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Identification of a Consensus DNA-Binding Site for the Arabidopsis thaliana SBP Domain Transcription Factor, AtSPL14, and Binding Kinetics by Surface Plasmon Resonance

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Abstract
Proteins with a conserved Cys- and His-rich SQUAMOSA promoter binding protein (SBP) domain are transcription factors restricted to photosynthetic organisms that possess a novel two Zn-finger structure DNA-binding domain. Despite the fact that altered expression of some SBP-encoding genes has profound effects on organism growth and development, little is known about SBP domain protein target genes. Misexpression of the Arabidopsis thaliana AtSPL14 SBP domain gene confers resistance to programmed cell death and modifies plant architecture. A consensus DNA-binding motif for AtSPL14 was identified by systematic evolution of ligands by exponential enrichment (SELEX) or random binding site selection (RBSS). DNA recognized by AtSPL14 contained the core binding motif (GTAC) found for other SBP domain proteins, but mutational analyses indicated that at least one additional flanking nucleotide is necessary for effective AtSPL14–DNA interaction. Comparison of several SBP domain amino acid sequences allows us to hypothesize which specific amino acids might participate in this sequence-specific DNA recognition. Electrophoretic mobility shift assays (EMSA) with mutant AtSPL14 DNA-binding domain proteins indicated that not all of the Zn²⁺ ion coordinating ligands in the second Zn structure are strictly required for DNA binding. Surface plasmon resonance (SPR) was used to evaluate AtSPL14 in vitro binding kinetics for comparison of equilibrium binding constants with other SBP domain proteins. These data provide a strong basis for further experiments aimed at defining and distinguishing the sets of genes regulated by the closely related SBP domain family members.

Abbreviations: CuRE, copper-responsive element; dsDNA, double-stranded DNA; DTT, dithiothreitol; EDTA, ethylenediaminetetraacetic acid; EMSA, electrophoretic mobility shift assay; FB1, fumonisin B1; fbr, FB1-resistant; HEPES, N-(2-hydroxyethyl)piperazine- N'-2-ethanesulfonic acid; IPTG, isopropyl β-D-1-thiogalactopyranoside; NMR, nuclear magnetic resonance; PCD, programmed cell death; PMSF, phenylmethylsulfonyl fluoride; RBSS, random binding site selection; RU, resonance (response) units; SA, streptavidin; SBP, SQUAMOSA promoter binding protein; SDS, sodium dodecyl sulfate; SELEX, systematic evolution of ligands by exponential enrichment; SPL, SQUAMOSA promoter binding protein-like; SPR, surface plasmon resonance; WT, wild type.

Proper growth and development of multicellular organisms depend on a delicate balance between cell proliferation and programmed cell death (PCD). In plants, PCD is required for tracheary element differentiation to form the water-conducting xylem tissue and accurate formation of various reproductive organs (1-5). PCD is also an important aspect of plant defense against pathogen attack (6-8). Despite the essential nature of PCD, large gaps remain in our knowledge of the mechanistic details and molecular components controlling plant PCD. Therefore, to identify novel genes that might participate in plant PCD, we developed a mutant selection scheme for the model plant Arabidopsis thaliana using a PCD-inducing fungal toxin fumonisin B1 (FB1), to identify FB1-resistant (fbr) mutants (9-11). Misexpression of the AtSPL14 gene in the fbr6 mutant confers the ability to proliferate in the presence of FB1 and modifies normal plant architecture, linking the insensitivity to cell death to altered plant development (9). AtSPL14 has features of a DNA-binding transcription factor, including a Cys- and His-rich SQUAMOSA promoter binding protein (SBP) domain predicted to bind DNA, ankyrin repeats that mediate protein–protein interactions, nuclear localization, and ability to bind to A. thaliana genomic DNA (9).

SBP domain proteins are defined by a conserved approximately 80 amino acid–protein domain (the SBP domain or SBP box) found only in proteins from photosynthetic organisms, ranging from single-celled algae (e.g., Chlamydomonas reinhardtii) to higher plants (e.g., A. thaliana and Oryza sativa). The SBP gene families are comprised of 16 genes in A. thaliana and 19 genes in O. sativa that encode proteins that share the highly conserved SBP DNA-binding domain but are diverse in overall domain structure (12, 13). To date, only a few functions for SBP domain proteins have been reported, perhaps due to genetic redundancy of closely related family members. In all cases, SBP domain proteins have been implicated in various aspects of plant growth and development, including metal sensing in algae and directing development of leaves, embryos, and floral organs in higher plants (9, 13-22). Some of these SBP do-
main proteins may also be posttranscriptionally regulated by noncoding microRNAs to control their spatial and temporal expression (23-25). Despite the obvious importance of these proteins, the specific genes targeted by individual SBP domain family members and the molecular consequences of their actions are largely unknown.

To fully understand the physiological functions of AtSPL14 in regulating plant PCD and/or development, AtSPL14 target genes, their modes of regulation, and the consequences of expression/repression need to be determined. As one step toward that goal, we used an affinity-based assay, referred to as systematic evolution of ligands by exponential enrichment (SELEX) or random binding site selection (RBSS) to screen a random pool of dsDNA fragments for sequences capable of binding to recombinant AtSPL14 protein. From this analysis, an AtSPL14-binding consensus DNA motif was derived. Mutual analyses indicated that predominantly the core motif, CGTAC, is essential for AtSPL14 protein binding to the DNA in vitro.

Recent structures of SBP domains determined by nuclear magnetic resonance (NMR) indicate that SBP domains form a unique two Zn-finger structure DNA-binding domain (26, 27). The SBP domain contains eight absolutely conserved Cys or His residues, some of which are critical for SBP domain DNA binding (28). We determined that all of the highly conserved cysteines in the two Zn$^{2+}$ ion binding structures of the AtSPL14 SBP domain binding are important for DNA recognition by electrophoretic mobility shift assays (EMSA) and surface plasmon resonance (SPR). Yet, an AtSPL14 SBP domain with one of the Cys residues in the second Zn-finger structure mutated retained some DNA-binding ability. Moreover, we monitored the kinetic features of the AtSPL14 SBP domain–DNA binding by SPR. We further compare and contrast the target sequences and the equilibrium binding constants we determined for AtSPL14 with those of other SBP domain proteins (18, 28, 29).

Materials and Methods

Recombinant AtSPL14 Protein Expression and Purification — Two different truncated and epitope-tagged versions of recombinant AtSPL14 proteins were used. The full-length AtSPL14 cDNA was generated by reverse transcription–polymerase chain reaction (RT-PCR) from RNA isolated from ecotype Col-0 (the reference genotype for the A. thaliana genome) using oligonucleotide primers SBPF (5'-GGATCCATGGAT

GAGGGTAGAGCTCAAGTG-3') and SBPR (5'-ACTAGTCCGGATCCGGATTGACCTATATCCACCTC-3') and verified by DNA sequencing (30). Engineered BamHI and existing internal Sall restriction enzyme recognition sites were used to subclone FBR6 short (FBR6s; aa 1–402) into BamHI/Sall-digested pET-28a(+) to produce a protein composed of an N-terminal His tag, thrombin cleavage site, and T7 tag fused to a region of AtSPL14 encompassing the DNA-binding domain (Novagen, EMD Chemicals, Inc., Darmstadt, Germany). The His-tagged FBR6s protein was 447 amino acids with a predicted molecular mass of 48.8 kDa.

For electrophoretic mobility shift assays (EMSA) and surface plasmon resonance (SPR) binding assays, recombinant epitope-tagged FBR6 supershort (FBR6ss; aa 111–200) protein was used. The conserved SBP domain of AtSPL14 was amplified by PCR using oligonucleotide primers oJS86 (5’-CC

GAATTCTCTCCGGAGGAATTATCCC-3’) and oJS87 (5’-CCGAATTCTTATGCAACCTCTCCGGCTGCCG-3’) and verified by DNA sequencing (30). The resulting PCR product was subcloned into pET-28a(+) to produce a protein composed of an N-terminal His tag, thrombin cleavage site, and T7 tag fused to a region of AtSPL14 encompassing the DNA-binding domain. The His-tagged FBR6ss protein was 126 amino acids with a predicted molecular mass of 14.3 kDa.

The recombinant proteins were expressed in Escherichia coli by inducing log phase cultures with 0.2 mM isopropyl β-d-1-thiogalactopyranoside (IPTG) at 37 °C for 2 h, and proteins were purified on Ni$^{2+}$-conjugated affinity resin according to the manufacturer’s instructions (ProBond Purification System; Invitrogen, Carlsbad, CA). For SELEX experiments proteins were retained on the resin.

Site-Directed Mutagenesis — The site-directed mutagenesis was performed according to the Stratagene’s Quick-Change site-directed mutagenesis kit instruction manual (Stratagene, Cedar Creek, TX). Oligonucleotide primers used to mutate the nucleotides in the AtSPL14-binding DNA and the cysteine residues of the AtSPL14 SBP domain are shown in Tables S1 and S2 of the Supporting Information, respectively.

Systematic Evolution of Ligands by Exponential Enrichment (SELEX) or Random Binding Site Selection (RBSS) — A random pool of oligonucleotides (76 nucleotides) of sequence 5’-GCTGCAGTTGACATCCTGATCCGGGACGACAGATCCGCTGAGCTGACGCGCT-3’, where N$_{26}$ represents 26 randomized nucleotides, was synthesized by equimolar incorporation of A, G, C, and T at each “N” position (Integrated DNA Technologies, Coralville, IA). The two sets of 25 nucleotides flanking the 26-nucleotide random core were designed for amplification by PCR. To make double-stranded DNA (dsDNA), the random pool of oligonucleotides (100 ng) was subjected to PCR using the forward primer and Taq polymerase enzyme (94, 68, and 72 °C, 1 cycle), and the PCR products were purified by agarose gel electrophoresis (1.5% MetaPhor agarose; Cambrex Bio Science Rockland, Inc., Rockland, ME) to yield ds-R76, the substrate in the initial binding reaction.

The binding reactions were carried out on ice essentially as described with a few modifications (31). The recombinant protein T7-His$_{6}$-FBR6s was purified using a His tag purification kit (ProBond; Invitrogen, Carlsbad, CA) and retained on the resin. The resin with immobilized protein (200 μL) was washed twice with binding buffer (20 mM Tris-HCl, pH 7.6, 50 mM NaCl, 1 mM MgCl$_2$, 0.2 mM EDTA, 5% glycerol, 0.5 mM DTT, 50 μM PMSF), mixed with 50 μg/mL poly(dIdC) (Amersham Biosciences, Cleveland, OH) to reduce nonspecific binding for 10 min, and incubated with ds-R76 DNA (200 ng) for 60 min, with gentle tapping every 10 min. The immobilized protein–DNA complexes were washed with TN buffer (10 mM Tris-HCl, pH 7.4, 150 mM NaCl) five times, and the DNA was eluted with 200 μL of dissociation buffer (0.5 M Tris-HCl, pH 7.4, 20 mM EDTA, 10 mM NaCl, 0.2% SDS). The bound DNA was amplified by PCR using 250 nM oligonucleotide primers that anneal to the defined terminal sequences of the 76-nucleotide oligonucleotide for 20 cycles. The resulting product was used as the substrate in the second round of SELEX. After five serial selection rounds, the amplified DNA was cloned into vector pGEM-TEasy (Promega, Madison, WI) and subjected to DNA sequencing.
The consensus target site for the AtSPL14 SBP DNA-bind

Electrophoretic Mobility Shift Assay (EMSA) — DNA fragments were fluorescently labeled by PCR amplification in a reaction containing 100 nM dNTPs, 1× PCR buffer, 1.5 mM MgCl2, 2 ng of DNA template (pGEM-TEasy subclones selected from SELEX), 300 nM 5′-IR-dye 700 GTACCTTC-GTTGCCGCTAG-3′ corresponding to the T7 promoter (Li-Cor, Lincoln, NE), 300 nM primer R or primer F, and Taq polymerase enzyme. The resulting PCR product was quantified using gel electrophoresis. Electrophoretic mobility shift assay binding reactions were performed in the same binding buffer used in the SELEX (exception: 1 mM DTT and no PMSF) in a total volume of 20 nL containing 60 nM protein and 4 nM labeled DNA and incubated for 30 min at 25 °C in darkness prior to electrophoresis. The protein−DNA binding mixture was electrophoretically resolved for 45 min at 4 °C in the dark on a prerun 8% nondenaturing polyacrylamide gel (polyacrylamide-bis ratio = 37.5:1) in Tris−borate−EDTA buffer (89 mM Tris, 89 mM boric acid, and 2.5 mM EDTA) at constant voltage (15 V/cm). Gels were analyzed using an Odyssey Infrared Imager (Li-Cor, Lincoln, NE).

Immobilization of Biotinylated DNA on the SA Sensor Chip — Cognate and noncognate DNA fragments (156 bp) containing dsR76 used in the SELEX experiments were biotinylated by PCR amplification with dsDNA cloned in pGEM-TEasy as templates, 5′-biotin-GTACCTTCGTTGCCGCTAG-3′ oligonucleotide corresponding to the T7 promoter (IDTDNA, Cedar Rapids, IA), and either primer R or primer F. The PCR products were purified by the QIAquick PCR purification kit (Qiagen, Valencia, CA), and DNA was immobilized on streptavidin (SA) sensor chips (Biacore AB, Uppsala, Sweden).

5′-Biotin-labeled DNA (the random binding site selected DNA 14 containing the consensus FBR6ss binding site, CGTAC, Figure 1) was immobilized onto the SA sensor chip surface [450 resonance (response) units, RU] to provide the maximal level of RU associated with protein binding in the range of 50–100 RU. SA chips were conditioned with three consecutive 1 min injections of 1 M NaCl and 50 mM NaOH prior to immobilization. Two reference cells were used as controls for background subtraction: reference 1 was the surface alone (no bound DNA), and reference 2 had a noncognate DNA (a random DNA that did not possess the consensus binding motif) immobilized to the reference cell at the same concentration as the cognate DNA.

Surface Plasmon Resonance (SPR) Analysis — The purified His-FBR6ss protein was desalted and equilibrated in HBS-EP buffer using Centricon-mediated centrifugal filtration (10 kDa cutoff; Millipore, Billerica, MA). The protein was diluted in HBS-EP buffer to yield several different concentrations ranging from 3.25 to 300 nM. Varying protein concentrations were injected at a 75 µL/min flow rate. The chip surface was regenerated by injection of 0.1% SDS and 3 mM EDTA buffer for 1 min after each protein injection. Responses from the reference cell(s) were subtracted to correct for refractive index changes and nonspecific binding.

SPR Data Analysis — Data were analyzed with BIAevalu- 

Figure 1. The consensus target site for the AtSPL14 SBP DNA-binding domain determined by SELEX. Double-stranded DNA molecules containing a 26-nucleotide completely randomized central region were subjected to repetitive cycles of binding to recombinant AtSPL14 bound to a Ni2+-chelating affinity resin and PCR amplification. The individual binders were subsequently tested by electrophoretic mobility shift assays (EMSA) and competition with unlabeled probe (as in Figure 2A), yielding 20 distinct dsDNA fragments capable of binding to the AtSPL14 SBP domain. (A) An alignment of the 20 individual binders based on the results of the web-based multiple expectation maximization for motif elicitation (MEME) analysis program (36). The core consensus sequence nucleotides are in color, the adjacent random nucleotides are in black, and the nonrandom nucleotides derived from the fixed sequence immediately flanking the random nucleotides in the dsDNA pool are in gray. (B) A representation of the consensus DNA target binding motif (disregarding the fixed nucleotides flanking the random core) using WebLogo (37). The degree of conservation is indicated by the height of the letters. The core sequence “CGTAC” was found in all dsDNA-binding fragments.
Results

Identification of a Consensus Binding Site for the AtSPL14 SBP Domain by Systematic Evolution of Ligands by Exponential Enrichment (SELEX) — SELEX or random binding site selection (RBSS), an in vitro oligonucleotide binding and PCR amplification method, was used to define consensus DNA-binding sequences for the AtSPL14 SBP DNA-binding domain (33-35). The AtSPL14-binding DNA was selected from a pool of 76 bp of double-stranded DNA (dsDNA) with a central core of 26 random nucleotides by repeated cycles of binding to the hexahistidine-tagged FBR6s protein (encompassing the SBP DNA-binding domain) immobilized on Ni²⁺-chelating affinity resin. Fifty-seven individual clones that were derived from five serial rounds of selection were subcloned and subjected to DNA sequencing, revealing that identical clones were identified multiple times. All of the unique clones were tested for binding to FBR6ss using EMSA assays and competition assays.

Figure 2. Specific AtSPL14 DNA binding by electrophoretic mobility shift assays (EMSA) and competition assays. (A) An example of the initial competition assays by EMSA used to determine binding of the individual dsDNA fragments identified by SELEX. The double-stranded DNA probe was generated by PCR with IRDye 700 fluorescently labeled (probe) or unlabeled (competitor) oligonucleotide primers. The probe (4 nM) was incubated without (−) or with (+) recombinant His-tagged AtSPL14 SBP domain protein (FBR6ss, 60 nM). For testing specificity, increasing amounts of unlabeled competitor were included in the binding reaction; the ratio of competitor:probe is indicated. “Complexes” were separated on an 8% nondenaturing polyacrylamide gel and visualized by infrared imaging. Lanes: (1) free probe; (2) probe plus AtSPL14 SBP domain protein; (3-6) probe plus increasing amounts of unlabeled competitor. (B) For EMSA competition assays, single nucleotide substitutions in the core consensus binding site of a selected dsDNA-binding fragment were generated by site-directed mutagenesis. The core consensus motif is in color with flanking nucleotides in black for the wild-type (wt) dsDNA, and individual nucleotide changes for the mutated (M1–M13) dsDNA are indicated. (C) EMSA competition assays with the wild-type (WT) or mutated (M1–M13) dsDNA fragments. For the binding reactions, “FP” indicates free probe with no protein, “0” indicates no competitor, and the triangles represent increasing amount of competitor in the binding reaction (20× or 40× molar ratios). (D) Band intensities corresponding to bound complexes were determined by infrared imaging (Odyssey; Li-Cor, Lincoln, NE). Binding efficiencies were normalized to a control binding reaction with no competitor on each gel (“0” in panel C). Binding levels with a 20× molar ratio (black bars) and a 40× molar ratio (white bars) of unlabeled competitor are shown. Error bars represent 95% confidence levels from experiments performed in triplicate.
Consensus DNA-Binding Site for the A. thaliana SBP Domain Transcription Factor, AtSPL14

Figure 3. Global analysis of SPR biosensor data for AtSPL14 SBP domain–DNA interaction kinetic analyses. Top panels: The dotted lines represent SPR sensograms (resonance units, RU) obtained by injecting different concentrations of the AtSPL14 SBP domain protein (5, 15, 20, 25, 30, and 50 nM; bottom to top) onto a SA sensor chip coated with a representative 156 bp cognate dsDNA fragment identified by SELEX that bound in EMSA competition assays. Binding data were collected at a flow rate of 75 µL/min. Signals from the control reference cell (coated with a non-cognate DNA) were subtracted. Bottom panels: Residual plot showing the difference between measured and calculated responses. (A) Best fits of the binding data to a simple bimolecular 1:1 Langmuir binding model are represented by solid black lines. A residual plot showing the difference between measured and calculated responses indicated that the 1:1 Langmuir binding model is not a good fit (±10 RU). (B) Best fits of the binding data to a simple bimolecular 1:1 mass transfer binding model are represented by solid black lines. A residual plot showing the difference between measured and calculated responses indicated that the 1:1 mass transfer binding model is a good fit (±3 RU). Kinetic binding constants were determined for the mass transfer binding model: the association rate constant $k_a = 2.5 \times 10^7 \text{ M}^{-1} \text{s}^{-1}$, the dissociation rate constant $k_d = 7.4 \times 10^{-2} \text{s}^{-1}$, and the equilibrium binding constant $K_s = k_a/k_d = 3.3 \times 10^8 \text{ M}^{-1}$.
ple 1:1 binding model (Figure 4), resulting in an equilibrium binding constant $K_A = 3.8 \times 10^8$ M$^{-1}$. Experimental error was determined by performing the experiment in triplicate; $K_A = 3.1 \times 10^8 \pm 1.0 \times 10^8$ M$^{-1}$ (SEM). Therefore, the binding affinities obtained from determining both kinetic rate constants and steady-state equilibrium analyses were similar.

**Amino Acids Required for AtSPL14 SBP Domain−DNA Interactions** — Multiple sequence alignments of SBP domains from *A. thaliana*, *Antirrhinum majus*, and *C. reinhardtii* indicate that this is a highly conserved protein domain, with sequence identity ranging from 50% to 96% (Figure S2 of the Supporting Information). The consensus sequence is $\text{CX}_4\text{CX}_8\text{YX}_3\text{HX}_4\text{CX}_{16}\text{RXCQQCX}_3\text{HX}_5\text{FDX}_3\text{SCRX}_2\text{LX}_2\text{H}-\text{NXRR}$, where “X” is not absolutely conserved. The six absolutely conserved Cys residues and two of the absolutely conserved His residues have been shown to coordinate two Zn$^{2+}$ ions to form a novel two Zn-finger structure (26, 27). The presumed coordinating amino acids of the AtSPL14 SBP domain are shown in Figure S2 of the Supporting Information. To explore whether these six Cys amino acids are required for the AtSPL14 SBP domain to bind DNA, all six amino acids were individually mutated to Ala and Ser, and the mutant proteins were assayed for DNA-binding efficiency by EMSA. All 12 Cys substitutions almost completely abolished the ability of the AtSPL14 SBP domain to bind DNA, with the exception of C164-A and C164-S, which exhibited very weak binding at high protein concentrations (Figure 5). We also tested the ability of the mutant proteins to bind DNA by SPR. The binding of the mutant AtSPL14 SBP domain proteins was markedly reduced compared to the wild-type protein (Figure 5). Therefore, all six highly conserved Cys amino acids are important for DNA binding, but weak affinity is retained when one of the Cys amino acids predicted to coordinate a Zn$^{2+}$ ion in the second Zn-finger structure is mutated.
Discussion

Our interest in determining the AtSPL14 SBP domain DNA-binding motif and binding kinetics derives from identification of an A. thaliana mutant with a disruption in the AtSPL14 gene (At1g20980). The A. thaliana FBJ-resistant (fbr6) mutant was originally identified by selecting for mutants capable of growth and development on media containing the fungal toxin fumonisin B1 (FBJ). FBJ disrupts sphingolipid metabolism in eukaryotes (38) and induces PCD, dependent on transcription, translation, reversible protein phosphorylation, light, and hormone signaling pathways in A. thaliana (10, 11). In addition to resistance to FBJ, the fbr6 mutant also exhibits altered plant architecture, including elongated petioles and enhanced leaf margin serration (9). Knowledge of the AtSPL14 DNA-binding site (and potentially regulated genes) is critical to fully understanding the physiological functions of this sequence-specific DNA-binding transcription factor.

The AtSPL14 SBP Domain Binds to the Core GTAC Consensus Target Site but Requires an Additional Flanking Nucleotide for Effective Interaction — AtSPL14 (At1g20980) encodes a 1035 amino acid–protein with an SBP DNA-binding domain (http://srs.ebi.ac.uk; IPR004333), a bipartite nuclear localization signal (aa 117–193), and ankyrin repeats that mediate protein–protein interactions (IPR002110; aa 821–941) in the C-terminal region of the protein (9, 13, 39, 40). The highly conserved SBP DNA-binding domain is a Cys- and His-rich region (consensus: CXX/CXX̅HXX̅HX̅4/CQQCX̅HX̅4(C)) found only in proteins from photosynthetic organisms (12, 13, 18, 28). The founding members of the SBP domain gene family (SBP1 and SBP2 from A. majus) were identified by their ability to bind to the upstream regulatory region of the SQUAMOSA floral meristem identity gene (29). Previous reports indicated that A. majus proteins and some A. thaliana SBP domain proteins bind DNA encompassing a ten nucleotide motif common to cis regulatory elements in the orthologous A. majus SQUAMOSA and A. thaliana APETALA1 gene promoters and a similar motif in the A. majus DEFH84 promoter in vitro (13, 17). Alignment of these sequences revealed a putative consensus DNA-binding motif of TNGTGACAA (13). It appears that GTAC is the core DNA-binding motif for all SBP domains described so far, but nucleotides flanking the core motif are preferred by different SBP domain proteins.

We used a completely random approach, referred to as systematic evolution of ligands by exponential enrichment (SELEX) or random binding site selection (RBSS), to identify the conserved DNA-binding site for AtSPL14. These analyses revealed CCGTAC(A/G) as the optimal binding site for the AtSPL14 SBP domain (Figure 1). We found no evidence for a palindromic binding motif, which would suggest that binding occurs as a dimer, consistent with a previous report that other SBP domains bind DNA with a 1:1 stoichiometry (26). To verify the importance of the consensus motif, we mutated the individual nucleotides of the predicted binding site CCGTAC(A/G) to obtain both conserved and nonconserved substitutions of each base except the 3′-end A/G (Figure 2B). Mutations in the CGTAC core markedly reduced competition for AtSPL14 binding, but the C at the 5′-end and the A/G at the 3′-end were dispensable (Figure 2). Therefore, each individual nucleotide of the core motif CGTAC is necessary for effective AtSPL14 SBP domain DNA binding (Figure 2).

Birkenbihl et al. (28) recently used a similar random binding site selection with G and T fixed at positions 7 and 8 (of 16 total “random” nucleotides) to identify binding sites for AtSPL3 and AtSPL8 SBP domains. A high preference for at least one more C at the 5′-end of the core consensus GTAC motif was found for AtSPL3 (83%), but not for AtSPL8. Therefore, the recognition site we determined for AtSPL14 is more similar to that of AtSPL3 than AtSPL8. The preference for a C flanking the GTAC core must be due to a specific amino acid–nucleotide interaction. The overall sequence identities between the AtSPL14 SBP domain and those of AtSPL3 (70%) and AtSPL8 (74%) are quite similar (Figure S2 of the Supporting Information), with only three amino acid differences in the Zn2 region proposed to interact directly with DNA (27). AtSPL14 and AtSPL3 have a Glu, Arg, and Gly whereas AtSPL8 has Asn, Lys, and Asp, respectively (Figure S2 of the Supporting Information). Whereas no simple universal code has been elucidated, specificity is imparted in most protein–DNA contacts by hydrogen bonding in the major groove (41). Statistical analysis of atomic interactions in 139 protein–DNA complexes analyzed from the Protein Data Bank (PDB) revealed favored amino acid–nucleotide pairs (42). The authors categorized direct amino acid–nucleotide contacts, including hydrogen bonds and electrostatic, hydrophobic, and other van der Waals interactions. Whereas contacts with the sugars or phosphates of DNA contribute to DNA–protein stability, only H-bonds to nucleotide bases can confer sequence specificity. In interactions with nucleotide bases, the most commonly observed interaction was Arg with G (42). Therefore, it seems most likely that the substitution of Lys in AtSPL8 for Arg in AtSPL14 and AtSPL3 renders the 5′ nucleotide flanking the core GTAC motif unimportant. This hypothesis will need to be tested experimentally.

Simple pattern matching searches for the consensus binding sequence reported for the A. majus SBP proteins “TNGTGACAA” (13, 29) upstream of annotated A. thaliana genes identified 331 and 640 putative SPL-regulated genes with the pattern within 500 and 1000 base pairs of the translation start site, respectively. Using the consensus DNA-binding motif, we identified for AtSPL14 (GTAC) more than 6000 genes with the motif within 500 base pairs of the translation start site as possible AtSPL14 targets, and one gene of unknown function has 16 occurrences. Combined with microarray gene expression data comparing the transcriptomes of the fbr6 mutant to wild-type plants, a subset of candidate target genes for AtSPL14 were identified (data not shown). Whereas some of these promoter sequences also bind to AtSPL14 in vitro, additional experiments, such as chromatin immunoprecipitation (ChIP), will be required to verify that these are true targets for AtSPL14 in vivo.

The Kinetic Binding Parameters of SBP Domains Differ — The SBP domain DNA binding is dependent on the presence of Zn2+ ions (18, 28). The NMR-resolved structures of SBP domains revealed that the SBP domain forms a novel two Zn structure DNA-binding domain (26). The two Zn2+ ions are coordinated by three Cys residues and one His residue, forming two Zn structures in CCCH (Zn1) and CCHC (Zn2) configurations (Figure S2 of the Supporting Information). We mutated all six highly conserved Cys to Ala and Ser and determined the SBP–DNA binding efficiency by EMSA and SPR. All 12 mutations markedly reduced the AtSPL14–DNA interaction (Figure 2), verifying that all six
conserved Cys in the AtSPL14 SBP DNA-binding domain are important for AtSPL14 protein–DNA interaction and consistent with the supposition that these Cys residues participate in Zn$^{2+}$ ion binding (26, 28). However, the C164-A and C164-S (corresponding to the second Cys in Zn2) mutant proteins retained some weak binding ability (Figures 2 and S2 of the Supporting Information). The second Cys and the His in Zn1 and the first and fourth Cys in Zn2 were reported to be critical for AtSPL1 DNA binding (28). In that study the second Cys in Zn2 was not mutated, but the His in Zn2 was found to be somewhat dispensable (28). Therefore, we independently determined that all of the Zn1 coordinating residues are essential, but two of the presumed Zn2 coordinating residues (the His and second Cys) are not absolutely required for SBP domain DNA binding. These findings support the conclusions derived from the NMR structure of a truncated SBP domain protein (AtSPL12). The Zn1 structure is critical for overall structure, and removal of part of the Zn2 domain affected DNA binding but had little effect on overall folding (27). It is not yet clear whether proteins with mutations of the Zn$^{2+}$ ion coordinating amino acids (His and second Cys) in the Zn2 structure can still bind a second Zn$^{2+}$ ion. Another His residue is absolutely conserved in SBP domains (His187 in AtSPL14, Figure S2 of the Supporting Information) and might serve as a substitute fourth ligand. Alternatively, acidic residues or a water molecule might serve as the fourth coordinating ligand (43).

The AtSPL14 DNA-binding kinetics was analyzed by surface plasmon resonance (SPR). The mass transfer limitation due to the rapid association (>10$^7$) and dissociation rates (>10$^{-2}$) made determination of reliable kinetic constants difficult using conventional SPR (32). Therefore, we also evaluated steady-state binding at equilibrium. The determined equilibrium binding constant, $K_a$ (3.8 × 10$^8$ M$^{-1}$), is similar to the reported values (9.6 × 10$^7$ M$^{-1}$ and >5 × 10$^8$ M$^{-1}$) for the AtSPL4 and AtSPL12 SBP domains, respectively, and 1 order of magnitude greater than the value (2.8 × 10$^7$ M$^{-1}$) reported for the AtSPL7 SBP domain (26, 27). Kinetic rate constants were not reported for other SBP domain proteins.

The different $K_a$ values determined for different SBP domains might be due to the different salt concentrations in the analyte. Our data were generated in HB$^+$-EP buffer (containing 150 mM NaCl), whereas the $K_a$ of binding for AtSPL4, AtSPL7, and AtSPL12 was assayed in the presence of 100 or 300 mM KCl. The $K_a$ for AtSPL12 and AtSPL4 at 100 mM KCl (>5 × 10$^8$ M$^{-1}$ and 2.8 × 10$^7$ M$^{-1}$, respectively) was higher than those determined at 300 mM KCl (3.2 × 10$^7$ M$^{-1}$ and 2.1 × 10$^7$ M$^{-1}$, respectively), suggesting that salt concentration affects SBP domain–DNA-binding affinity. Therefore, electrostatic forces contribute to the SBP–DNA interaction, as was also observed in other protein–DNA interactions (44, 45).

Conclusions — In summary, different SBP domain proteins display different binding affinities to the same DNA and different selectivity for DNA targets. Even though they contain the same core consensus binding motif (GTAC), the A. thaliana APETALA1 gene-derived DNA and DNA containing the C. reinhardtii copper-responsive element (CuRE) had different affinities for several SBP domain proteins (28). Our results revealed that AtSPL14–DNA binding is highly sequence selective and allow us to hypothesize which particular amino acids may confer that specificity. Moreover, the well-conserved SBP domains possess diverse DNA-binding affinities for similar DNA sequences. These observations will be useful, in conjunction with additional experimentation, to identify the gene targets for individual SBP domain family members to understand their physiological functions in the context of whole organisms.

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Supporting Information

SPR data analysis comparing the subtraction of the signals from a blank reference cell and a reference cell coated with a noncognate DNA indicates that subtracting data from a reference cell coated with noncognate DNA is the preferred control (Figure S1). An amino acid alignment of the SBP DNA-binding domains encoded by the 16 A. thaliana genes, the 2 A. majus genes, and the CuRE-binding C. reinhardtii gene is provided with a schematic of the Zn$^{2+}$ ion coordinating residues (Figure S2). The oligonucleotide primers used for site-directed mutagenesis of both the DNA target and the AtSPL14 SBP domain protein are shown in supplementary tables (Tables S1 and S2). These materials are presented following the References.

References


Supporting Information follows.
Figure S1. Comparison of surface and noncognate dsDNA-coated reference cell subtraction methods. SPR sensorgrams of different concentrations of the AtSPL14-SBP domain protein (3.125, 6.25, 12.5, 25, 50, 100, 200 and 300 nM; bottom to top) injected onto a SA sensor chip coated with a representative 156 bp cognate dsDNA fragment identified by SELEX that bound in EMSA competition assays. Binding data were collected at a flow rate of 25 μL/min. **A)** Sensorgram with signals from the control surface reference cell (no DNA) subtracted. **B)** Sensorgram with signals from the control reference cell (coated with a noncognate dsDNA) subtracted.

Figure S2. Multiple sequence alignment of several SBP DNA-binding domains and Zn$^{2+}$ ion coordination. **A)** SBP domain sequences were aligned using the Clustal method with DNASTar software (Madison, WI). Residues that match the consensus sequence are shaded in black, numbers on the left refer to the starting amino acid, protein designations are indicated on the right (organisms: At, Arabidopsis thaliana; Cr, Chlamydomonas reinhardtii; and Am, Antirrhinum majus), and asterisks and amino acid numbers indicate the positions of the six highly conserved cysteines that were mutated in the AtSPL14 SBP domain. **B)** The coordinating Cys and His residues for Zn$^{2+}$ ions are indicated for the Zn1 (CCCH) and Zn2 (CCHC) structures.
### Table S1. Oligonucleotide primers used for site-directed mutagenesis of SELEX binders

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Sense oligonucleotide primer (5’→3’)</th>
<th>Antisense oligonucleotide primer (5’→3’)</th>
</tr>
</thead>
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<tr>
<td>M1</td>
<td>CGGGATCAGATATTAGACTAAGTGCAGGTGCAGGCAATTCCCGG</td>
<td>GACCTGAATTCGCTACGGCTACCTGAAATATCTGTCATCCCG</td>
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<tr>
<td>M2</td>
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<td>GACCTGAATTCGCTACGGCTACCTGAAATATCTGTCATCCCG</td>
</tr>
<tr>
<td>M3</td>
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<tr>
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</tr>
<tr>
<td>M5</td>
<td>CGGGATCAGATATTAGACTAAGTGCAGGTGCAGGCAATTCCCGG</td>
<td>GACCTGAATTCGCTACGGCTACCTGAAATATCTGTCATCCCG</td>
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<tr>
<td>M6</td>
<td>CGGGATCAGATATTAGACTAAGTGCAGGTGCAGGCAATTCCCGG</td>
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<tr>
<td>M7</td>
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<td>GACCTGAATTCGCTACGGCTACCTGAAATATCTGTCATCCCG</td>
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<tr>
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<td>GACCTGAATTCGCTACGGCTACCTGAAATATCTGTCATCCCG</td>
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### Table S2. Oligonucleotide primers used for site-directed mutagenesis of AtSPL14 SBP domain

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<th>Antisense oligonucleotide primer (5’→3’)</th>
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</thead>
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<td>CAATTATCAACCTGAACTTGCTCCTTGCCG</td>
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<tr>
<td>C125A</td>
<td>CCCATGTGTTGCTGTATGCTCAGTTCAGGCTAGG</td>
<td>GCATGAGACAATCATCTGAACTTGCTCCTTGCCG</td>
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<tr>
<td>C142A</td>
<td>CATGAAGGCATTAAGTGGCTGAACCTCATAGTAAGCTA</td>
<td>GTAGCTTTATTAGACCTTGAGACAATCTTGCC</td>
</tr>
<tr>
<td>C161A</td>
<td>GGTAAACAGATGCAGAGGTTTCATGTCGTGTCGCTC</td>
<td>CCTGCATACTGTCGCTTGCACATCGTCTTACC</td>
</tr>
<tr>
<td>C164A</td>
<td>CAGATGCAGAGGTTTCACAGTGCAGTTCGCTC</td>
<td>GCAGATGAAACCTGAGCTCCTTGCACTTGTTACC</td>
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<tr>
<td>C180A</td>
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<td>CCAGGCAACATCTGCTGACTGCTCCTTGCC</td>
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<tr>
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<tr>
<td>C125S</td>
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<tr>
<td>C142S</td>
<td>CATGAAGGCATTAAGTGGCTGAACCTCATAGTAAGCTA</td>
<td>GTAGCTTTATTAGACCTTGAGACAATCTTGCC</td>
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