DNA binding mechanism revealed by high resolution crystal structure of Arabidopsis thaliana WRKY1 protein

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ABSTRACT
WRKY proteins, defined by the conserved WRKYGQK sequence, are comprised of a large superfamily of transcription factors identified specifically from the plant kingdom. This superfamily plays important roles in plant disease resistance, abiotic stress, senescence as well as in some developmental processes. In this study, the Arabidopsis WRKY1 was shown to be involved in the salicylic acid signaling pathway and partially dependent on NPR1; a C-terminal domain of WRKY1, AtWRKY1-C, was constructed for structural studies. Previous investigations showed that DNA binding of the WRKY proteins was localized at the WRKY domains and these domains may define novel zinc-binding motifs. The crystal structure of the AtWRKY1-C determined at 1.6Å resolution has revealed that this domain is composed of a globular structure with five β strands, forming an antiparallel β-sheet. A novel zinc-binding site is situated at one end of the β-sheet, between strands β4 and β5. Based on this high-resolution crystal structure and site-directed mutagenesis, we have defined and confirmed that the DNA-binding residues of AtWRKY1-C are located at β2 and β3 strands. These results provided us with structural information to understand the mechanism of transcriptional control and signal transduction events of the WRKY proteins.

INTRODUCTION
WRKY proteins comprise a large group of transcription factors identified specifically from the plant kingdom, ranging from lower plants such as mosses to higher plants (1,2). Members of the WRKY superfamily play important roles in a variety of developmental and physiological processes in plants. The most documented function for this superfamily of genes is their involvement in salicylic acid (SA) signaling and disease responses (3–8). SA is an important signal molecule in both local defenses and systemic acquired resistance (SAR). Endogenous accumulation or exogenous application of SA activates SAR and pathogenesis-related (PR) gene expression. In Arabidopsis, 49 out of 72 tested WRKY genes can be induced by pathogen infection or SA treatment (7), suggesting that a broad range of WRKY members could be involved in playing regulatory roles in disease resistance. This role of disease resistance was further confirmed by the finding of an over-presentation of the cis-element activated by the WRKY proteins, the W-box, within the promoters of a number of genes that are co-expressed in SAR (9,10). Arabidopsis NON-EXPRESSOR OF PR1 (NPR1) has been demonstrated as a key protein in SAR signaling pathway (11). SAR and expression of PR1 are prevented in Arabidopsis npr1 mutants even after SA treatment (12). Previous reports also showed that over-expression of WRKY18 and WRKY70 in Arabidopsis resulted in increased resistance to pathogens and enhanced expression of PR genes (3,13).

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The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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MATERIAL AND METHODS

Plant material and growth conditions

Wild-type *Arabidopsis* plants (ecotype Col-0) were grown in the soil at a growth room (23/20°C, 12-h-light/12-h-dark cycle). All the seeds were vernalized at 4°C for at least 2 days before placement in a growth environment. The non-functional NPR1 mutant *npr1–3* (12) and the transgenic line overexpressing NPR1 gene (33) were kindly provided by Dr Frederick M. Ausubel (Harvard Medical School, USA) and Dr Xin-Nian Dong (Duke University, USA), respectively.

SA treatment and northern blotting

*Arabidopsis* plants were sprayed with 2 mM SA diluted from stock solution (100 mM, adjusted to pH 6.5 with KOH). Leaves from different 4-week-old plants were harvested at indicated time points for total RNA preparation with RNAzol (Vigrous, Beijing) according to the manufacturer’s instruction. Ten micrograms of total RNA was separated on 1.5% agarose–formaldehyde gels and transferred to a nylon membrane to hybridize with digoxigenin (DIG) labeled DNA probes. DIG DNA probes labeling, hybridization and detection were performed according to DIG Application Manual for Filter Hybridization (Roche, Penzberg, Germany). The following primer combinations were used to amplify DNA probes by PCR: *PR1*: forward 5′-ATGATTTTACTGGCTTTCTCG-3′, reverse 5′-TTATGATGCGCTTCTCGTGCAC-3′; *AtWRKY1-C*: forward 5′-ATGGCTGAGGTGGAAAGTTCGTCG-3′, reverse 5′-GCTTTGGGCGAGCTCTGTCTTGGG-3′. Equal loading was confirmed by staining the gel with ethidium bromide (EB).

Protein preparation and crystallization

The preparation and crystallization of the *AtWRKY1-C* protein was described previously (34). Briefly, the cDNA fragment covering the C-terminal WRKY domain of *AtWRKY1* was amplified by polymerase chain reaction (PCR) from a full-length cDNA clone of *AtWRKY1* obtained from *Arabidopsis* (32) and cloned into the expression vector pET21aDEST. The protein was expressed in *E. coli* strain Rosetta and was purified to homogeneity in a two-step procedure of Ni²⁺ chelating and size exclusion chromatography. The mutant proteins were prepared similarly. The protein sample was kept in 20 mM Tris, 200 mM NaCl, pH 7.5 prior to crystallization. Crystals of the *AtWRKY1-C* were obtained at 277 K from condition containing 1.2 M succinic acid, 0.1 M Tris–HCl pH 7.0 and 1% w/v PEG MME 2000, by using the hanging-drop vapor diffusion methods.

Data collection and structure determination

Two crystal forms were obtained from the same growth condition. One belongs to the space group *P2_1*, with diffraction to 2.5 Å resolution at home X-ray sources as published before (34), whereas the other crystal form...
belongs to the space group P321, and diffracted to better than 1.6 Å at beamline 3W1A of Beijing Synchrotron Radiation Facilities (BSRF). Diffraction data were collected on a MAR165 CCD camera (MARresearch GmbH, Hamburg). The wavelength was set to 1.24 Å, at the higher energy side of the absorption peak of zinc. The data with anomalous signals were processed using the program DENZO and SCALEPACK (35). The statistics for the X-ray data collection and processing of the P321 crystal form are summarized in Table 1.

The zinc site was located using the program SHELXD (36) and the heavy atom parameters were refined using the program SOLVE (37). The initial phases were obtained by the program OASIS-2004 (38), then refined by using the program DM (CCP4) (39). Finally, ARP/warp (40) was used for the model auto-building.

Since the initial phases and auto-traced model were of good qualities, the subsequent refinement work could be easily carried out by using the program CNS (41). Manual adjustment of the structure was done with the program O (42), and a succinic acid as well as water molecules were added into the positive difference densities. The final model was refined by REFMAC5 (43).

**Mutant design and site-directed mutagenesis**

The design of mutants was based on a DNA–protein complex model between the W-box with the sequence 5’-ATCGTTGACCGAGTTGA-3’ and AtWRKY1-C, which was generated from the known GCM–DNA complex (44) with the PDB ID 1ODH. GCM in the complex was replaced by AtWRKY1-C with least square (LSQ) fitting between AtWRKY1-C and GCM using the program O (42). The root mean-square standard deviations (RMSD) after LSQ was 2.3 Å for 68 C atoms. A B-DNA model for specific binding was prepared using the program CNS (41) and NAB (45) with the sequence 5’-ATCGTTGACCGAGTTGA-3’. DNA in the GCM–DNA complex was then replaced using the function of LSQ fit in XtalView (46).

The wild-type AtWRKY1-C cDNA was used as a template for the mutagenesis primers. Site-directed mutagenesis by PCR was introduced into AtWRKY1-C. DNA sequencing verified the introduction of the desired mutations and demonstrated that no unwanted mutations were present in the mutated protein sequences.

**Electrophoretic mobility shift assays (EMSA)**

The oligonucleotides 5’-ATCGTTGACCGAGTTGA-3’ and 5’-TCAACTCCGTCAACCGAT-3’ containing the optimal binding site for AtWRKY1 were annealed to form DNA duplex. The typical binding reaction (20 µl) contained 3 µg dsDNA, 20 mM HEPES/KOH (pH 7.2), 40 mM KCl, 1 mM EDTA, 0.5 mM DTT, 10% glycerol, and 5 µg purified wild-type AtWRKY1-C protein or mutant proteins. The binding reaction mixture was incubated at room temperature for 20 min and the complex was separated from the free duplex on a 12% non-denaturing polyacrylamide gel electrophoresis (PAGE) in 0.5× TBE at 70 V for 3.5 h. The gel was stained with EB. Images were captured using the fluorescence imaging system (Alpha Innotech Corporation, USA). A more ‘standard’ EMSA analysis was also performed. The standard binding reaction (20 µl) contained 0.5 µg of poly(dI-dC), 20 mM HEPES/KOH (pH 7.2), 40 mM KCl, 1 mM EDTA, 0.5 mM DTT, 10% glycerol, 0–180 ng purified proteins were added and 10 ng labeled double-stranded synthetic oligonucleotide, labeled with [γ-32P]ATP using T4 polynucleotide kinase (NEB). For the competition assay, 100-fold molar excess of the unlabeled specific probe was added. DNA–protein complexes were allowed to form at room temperature for 30 min and then resolved on a 12% non-denaturing polyacrylamide gel in 0.5× TBE. The gel was dried and exposed for 2 h on X-ray films.

**CD spectroscopic assays**

All the CD spectra of the wild-type and mutant AtWRKY1-C proteins were recorded on a Jobin Yvon CD 6 spectrometer (Longjumeau, France) at 298 K. The CD spectra of all proteins were recorded in phosphate buffer saline (PBS) buffer, pH 7.5. For near-UV–CD spectrum, a cell with a path length of 1 mm was used. Each spectrum was the average of four scans corrected by subtracting a spectrum of the buffer solution in the same condition. Each scan in the range of 195–260 nm was obtained by exposure for 2 h on X-ray films.

**RESULTS AND DISCUSSION**

**AtWRKY1 is involved in the SA signaling**

SA is an important signal molecule in plant disease resistance. It has been previously shown that SA treatment can induce the expression of PR genes and activate systemic acquired resistance (SAR), which makes plants resistant against a spectrum of pathogens (47). To study the potential involvement of AtWRKY1 in SA signaling pathway and its possible relation to NPR1, a key protein in SAR, we characterized the expression profile of AtWRKY1 by RNA gel blot in wild-type Arabidopsis seedlings, npr1-3 mutant (12) and NPR1-H line (33) in
response to exogenous SA treatment. As an important marker of SAR, *PR1* was also examined to monitor the SA treatment procedure. In consistence with previous reports, *PR1* began to accumulate at 4 h post-treatment and the induction of *PR1* was abolished in *npr1-3* while increased in NPR1-H than that in wild type (Figure 1). The induced expression of *AtWRKY1* was observed after 8 h post-treatment and became obvious later than that of *PR1*. In *npr1-3*, *AtWRKY1* can still be expressed but at a lower level than that in wild-type plants. The opposite trend appeared in NPR1-H with higher expression level of *AtWRKY1* than in wild type (Figure 1). These data suggested that *AtWRKY1* was involved in SA signaling pathway and its induced expression was partially dependent on NPR1.

**The nuclear localization of *AtWRKY1-*C**

As transcriptional regulators, WRKY proteins are shown to contain functional nuclear localization signal (NLS) and are targeted to nucleus (27,48). However, the NLS sequences of WRKY proteins are not well conserved, and may be distributed at different locations. The analysis of protein sequences of *AtWRKY1* revealed a potential NLS motif KRRKK between residues 273 and 277 near the C-terminal WRKY domain, which is a conserved feature for at least nine members of the group I WRKY proteins in *Arabidopsis* (Figure 2D). We have thus included this potential NLS motif in the *AtWRKY1-C* sequence for this study. Introducing the fusion construct of *AtWRKY1-C* with dimeric GFP (dGFP) into onion epidermal cells, the green fluorescent signal targeted specifically to the nucleus. However, the fluorescence of dGFP alone and fusion protein of *AtWRKY1-C*NLS with dGFP were unevenly distributed throughout the cells (data not shown), which suggested that the potential NLS motif of *AtWRKY1-C* is indeed responsible for the nuclear localization of *AtWRKY1*.

**Structure determination of *AtWRKY1-C* crystal**

Although in the same mother liquor and temperature, *AtWRKY1-C* was crystallized in a different space group, P321, from the one published before (34). The structure was determined to 1.9 Å resolution by using the Single-wavelength Anomalous Dispersion (SAD) method. The average ratio of Bijovet pairs (<| Delta F|>/<F>) was 4.6%, in accordance with the high correlation coefficients (CC/CC all and CC/CC weak) 35.94/20.70, given by SHELXD (36) in zinc substructure locating. Phasing and density modification (DM) (39) were carried out consecutively using the software of OASIS-2004 (38) and DM (CCP4). With a figure of merit (FOM) value of 0.771 and good quality of density maps, 75 residues could be docked automatically by ARP/wARP (40). However, no density could be observed between residues 266 and 292 and between residues 369 and 371. The final model was refined manually to R-factor/R-free of 17.9 and 20.4%, respectively at 1.6 Å resolution (Table 1). The final model includes 76 residues corresponding to 293–368, one zinc ion, one succinic acid and 100 water molecules.

**Description of the overall structure of *AtWRKY1-C***

The *AtWRKY1-C* structure is mainly composed of a five-stranded antiparallel β-sheet (β1, 294–300; β2, 312–318; β3, 327–332; β4, 340–345; β5, 352–358, see Figure 2D), with the disordered N-terminal 27 (including the potential NLS motif) and C-terminal three residues missing from the structure (Figure 2). WRKY protein’s defining sequence, ‘WRKYGQK’, spans the entire β2 strand. Due to the long bridging loop (Thr301–Arg311) between β1 and β2 which covers one side of the β-sheet, the

### Table 1

<table>
<thead>
<tr>
<th>Model</th>
<th>Resolution (Å)</th>
<th>R-factor (%)</th>
<th>R-free (%)</th>
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<tr>
<td>Final</td>
<td>1.6</td>
<td>17.9</td>
<td>20.4</td>
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Figure 1. *AtWRKY1* is partially NPR1-dependent in SA pathway. Four weeks old wild type (Col-0), *npr1-3* and 35S::NPR1 were sprayed with 2 mM SA and harvested at indicated time points. *PR1* transcripts were detected the same as *AtWRKY1* except for the difference of probe. The ethidium bromide stain of rRNA is shown for each lane to allow assessment of equal loading.

Figure 2. Structure of *AtWRKY1-C* and multi-sequence alignment. (A) Ribbon representation of *AtWRKY1-C* domain. The *AtWRKY1-C* is composed of five β-strands (yellow ribbons), which are numbered from the N-terminus. The zinc ion is shown as a purple sphere and the zinc-coordinating residues are represented by sticks (yellow for C, red for O, blue for N, orange for S). (B) 3D-superimposition of the structure of *AtWRKY1-C* and the best representative NMR structure of *AtWRKY4-C* (model 15), using LSQ Fit in O. Macromolecular structures are shown by cartoons, with the *AtWRKY1-C* colored in yellow and *AtWRKY4-C* in cyan. The zinc ions are represented as spheres shown by magenta in *AtWRKY1-C* and orange in *AtWRKY4-C*. (C) Charge distribution on *AtWRKY1-C* structure surfaces given by GRASP. Positive charges are represented by blue, negative charges are represented by red. (D) Structure-based sequence alignment of both N-terminus and C-terminus of the nine *Arabidopsis* WRKY proteins from group I. The zinc-coordinating residues are shown on blue background. Conserved residue elements for stabilizing the structure and recognizing DNA are drawn on red and yellow background, respectively. Residues of β1, highlighted in green but missed in *AtWRKY4-C* structure, are rather conserved in all C-terminal domains of group I WRKY proteins. Residues marked by stars and triangles are scaffolds of two stable regions. Details of the interactions of the two regions are represented in Figure 3C and D.
structure of *AtWRKY1-C* looks rather globular and stable (Figure 2A).

The crystal structure of *AtWRKY1-C* differs from the NMR structure of a similar protein *AtWRKY4-C*-terminal domain (*AtWRKY4-C*, which shares about 53% sequence identity with *AtWRKY1-C*) determined recently (31). As shown in Figure 2B and D, the $\beta$1 strand of *AtWRKY1-C* was missing in the NMR *AtWRKY4-C* model, thus the strands $\beta$2 to $\beta$5 in the crystal structure of *AtWRKY1-C* corresponding to the strands $\beta$1–$\beta$4 of the NMR structure, and the RMSD corresponding to the similar regions of the two structures (*AtWRKY1-C* residues 309–367, *AtWRKY4-C* residues 411–469) is 1.5 Å, indicating a very similar structure for a common DNA-binding mechanism. Since the *AtWRKY4-C* domain was constructed to start from VQTTS which is located in the middle of the $\beta$1 strand (Figure 2B and D with the alignment), the discrepancy in the organization of the $\beta$-sheet is most likely caused by the different ways of preparing the NMR and crystallization samples. In order to have a fair comparison, we have repeated and constructed the NMR samples in this work and compared the CD spectra for the two proteins, *AtWRKY4-C* (Asp365–Ala469) similarly constructed as *AtWRKY1-C*, and *AtWRKY4-C$\Delta$1b* (Val399–Ala469) similar to the NMR sample without the N-terminal $\beta$ strand. The CD spectra showed similar pattern for zinc binding for both *AtWRKY4-C* and *AtWRKY4-C$\Delta$1b*, and characteristics in both cases of a folded $\beta$-sheet structure. However, there was a reduction in the CD signal of *AtWRKY4-C$\Delta$1b* near the negative peak at 210–220 nm, indicating a reduction of $\beta$-sheet signal of the truncated protein (data not shown). We thus predict that the *AtWRKY4-C* would have a similar globular structure like *AtWRKY1-C*, had the *AtWRKY4-C* sample prepared several residues longer at the N-terminus.

The conserved $\beta$1 and B-factor distribution of *AtWRKY1-C*

As shown in Figure 2D, in addition to the conserved WRKY defining residues (in red, yellow and blue), the sequences in the $\beta$1 region (colored green) are well conserved in the C-terminal domains of the group I WRKY family members. We can thus conclude that all the C-terminal domains of the group I WRKY protein possess a very similar five-stranded $\beta$-sheet architecture as in *AtWRKY1-C*.

Figure 3 describes the B-factor distribution over the *AtWRKY1-C* structure. Figure 3A shows the B-factor plot against the residue numbers, and Figure 3B is the overall structure overlaid with the B-factors color coded as defined in Figure 3A. The regions with high B-factors (higher than 25) are in both the N- and C-terminal ends and on the loop between $\beta$2 and $\beta$3, this loop may be involved in conformational changes upon DNA binding. The middle part of the long loop connecting $\beta$1 and $\beta$2, and a few residues after $\beta$5 also showed slightly high B-factor between 20 and 25, whereas most of the structure are quite stable and ordered with an average B-factor of 21, particularly the zinc-binding site and the central $\beta$ sheet are all with B-factors below 20.

The conserved salt-bridges and the D308-W312-K341 triad

Interestingly, both ends of the connecting loop between $\beta$1 and $\beta$2 show conserved pattern with ordered salt-bridge interactions and extensive hydrogen-bonding networks (Figure 3C and D). Figure 3C shows that in the C-terminal end of the loop, Asp308 forms a well-defined salt-bridge with Lys341, extensively H-bonding with side chains of Thr312, Tyr357 and backbone of Tyr310. The three key residues Asp308, Thr312 and Lys341, components of D308-W312-K341 triad, and their hydrogen bonded residues Gly309, Tyr357 are marked by stars in Figure 2D. The stable N-terminal end of the loop between $\beta$1 and $\beta$2, Arg345 is hydrogen bonded with the backbones of Thr301, Phe303 and Thr350. A potential salt-bridge between Arg345 and Asp304 was also observed as shown, and these two conserved residues are marked by triangles in Figure 2D.
the lowest $B$-factors, we thus name this constellation as D308-W312-K341 triad. In the NMR studies, the interactions around D308 could not be seen due to the undefined N-terminal of that structure, only the hydrophobic interaction of AtWRKY4-C W414-K443 (AtWRKY1-C residues W312-K341), which is part of the features of the previously defined triad constellation, has been mentioned (31). Figure 3D depicts another well conserved salt-bridge between Asp304 and Arg345 in the N-terminal end of the loop, and Arg345 is hydrogen bonded with the main chain of Thr301, Phe303 and Thr350.

The zinc-binding site in AtWRKY1-C

One of the important features resolved by the high-resolution crystal structure of AtWRKY1-C is the well-ordered zinc-binding site. As shown in the electron density map in Figure 4A, the high-resolution data of 1.6 Å made it possible to distinguish the zinc coordination with all the ligand atoms, and it was clearly shown that the Nδ of His361 and the Nε of His363, coordinate to the zinc ion (Figure 4A and B). The distances between the ligand atoms and the zinc ion are 2.06 Å (Zn–Nδ of His361), 2.07 Å (Zn–Nε of His363), 2.28 Å (Zn–Sγ of Cys332), 2.27 Å (Zn–Sγ of Cys337), all the distances are well within the ideal distances of a zinc-binding site in proteins (49).

The whole zinc-binding site is situated in the low $B$-factor regions (Figure 3), and the zinc ligand residues have almost the lowest $B$-factors in the structure, the zinc-binding site is right next to the low $B$-factor D308-W312-K341 triad described in the Figure 3C. Together with the D308-W312-K341 triad, the zinc-binding site is shaping up and stabilizing the AtWRKY1-C structure, since the zinc binding has been shown to be crucial for the stability of WRKY proteins (31).

The sequence signature of this zinc-binding site, C–X$_4$–C–X$_{23}$–H–X–H, somewhat resembles a classic C2H2 profile (50), the feature of a zinc-finger. In the crystal structure, it has been clearly shown that the zinc ion is coordinated by two cysteines (Cys332, Cys337) and two histidines (His361, His363) (Figure 4), but structurally dissimilar to classic zinc fingers, instead the structure is similar to a sequence unrelated DNA-binding protein Drosophila GCM (glia cell missing) (44) (PDB ID 1ODH). In contrast with the classic C2H2 zinc finger, AtWRKY1-C does not contain any helix, so it cannot bind DNA through the helix as most zinc-finger proteins do. The DNA-binding ability of AtWRKY1-C is mediated through the beta-hairpin regions between β2 and β3, and this mode of DNA–protein interaction is similar to the insightful prediction by Church, Sussman and Kim long time ago for non-helical DNA binding (51). Previous work on DNA binding of WRKY family has also pointed out that the conserved WRKYGQK region located on β2 strand is important for DNA binding (29).

Conserved residues responsible for DNA binding

There are more than 70 members in the Arabidopsis WRKY family grouped into three major groups (26), nine members from the group I with two WRKY domains are selected for multiple sequence alignment based on secondary structure information (Figure 2D) (sequence data were downloaded from http://rarge.gsc.riken.go.jp/rartf/tf_info/fasta_p1/TF_9_P1.fasta). It was shown from the alignment that at least 18 residues are strictly conserved, including the four residues coordinating with the zinc ion, and three residues in the conserved D308-W312-K341 triad.

As shown in Figure 2D, more than 10 well-conserved residues are positioned in the region between β2 and β3 including the WRKYGQK motif. Particularly, the five consecutive residues RKYGQ, distinct components of the WRKY family scaffold, belong to the β2 strand; while the other five conserved residues located on β3 strand, in the pattern of PRSYYR/K. Considering the specific DNA sequence that all WRKY members could recognize, these conserved residues are good candidates for the specific DNA-binding process.

As shown in Figure 2C, the surface charge distribution of AtWRKY1-C was uneven, with a calculated PI (point of iso-electricity) of 9.5. On one side, there were consecutive regions of positively charged residues, including Arg313, Lys314, Lys318, Lys321, Arg327 and Arg331, all belonging to the strictly conserved residues and located on β2 and β3 strands. Well-conserved Lys340 located at the beginning of strand β4 also belongs to this positively charged region. Since the WRKYGQK sequence falls nicely into this positively charged region and it has been shown that the WRKY motif is responsible to bind to the W-box (29), it can thus be deduced that the β2 and β3 strands are most likely to be involved in the process of DNA binding.

Identification and confirmation of the DNA-binding residues

After searching for similar structures of AtWRKY1-C using the DALI server (52), a DNA–protein complex
structure, the Drosophila GCM (44) (PDB ID 1ODH, Z-score > 6.2) was returned. When the GCM protein was used to perform a least-square fitting with *AtWRKY1-C*, the resulting RMSD between the aligned parts (68 residues) was 2.3 Å, a DNA complex model of W-box DNA and *AtWRKY1-C* was thus constructed as shown in Figure 7A. By comparison of the potential DNA-binding residues (which are close to the DNA major groove as shown in the complex model), and previous work on the DNA-binding related residues on the WRKYGGQK motif (29), the following mutants with a single residue change were constructed: R313E, K314A, K314R, Y315F, Y315R, G316F, Q317A, Q317K, R327A, R327E, Y330F, R331K, R331A and K340A. In Maeo *et al.’s* work (29), it has been shown that all residues in the WRKYGGQK (corresponding to residues 312–318 in *AtWRKY1-C*) motif are important for the DNA binding. Particularly for W312, K314, Y315, K318, a change to Ala for any of those residues would completely abolish the DNA binding. On the other hand, the R313, G316 and Q317 would only reduce the amount of DNA binding (29). We have thus constructed multiple mutants for R313, K314, Y315, G316 and Q317. Furthermore, the well-conserved residues R327, Y330, R331 and K340 are quite close to the DNA-binding sites, these residues were also selected for mutagenesis studies. Thirteen out of the fourteen mutants are located on strands β2 and β3. This region, particularly the connecting loop, showed flexible feature (high B-factor) in Figure 3A and B, indicating conformational changes might occur upon DNA binding. EMSA were used to test the DNA-binding abilities of these mutants with a fragment of 17-bp duplex DNA containing the W-box of TGAC as shown in Figure 5. The EMSA assay using 32P-labeled probe according to standard protocol was also performed with similar results (not shown). To examine the specificity of the binding, a competition experiment was performed by using an excess amount of unlabeled probe of the same sequence. When a 100-fold excess of unlabeled oligonucleotides were included in the reaction, binding of *AtWRKY1-C* to the labeled probe was abolished completely (data not shown). All the mutants could be readily expressed and purified as the wild-type *AtWRKY1-C*, the circular dichroism (CD) spectra of these mutant proteins were used to monitor the correct folding of these proteins, thus the DNA-binding ability changes are not due to the structural change caused by the mutations, see Figure 6.

As shown in Figure 5, K314A, Y315R, G316F, R331A, R327A and R327E have completely abolished the DNA-binding abilities of these mutants, and the CD spectra in Figure 6 have shown these mutants were well-structured proteins. Thus the inabilities of DNA binding must be caused by the point mutations on the correct structural scaffold. The DNA-binding activity of the mutations R313E, K314R, Y315F and Q317K were reduced, whereas the mutations of Q317A, R331K, K340A and Y330F did not show any decrease of the DNA-binding ability, indicating that these mutations did not affect the DNA binding of *AtWRKY1-C*.

These results are in good agreement with the previous work (29), and conclusions from the NMR structure (31).

In the crystal structure as well as in the NMR structure, a kink was formed in the middle of β2 around Gly316. The formation of this kink was considered helpful for the protruding of Gly316 into DNA major groove deeply and attending specific binding. The assumption was confirmed by the mutation G316F, which has lost almost all the DNA-binding ability. The complete loss of DNA binding of the mutants R327E, R327A and R331A are of great interests, since these two positively charged residues localized on the β3 strand that has been shown to be crucial for the DNA binding for the first time by site-directed mutagenesis. Furthermore, these two arginine residues, which are separated by four residues, is consistent with Tateno’s model of DNA binding at the major groove with a pair of β strands (53).

As a summary, all the mutation results further confirmed that strands β2 and β3 participate in DNA binding. It also suggested that residues Lys314, Gly316, Lys318 on β2, and Arg327, Arg331 on β3 are the key residues for DNA-specific recognition and substitution any one of them resulted in no specific binding. In addition, the distribution of these residues is *i, i+2, i+4* on one strand and *j, j+4* on the other strand. This distribution is in agreement with Tateno’s model (53), which proposed that the binding of DNA to proteins with two β strands should happen on the DNA major groove in a convex manner. Based on such a model, it has been found that normally six consecutive base pairs are
involved in specific recognition, and the large-sized residues (not only charged), such as lysines at i and i+4, could bridge the large gap separating the DNA.

The structure model of DNA–AtWRKY1-C complex we made in this study has been based on the GCM–DNA structure (44), which is different from the Tateno’s DNA-binding model (53) in details. The β-strands of the GCM protein adapted a previously undescribed binding mode to interact with DNA (44). Despite of the absence of sequence and topology similarity, AtWRKY1-C’s five-stranded β-sheet overlaps with the corresponding strands of GCM with unexpected high accuracy (Figure 7A), except for the β2–β3 region including the flexible loop, which will be subjected to conformational changes upon DNA binding. We thus refined our DNA–AtWRKY1-C complex according to the mutagenesis results, and the resulting model is quite similar to the GCM–DNA complex in topology. A schematic description of this refined model for AtWRKY1-C with the W-box, combining all information obtained from the structural and mutagenesis experiments presented in this work.

Figure 7. A model of the AtWRKY1-C and W-box interaction. (A) Superimposition of AtWRKY1-C and the overlapping structure of the Drosophila GCM domain using LSQ Fit in the program O. Macromolecular structures are shown by cartoon loops, with the GCM domain colored yellow, AtWRKY1-C blue and zinc ions as orange and magenta spheres. The DNA double-helix is shown with sticks wrapped inside surface rendering. (B) The final schematic model of the specific interaction of AtWRKY1-C with the W-box, combining all information obtained from the structural and mutagenesis experiments presented in this work.

**ACCESSION NUMBER**

The Protein Data Bank accession number for the AtWRKY1-C crystal structure discussed in this paper is PDB ID 2AYD.

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Conflict of interest statement. None declared

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