.washing-</p>
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2.3. Exon–intron structure and motif analysis

The diagrams of exon–intron structure were obtained using the online Gene Structure Display Server we developed (GSDS: <http://gsds.cbi.pku.edu.cn>) with either GenBank accessions, or both CDS and genomic sequences. The sequence logos were generated using the online Weblogo platform (<http://weblogo.berkeley.edu/>). We downloaded the MEME package (<http://meme.sdsc.edu/meme/>) and installed it locally for motif search. The SBP-domains and the long C-termini of proteins of group I, subgroup IIa and *OsSBP1* were excluded in the motif search since they were very similar among themselves.

3. Results

3.1. Identification of SBP-box genes in all lineages

We searched the NCBI non-redundant database and dbEST for members of the SBP-box gene family. All positive hits were from green plants except for a suspicious entry from fungi. This entry is an EST from *Phytophthora infestans*, an oomycete that causes the late blight disease in potato and tomato [GenBank: CV968636]. We doubt that this EST might be a contaminated sequence from tomato since it has 99% identity with two tomato EST sequences [BI934749, BF096268], and we did not find it in the *Phytophthora* functional genomics database (<http://www.pfgd.org/>).

BLAST search results showed that SBP-box genes existed in various green plants from unicellular green algae, mosses, lycophytes, to gymnosperms and angiosperms, but not in brown alga (Phaeophyceae), red alga *Rhodophyta* and blue-green alga (*Cyanobacteria*) and golden algae (*Chrysophyceae*). To explore the origin and evolutionary process of the SBP-box gene family, we characterized SBP-box genes from species representing the main lineages of the plant kingdom: the green alga *C. reinhardtii*, the moss *P. patens*, the lycophyte *S. moellendorffii*, the gymnosperms pine and spruce, the dicotyledonous angiosperms *Arabidopsis* and poplar, the monocotyledonous angiosperms rice and maize. Either complete or draft genome sequence was used in our searches except for the two gymnosperms (pine and spruce) whose genome sequences were not yet available. Finally, we obtained a dataset of 120 SBP-box genes from the above 9 plants (Table 1; see Supplementary material, Table S1). Potential SBP-box genes from green alga, pine, spruce, maize and poplar with partial SBP-domains were excluded in this dataset.

Among the 16 *Arabidopsis* SBP-box genes (*AtSPL1-15* and *AtSPL17*), *AtSPL13* (At5g50670) and *AtSPL17* (At5g50570) were located on two adjacent BACs. The coding sequences of these two genes were the same, but the upstream and downstream regions of these two genes

Table 1
Number of SBP-box genes in nine representative plants

Lineage	Organism	Number	Nomenclature
Alga	<i>Chlamydomonas reinhardtii</i>	7	CrSPL
Moss	<i>Physcomitrella patens</i>	14	PpSBP
Lycophyte	<i>Selaginella moellendorffii</i>	13	SmSBP
Gymnosperm	<i>Pinus taeda</i>	3	PtSBP
	<i>Picea glauca</i>	2	PgSBP
Dicots	<i>Arabidopsis thaliana</i>	16	AtSPL
	<i>Populus trichocarpa</i>	26	PtSBP
Monocots	<i>Oryza sativa</i>	18	OsSBP
	<i>Zea mays</i>	21	ZmSBP
Total		120	

were different. (TAIR, <http://www.Arabidopsis.org>). We considered them as duplicates generated by recent duplication event. The locus *At1g76580* which was named as *AtSPL16* previously did not encode an SBP-domain in either cDNA clone (GenBank: NM_106308 and AY062670) or TAIR annotation. Interestingly, a complete SBP-domain was found by BLASTX at the upstream of the cDNA sequence. It was reported that *AtSPL14* (*At1g20980*) and *AtSPL16* (*At1g76580*) were duplicated gene pairs (Bowers et al., 2003; Blanc and Wolfe, 2004). We excluded *At1g76580* since its SBP-domain was lost due to possible frame shift mutation. A complete SBP-domain was distributed in two

neighboring *japonica* loci (*LOC_Os11g30380* and *LOC_Os11g30370*) separated by a stop codon at the conserved intron position of the SBP-domain. We excluded these two genes in our analysis. We removed the extra fragment at the conserved intron position of the SBP-domain of *OsSBP17* (*LOC_Os09g31438*) according to the available rice cDNA sequences (Kikuchi et al., 2003).

3.2. Phylogenetic relationships of SBP-box genes in all lineages

We constructed an unrooted maximum-likelihood (ML) phylogenetic tree for the 120 SBP-box genes from 9 species (Fig. 1) based on the amino acid sequences of their SBP-domains (see Supplementary material, Figure S1 and S2). In addition, using the neighbor-joining (NJ) and maximum-parsimony (MP) methods, we obtained trees with similar topology (data not shown). The tree topology and the corresponding phylogenetic relationships indicated that all proteins from green alga were grouped into the same clade (CR group), while those from land-plants were grouped into several other clades. Group I contained 6 land SBP-domains with a distinct feature that the zinc finger at the N-terminal consisted in four Cys residues while the N-terminal zinc finger of group II SBP-domains had a Cys3His motif (Fig. 2). In addition to the special zinc finger pattern, SBP-box genes in group I showed some features different from the other two groups.

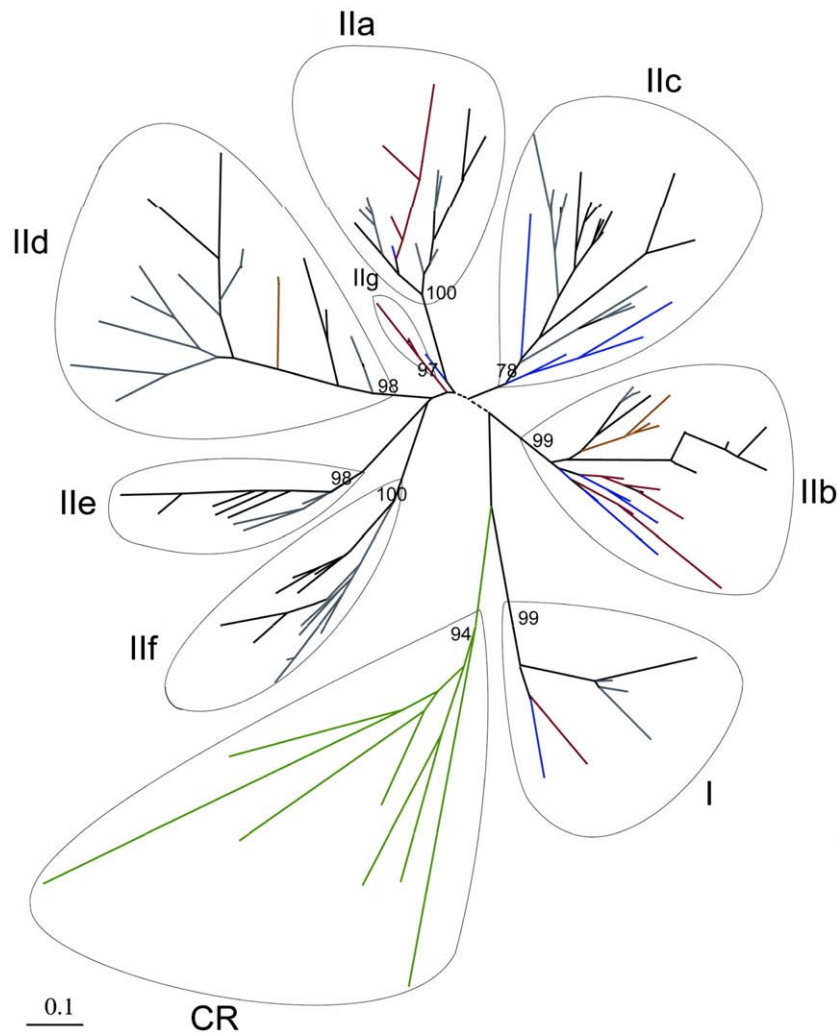


Fig. 1. Unrooted maximum-likelihood tree of 120 complete SBP-domains. The tree was inferred by the maximum-likelihood (ML) method implemented in Molphy based on amino acid sequences of the SBP-domains. The dash line connecting the clades of group II indicates the uncertain relationship among different subgroups (bootstrap probability <70%). Scale bar corresponds to 0.1 amino acid substitution per residue. Colors denote different lineages, green: green alga, red: moss, blue: lycophyte, brown: gymnosperms, black: monocots, gray: dicots.

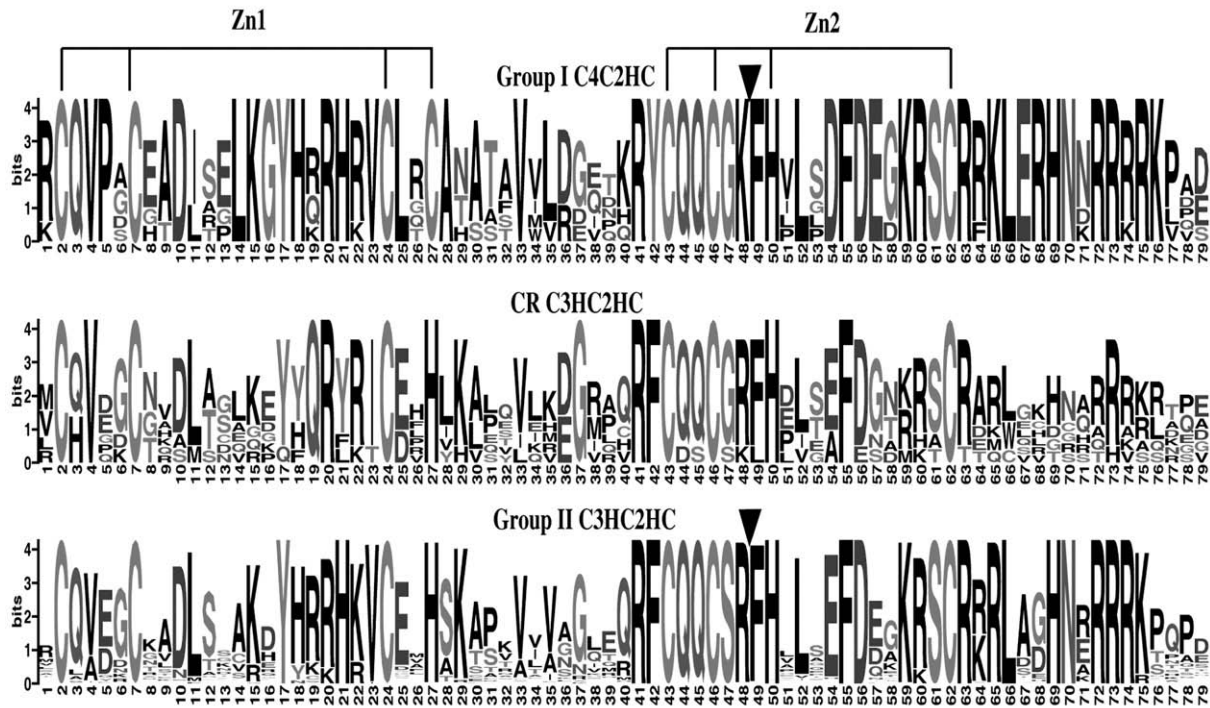


Fig. 2. Sequence logo of the DNA-binding domains of three different groups of SBP-box genes. The two zinc finger motifs are indicated by connected lines. The first zinc finger consists of four Cys residues in group I and Cys3His in group II and the CR group. The second zinc finger in all three groups has a C2HC motif. The triangle between residues 48 and 49 of group I and II refers to the intron splicing site. The overall height of each stack indicates the sequence conservation at that position, whereas the height of symbols within each stack reflects the relative frequency of the corresponding amino acid.

Each species from *P. patens*, *S. moellendorffii*, spruce, to *Arabidopsis*, rice and maize had only one group I SBP-box gene except for poplar which had 2 members.

Group II contained the majority of the SBP-box genes from land-plants and it was further grouped into 7 subgroups (IIa–IIg) with high statistical supports (Fig. 1). Subgroup IIa and IIb contained genes from all land-plants. About half moss and lycophyte SBP-box genes fell into subgroup IIb which contained only one *Arabidopsis* and three rice genes. Subgroup IIc contained only vascular plants genes, and genes in subgroup II d–II f were all from seed plants. All three genes in subgroup IIg were from moss. The five gymnosperm SBP-box genes were classified into subgroup IIb and II d. In most subgroups of group II, we could find two or more poplar SBP-box genes corresponding to one *Arabidopsis* gene, and more than one maize SBP-box genes grouped with one rice gene. SBP-box genes from the same lineage such as moss, lycophyte, gymnosperms and angiosperms tended to be clustered together.

3.3. Phylogenetic and gene structure analyses of *Arabidopsis* and rice SBP-box genes

We used the ML method to construct phylogenetic tree based on the SBP-domain amino acid sequences (Fig. 3a) of the 16 *Arabidopsis* and 18 rice SBP-box genes. The topology was similar to that constructed with 120 SBP-box genes from all lineages. The two genes of group I were grouped as one clade with high statistical support and the genes of group II were clustered into 6 subgroups (IIa–II f). Furthermore, we made an analysis for the exon–intron structure of the *Arabidopsis* and rice SBP-box genes (Fig. 3b). Our results showed that genes in the same subgroup had similar exon–intron structure except for *OsSBP1* in group IIc and *OsSBP4* in subgroup II f. Genes in group I and subgroup IIa, except for *OsSBP6*, all had 10 exons and genes in other subgroups, except for *OsSBP1*, all contained less than 4 exons; while *OsSBP1* and *OsSBP6* contained 11 exons. The gene structure of *OsSBP1* was similar to that of subgroup IIa rather than subgroup IIc

(Fig. 3b). Proteins of subgroup IIa and *OsSBP1* all had a long C-terminus with conserved sequences of more than 600 amino acid residues (Fig. 3c). Sequence comparison showed that amino acids encoded by the third and fourth exon of *OsSBP1* and *OsSBP6* were similar to the sequence encoded by the third exon of other genes in subgroup IIa. Genes in subgroup II f all had 4 exons except for *OsSBP4*, which missed one exon and one intron.

Our analyses demonstrated that the DNA-binding domain of all land-plants was encoded by two exons except for moss *PpSBP2*. The intron position was highly conserved with the splicing site before the dipeptide Phe-His of the conserved CQQC[S/G][R/K]FH octapeptide. The intron phases of the two exons encoding the SBP-domain were also conserved. We found that most of the rice SBP-box genes had long introns. The average intron length of *Arabidopsis* SBP-box genes was 124bp, shorter than that of the whole *Arabidopsis* genome (168 bp) calculated from *Arabidopsis* genome TAIR6 release, while the average intron length of rice SBP-box genes was 520 bp, longer than that of the whole rice genome (393 bp) calculated from TIGR rice genome release 4.0. Most of them are putative miniature inverted-repeat transposable elements (MITEs) or retrotransposons which can be found in the TIGR *Oryza* Repeat Database (<http://rice.tigr.org/tdb/e2k1/osa1/blastsearch.shtml>).

3.4. MicroRNA target site and conserved motifs in SBP-box genes

Rhoades et al. (2002) reported that a complementary site of miRNA *miR156/157* in some SBP-box genes and predicted 10 *Arabidopsis* SBP-box genes as potential *miR156* targets. We also found this conserved functional site either in the last coding exon or the 3′-UTR of subgroups IIc–II f *Arabidopsis* and rice SBP-box genes (Table 2). The miRNA target site was almost completely conserved except for 1 mismatch in *OsSBP3*. The miRNA target sites in exons of different genes all encoded a conserved peptide ALSLLS. BLAST search indicated that this conserved hexapeptide also existed in SBP-box genes of other species, such as maize and *Medicago truncatula*. *AtSPL3/4/5* and *OsSBP13* in subgroup II d had only two exons and this miRNA target

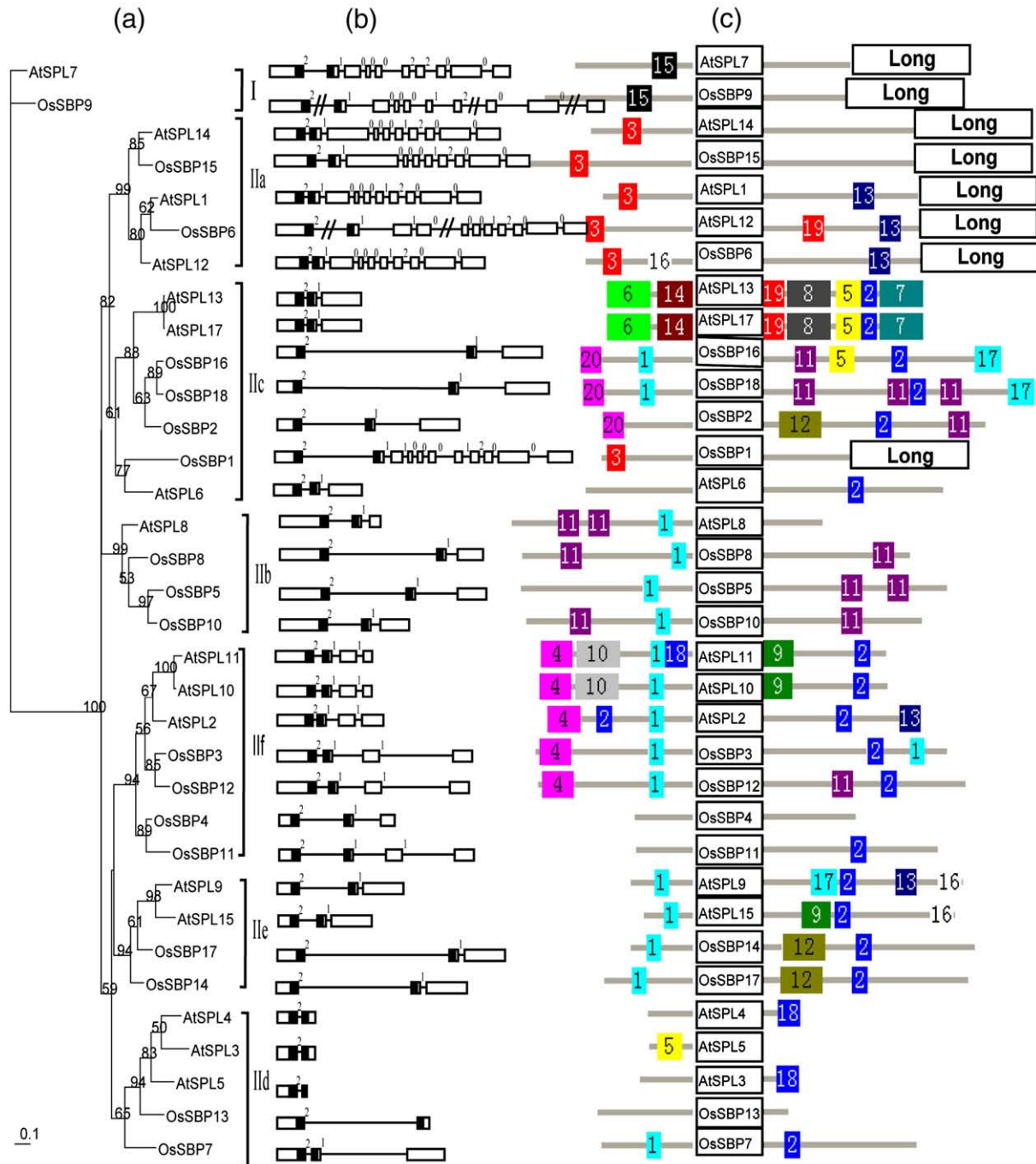


Fig. 3. Clustering of 16 *Arabidopsis* and 18 rice SBPs by three different approaches (a) Maximum-likelihood phylogenetic analysis reconstructed by Molphy. Scale bar corresponds to 0.1 amino acids substitutions per residue. (b) Exon and intron structure. Filled boxes: SBP-domains; white boxes: other exon regions; lines: introns. Numbers 0, 1, and 2: intron phases. The length of the boxes and lines are scaled based on the length of genes except for *OsSBP6* and *OsSBP9* with long introns denoted by “//”. (c) MEME motif search results aligned based on the DNA-binding domain represented as white boxes with gene names. Conserved motifs are indicated in numbered color boxes. Boxes marked “Long” indicate the long C-terminus.

site located in their 3'-UTRs. *OsSBP4* and *OsSBP11* was a duplicated gene pair and the miRNA target site located in the 3'-UTR of *OsSBP4* and the last exon of *OsSBP11*.

We made MEME search for conserved protein motifs flanking the SBP-domains and the results revealed some conserved motifs among several subgroups (Fig. 3c). Proteins in the same subgroup shared similar number and pattern of conserved motifs. Fig. 4 shows the sequence logo of the top four conserved motifs. Table 3 lists the occurrence of these 4 motifs in *Arabidopsis*, rice, moss and lycophyte. Motif 1 was found in most subgroups except for subgroup IIa of *Arabidopsis* and rice. Motif 2 encoded by the miRNA complementary

sequence also existed in moss subgroup IIg and lycophyte subgroup IIc. Motif 3 was only found in subgroup IIa of *Arabidopsis* and rice. Motif 4 was predicted in subgroup IIg of moss and subgroup IIIf of *Arabidopsis* and rice. Physiological function of these motifs remains to be investigated though the potential miRNA target site of motif 2 was reported.

Proteins in group I, subgroup IIa and *OsSBP1* contained a long C-terminus. A database search revealed that the protein sequences of the long terminus were SBP specific. All these SBP-box genes with long C-terminus including moss and lycophyte SBP-box genes in group I, together with CrSPL1 shared a highly conserved 59 AA motif about 120

Table 2
Location of miRNA target site on subgroups in group II of *Arabidopsis* and rice SBP-box genes

Gene name	Subgroup	No. of exons	Location
<i>AtSPL6</i> , <i>AtSPL9</i> , <i>AtSPL13</i> , <i>AtSPL15</i> , <i>AtSPL17</i>	c, e	3	Exon 3
<i>OsSBP2</i> , <i>OsSBP14</i> , <i>OsSBP16</i> , <i>OsSBP17</i> , <i>OsSBP18</i>	f	4	Exon 4
<i>AtSPL2</i> , <i>AtSPL10</i> , <i>AtSPL11</i>	f	4	Exon 4
<i>OsSBP3</i> , <i>OsSBP11</i> , <i>OsSBP12</i>	f	3	3'-UTR
<i>OsSBP4</i>	f	3	3'-UTR
<i>OsSBP7</i>	d	3	Exon 3
<i>AtSPL3</i> , <i>AtSPL4</i> , <i>AtSPL5</i> , <i>OsSBP13</i>	d	2	3'-UTR

AA downstream the SBP-domain (see Supplementary material, Figure S3). This conserved motif had an intron at a conserved position. We observed an Ankyrin (IPR002110) motif about 500 AA downstream to the SBP-domain in subgroup IIa and *OsSBP1*, indicating that those domains might interact with other proteins to regulate gene transcription.

4. Discussion

4.1. Evolution of gene structure

The exon–intron structure of three moss SBP-box genes (*PpSBP1*: AJ968320; *PpSBP3*: AJ968318; *PpSBP4*: AJ968319) gave an evidence of exon–intron loss in the evolution of the SBP-box gene structure. *PpSBP1* contained two exons at the 5'-end flanking the SBP-box and *PpSBP4* had a short exon in the same region (Fig. 5c). On the other hand, *PpSBP3* of subgroup IIg did not have the corresponding exon at the 5'-terminus. Furthermore, *PpSBP3* had only two exons at the 3'-end flanking the SBP-box while both *PpSBP1* and *PpSBP4* had three exons in the same region. The gene structure of these three moss SBP-

Table 3
Distributions of 4 most conserved motifs in subgroups within group II of *Arabidopsis*, rice, moss and lycophte

	Subgroup ^a	Motif 1	Motif 2	Motif 3	Motif 4
<i>AtSPL</i> , <i>OsSBP</i>	a, b, c, d, e, f	b, d ^b , e, f	c, d, e, f	a	f
<i>PpSBP</i>	a, b, g	a, b, g	g		g
<i>SmSBP</i>	a, b, c	a, b, c	c		

^a All existed subgroups in each species.

^b In subgroup d, motif 2 represents the miRNA target site in the 3'-UTR but not to the translated version as depicted in Fig. 4.

box genes may provide some hints for the gene structure of group IIa and other subgroups in group II in angiosperms. The ancestor SBP-box gene in land-plants might have some exons at both 5'- and 3'-termini flanking the SBP-box. One or two exons upstream of the SBP-box coding region had remained in some moss genes, but they were all lost in angiosperms. The downstream exons of the SBP-box coding region after the SBP-box might also have suffered from exon loss events and the number of exons was reduced from 8 (group I and IIa) to 2–3 in moss group II and 0–2 in angiosperms. Genes of angiosperm subgroup IIc and moss *PpSBP3* had the same exon–intron structure and conserved motifs. Genes with three exons in subgroup IIe might have evolved by intron loss, while genes with two exons in subgroup IIe might have been formed by degenerating the last exon to the 3'-UTR which retained the miRNA target site. In subgroup IIc, *OsSBP4* was a duplicated gene of *OsSBP11* and its last exon was degenerated to the 3'-UTR which retained the miRNA target site. Based on the above evidence, we propose that the diversity of SBP-box gene structure was mainly caused by gene duplication followed by intron and exon loss and it is still an undergoing process in angiosperm SBP-box genes.

4.2. Origin and evolution of SBP-box genes

It has been proposed that SBP-box genes are plant specific (Cardon et al., 1999). Sequence similarity search against available EST databases and genome sequences including brown algae (taxid:2870), red algae (taxid:2763), blue-green algae (taxid:1117), and golden alga (taxid:2825) suggested that SBP-box genes existed only in green plants. The earliest SBP-box genes we identified in this study were from the genome sequence of *C. reinhardtii*, a model organism representing the Chlorophyta (green algae). Our results indicated that SBP-box genes were plant specific and might originate predating the divergence of the green algae and the ancestor of land-plants. Based on the results obtained from phylogenetic analysis, gene structure comparison and motif search, we propose a model to account for the evolution of the SBP-box gene family (Fig. 5a).

Our results showed that all SBP-box genes from land-plants were clustered into group I and II while all seven SBP-like genes identified from green alga fell into a separate clade (Fig. 1). In SBP-box genes of the land-plants, the intron position in the SBP-domain as well as the intron phases of the two exons encoding the SBP-domain was conserved. This conserved intron position of SBP-domain indicated that all land-plant SBP-box genes might have originated from a common ancestor. Phylogenetic analysis suggested that SBP-box genes diversified into group I and II before the land-plants started to diverge but after the divergence of green algae from the last common ancestor of land-plants. Interestingly, it has also been reported that the plant specific DOF TF family and the AP2 TF subfamily have the similar pattern of origination and evolution (Moreno-Risueno et al., 2007; Shigyo et al., 2006).

CrSPL1 and proteins of group I and subgroup IIa shared the similar pattern of a long C-terminus and a highly conserved motif (see Section 3.4 and Supplemental material, Figure S3). Genes of group I and IIa had many more exons than that of other subgroups with several continuous zero phase introns (Fig. 5b), which might be lost more easily (Roy and Gilbert, 2005). Based on the above evidence, we

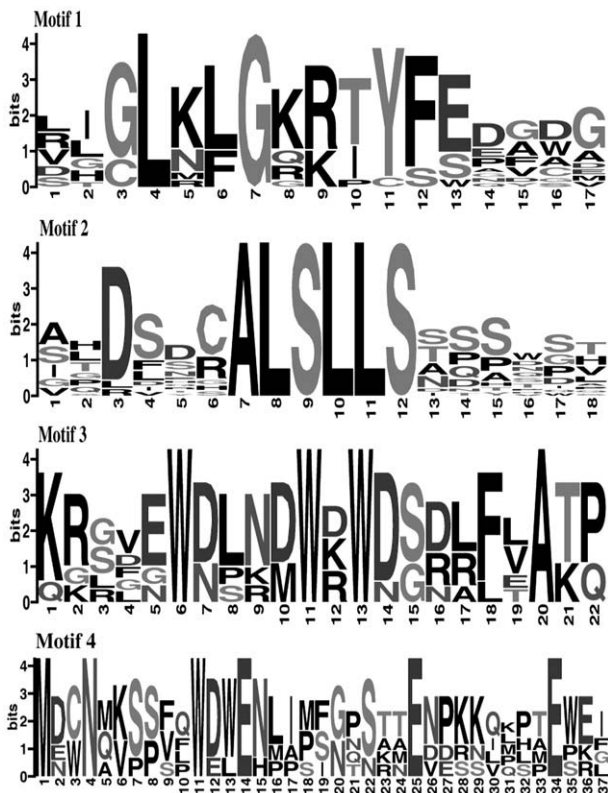


Fig. 4. Sequence logos of the top four conserved motifs. The overall height of each stack indicates the sequence conservation at that position, whereas the height within each stack reflects the relative frequency of the corresponding amino acid.

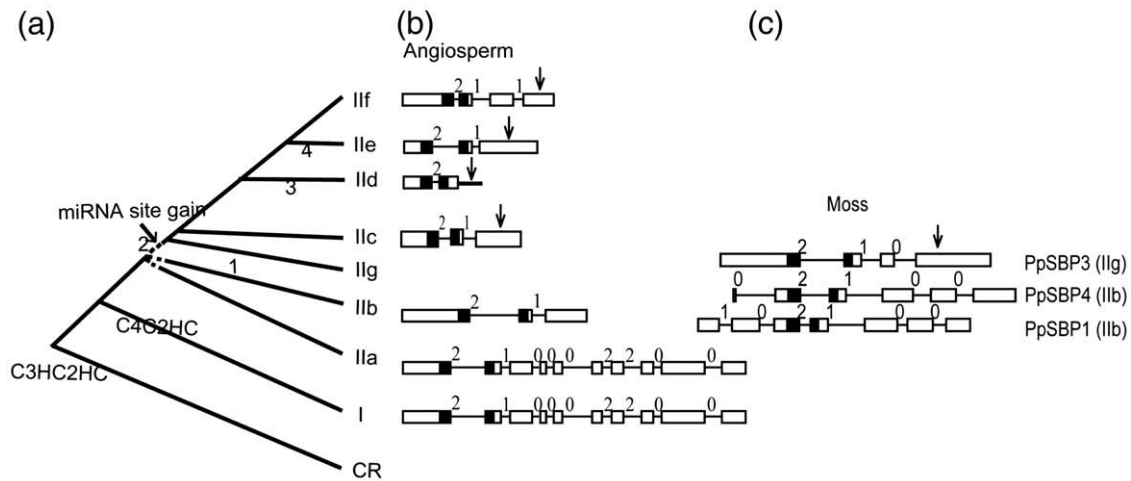


Fig. 5. A model for the evolutionary process of the SBP-box gene family (a) The three major groups (CR, I and II) and subgroups within group II are represented as branches. The CR clade consists of genes from green alga and the other clades include genes from land-plants only. C4C2HC and C3HC2HC represent two different zinc finger patterns. The gain of miRNA target site is marked by arrow. The numbers 1, 2, 3 and 4 on branches denote the intron–exon loss events. The dash line indicates the uncertain relationship among these clades. (b) Gene structure of group I and group II SBP-box genes of angiosperms. (c) Gene structure of 3 moss SBP-box genes (subgroup Ilb and Ilg). In (b) and (c), numbers 0, 1 and 2 indicate the intron phases. The arrow shows the microRNA target site.

assume that the SBP ancestor gene of land-plants might have a complex gene structure with many exons and two zinc fingers C3H and C2HC. It then duplicated and diverged into two ancestor genes of group I and II. The ancestor genes of group I have evolved to the group I genes by altering the first zinc finger from C3H to C4. The group II ancestor gene might have duplicated into three copies. The first copy might evolve to subgroup Ilb by losing exons and introns. The second copy might evolve to subgroup Ila and the third copy might be the ancestor of subgroups Ilc–Ilg. As group I and subgroup Ila and Ilb contain genes from all land-plants (Fig. 1), we infer that these duplication events might predate the divergence of moss and vascular plant lineage.

Genes of subgroups Ilc–Ilg all had a conserved miRNA target site in their exon or 3'-UTR. We suggest that the miRNA target site might exist in the ancestor of these subgroups. The ancestor gene of subgroup Ilc–Ilg might suffer intron and exon loss and obtain a miRNA target site, then evolve into three clades: Ilc, Ilg and Ild–Ilf (Fig. 1). Subgroup Ilc contained genes of both lycophyte and angiosperms, while subgroups Ild–Ilf contain genes from seed plants only and subgroup Ilg consisted of moss genes only. Both subgroup Ilf and Ilg had motif 4, whereas motif 1 was found in subgroup Ile–Ilg (Table 3). For *Arabidopsis* and rice, genes of subgroup Ild–Ilf had 2–3, 3 or 4 exons, respectively. We deduce that the ancestor gene of subgroup Ilc–Ilg evolved into subgroup Ilg in moss, subgroup Ilc in vascular plants and subgroup Ild–Ilf angiosperms by exon and intron loss.

4.3. Duplication of SBP-box genes

Duplication at both gene and genome levels has been and continues to be a pervasive process and contributes to the origin of biological novelty in evolution (Adams and Wendel, 2005). Gene duplication in angiosperm has been reported in many TF families, such as AP2, MADS, DOF, etc. (Zahn et al., 2005; Moreno-Risueno et al., 2007; Shigyo et al., 2006). Some duplicated SBP-box gene pairs (*AtSPL10* and *AtSPL11*, *AtSPL4* and *AtSPL5*, *AtSPL1* and *AtSPL12*, *OsSBP10* and *OsSBP5*, *OsSBP11* and *OsSBP4*, *OsSBP12* and *OsSBP3*) in *Arabidopsis* and rice have been found in genome analyses (Bowers et al., 2003; Blanc and Wolfe, 2004; Paterson et al., 2004; Wang et al., 2005b). Our analyses demonstrated that SBP-box genes duplicated and diversified in all species during their evolution. SBP-box genes from the same lineage tended to be clustered together in the phylogenetic tree, suggesting that they duplicated after the divergence of the lineages such as moss, lycophyte, gymnosperms and

angiosperms (Fig. 1). In most subgroups of group II, two or more poplar SBP-box genes were found along with one *Arabidopsis* gene indicating that SBP-box genes in poplar experienced duplications after the divergence of poplar and *Arabidopsis*.

4.4. Function divergence of SBP-box genes

The difference of exon–intron structure and the divergence of amino acid sequence among different subgroups provide us with some hints that SBP transcription factors may have a variety of physiological functions. To date, several important and divergent biological processes regulated by SBP-box genes have been reported, such as flower and fruit development (Klein et al., 1996; Cardon et al., 1997; Wang et al., 2005a; Manning et al., 2006), architecture formation (Becraft et al., 1990; Unte et al., 2003; Stone et al., 2005), sporogenesis (Unte et al., 2003), response to copper and fungal toxin (Eriksson et al., 2004; Stone et al., 2005), as well as control of GA level (Zhang et al., 2006).

Each species in land-plants had only one SBP-box gene in group I except for poplar in which two members were predicted. All group I genes were very conserved and clustered into a separate clade with high statistical support on the phylogenetic tree. The *Arabidopsis* member *AtSPL7* in group I was found to have the highest expression intensity in xylem obtained by Gene Atlas on Genevestigator (<https://www.genevestigator.ethz.ch/at/>). It would be interesting to explore the exact role of group I SBP-box genes in all green plants by functional characterization.

The variety of subgroups within group II reflected a big spectrum of structural and functional diversity of this group. We found 6 moss and 6 lycophyte SBP-box genes but only one *Arabidopsis*, 2 poplar and 3 rice genes in subgroup Ilb. *AtSPL8* was the only member of *Arabidopsis* gene in subgroup Ilb and was reported to involve in sporogenesis (Unte et al., 2003). *AtSPL3*, a member of subgroup Ild is a putative regulator of the MADS-box TF genes and constitutive expression of *AtSPL3* results in early flowering (Cardon et al., 1997). Three *AtSPL3* homologs in *A. majus* (*AmSBP1* and *AmSBP2*) and silver birch (*BpSPL1*) bind to MADS-box gene regulating flower development (Huijser et al., 1992; Klein et al., 1996; Lannenpaa et al., 2004). A tomato *AtSPL3* homolog (*LeSPL-CNR*) is critical for normal fruit development and ripening (Manning et al., 2006). These evidences suggest that SBP-box genes in this subgroup of seed plants play a critical role in flower and fruit development through regulating MADS-box genes.

4.5. Conservation of miRNA target site in SBP-box genes

MicroRNAs play important roles in gene expression regulation and miRNA targets have been found in many TF families, including SBP, MYB, NAC, ARF, CCAAT, GRAS, and AP2 (Rhoades et al., 2002). For example, in the SBP-box gene family, tissue-specific interactions between *OsmiR156* and *OsSBP* target genes were found in rice (Xie et al., 2006), and moss *PpSBP3* has also been reported to contain a *miR156* target site (Arazi et al., 2005). *MiR156* over expression causes a moderate delay in flowering and a severe decrease of apical dominance through regulating SBP-box genes (Schwab et al., 2005). In the AP2 TF family, the *miR172* target site was conserved in gymnosperm and angiosperm of AP2 homologs (Shigyo et al., 2006). In our case, a *miR156* target site was found in many SBP-box genes of moss, lycophyte, and angiosperms (Table 3), suggesting that the regulatory interaction between *miR156* and SBP-box genes exists before the divergence of moss from the vascular plants.

Our analysis showed that the SBP-box genes with the miRNA target site existed across many subgroups (IIc–IIlf) in angiosperms, suggestive of the conservation of the miRNA target site because of its functional importance. More importantly, this miRNA target site would move to 3'-UTRs of genes when exons with this site degenerated. Interestingly, we found only one or few genes in moss and lycophyte but many in angiosperms with this target site, indicating that miRNA regulation is more prevalent in angiosperms than other lineages. All these suggest that the regulation of miRNA on TFs is an ancient and important regulatory mechanism.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.gene.2008.03.016.

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