AvrXa27 binding influences unwinding of the double-stranded DNA in the UPT box

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Transcription-Activator Like (TAL) effectors, delivered by Xanthomonas pathogens bind specifically to UP-regulated by TAL effectors (UPT) box of the host gene promoter to arouse disease or trigger defense response. This type of protein-DNA interaction model has been applied in site-directed genome editing. However, the off-target effects of TAL have severely hindered the development of this promising technology. To better exploit the specific interaction and to deepen understand the TAL-induced host transcription rewiring, the binding between the central repeat region (CRR) of the TAL effector AvrXa27 and its UPT box variants was studied by kinetics analysis and TAL-blocked helicase unwinding assay. The results revealed that while AvrXa27 exhibited the highest affinity to the wild type UPT box, it could also bind to mutated UPT box variants, implying the possibility of non-specific interactions. Furthermore, some of these non-specific combinations restricted the helicase-elicited double-stranded DNA (dsDNA) separation to a greater extent. Our findings provide insight into the mechanism of TAL transcriptional activation and are beneficial to TAL-mediated genome modification.

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1. Introduction
Pathogen derived Transcription-Activator Like (TAL) effectors alter host plant target gene expression by imitating eukaryotic transcription factors [1–5]. TAL effectors share a highly conserved structure, containing an N-terminal Type III secretion signal, a central repeat region (CRR), C-terminal nuclear localization signals, and a transcriptional activation domain [6]. CRR consists of repeats with similar amino acid sequences. Crystal structure studies show that each repeat forms a helix-loop-helix motif, and several repeats fold into a right-handed superhelix wrapping around the target promoter element. The repeat-variable diresidue (RVD) in the loop specifies the nucleotides-amino acid contact [7,8]. A series of results from GUS reporter gene assays and electrophoretic mobility shift assays have been statistically analyzed to decipher the code for recognition between the RVDs and their target DNA nucleotides. Some RVDs are found to interact preferably with a particular nucleotide, while others can bind to multiple nucleotides but with different frequencies [9–14].

The unique nucleotide binding property of TAL effectors has been widely used in customizing proteins for DNA editing (including cleavage, modification and mutation of target gene), manipulation of gene transcription, and creation of broad-spectrum disease resistance crops [4,15–19]. However, some off-target issues have been reported when using this TAL-DNA binding model [20–24], which severely hinders the application of this promising technology.

Recent studies resorted to detecting the specificity of the TAL and UPT box interaction by measuring binding affinity. Meckler et al. [25] revealed that different RVDs contributed to TAL-DNA binding affinities in various orders. In another case, excessive DNA binding affinity was found to cause reduced TALEN specificity [26]. Moreover, a study concerning a Zinc-finger protein demonstrated that for a particular DNA site, the protein with the highest affinity did not show the highest specificity [27]. Therefore, details regarding the activation of host gene transcription by TAL effectors remain largely unknown, and more work should be conducted to elucidate
2.2. Protein expression and purification of AvrXa27CRR protein

A 2.2-kb fragment was PCR amplified from a vector carrying full-length AvrXa27 DNA sequence using PrimerSTAR (TaKaRa) and primers 5′-GGAATTCCATATGGCCGAGTTGATCTA (Ndel site underlined), and 5′-GCCCCCTGCAGTTGCGACAAAGGGATGGGACGTGCCGT (Xhol site underlined). The product was digested with Ndel and Xhol and inserted into protein expression vector pET15b (Novagen) to create pET15b-AvrXa27CRR, which encoded an N-terminal 6X-His affinity tag and the CRR of AvrXa27.

2.2. Protein expression and purification

The pET15b-AvrXa27CRR plasmid was transformed into E. coli strain BL21 (DE3) (Novagen). The overexpression of AvrXa27CRR was induced during the log phase (OD600 = 0.5) by 0.4 mM isopropl-1-thio-β-D-galactopyranoside for 14 h at 18 °C. Cells cultures were collected by centrifugation. The pellets were resuspended in lysis buffer (20 mM Tris-HCl pH7.6, 300 mM NaCl, 5 mM Imidazole) and lysed via sonication. After centrifugation, the supernatant containing AvrXa27CRR was first purified by Ni2+ affinity chromatography (Qiagen), then loaded on a S20 column (GE Healthcare) and eluted by a NaCl gradient (100–1000 mM). The target protein-containing fractions were concentrated and dialyzed against the storage buffer (20 mM Tris-HCl pH7.5, 400 mM NaCl, 10% Glycerol and 1 mM DTT). The final AvrXa27CRR had the purity of >90% (Fig. 1A) and was stored at −80 °C.

2.3. DNA substrates and DNA labeling

All oligonucleotides were purchased from Shanghai Sangon Biotech (Shanghai, China). The sequences and fluorescent labels of the oligonucleotides were listed in Supplementary Table 1. All the oligonucleotides were purified by high performance liquid chromatography. Each duplex DNA substrate with a 5′-end 15-nt poly (dT) single-strand tail, a central double-stranded UPT box and variants. The mutated bases were presented in red and bold font. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

2.4. DNA-binding reactions

The binding reactions between AvrXa27CRR and UPT box were performed in a 150 μl buffer containing 20 mM Tris-HCl pH8.0, 50 mM KCl, 2 mM MgCl2, 1 mM DTT. The concentration of fluorescently labeled DNA substrates was 5 nM, while the AvrXa27CRR concentration ranged from 0 to 100 nM. The reactions were incubated at 25 °C for 5 min, after which the steady-state fluorescence polarization was recorded by Infinite F200 (TECAN). The binding dissociation constant ($K_d$) of the protein-DNA complex was calculated according to previously published method [28].

2.5. TAL-blocked BsPif1 unwinding assay

The TAL-blocked BsPif1 unwinding assay was implemented by the stopped-flow fluorescence measurements. Each DNA substrate in this assay contained a 5′-end poly (dT) single-strand tail, a central double-stranded UPT box and a 3′-end quenched fluorescence signal. 100 nM AvrXa27CRR and 4 nM duplex DNA substrates were pre-incubated at 25 °C for 5 min. 100 nM BsPif1 helicase was added to incubate for another 5 min. The mixture was drawn into syringe one. A second syringe was filled with the same reaction buffer (25 mM Tris-HCl pH7.5, 50 mM NaCl, 2 mM MgCl2, 2 mM DTT) as syringe one and 1 mM ATP. After incubation, the liquids in two syringes were rapidly mixed to initiate the unwinding reaction. The fluorescence enhancement of fluorophore due to the BsPif1-catalyzed strand separation was continuously monitored by a Bio-Logic MOS-450/AF-CD spectrometer (Bio-Logic, France). Control experiments were performed under the same conditions in the absence of AvrXa27CRR. All the stopped–flow kinetic traces derived from an average of 8 individual traces. The kinetic traces were analyzed as described previously [29,30].

3. Results

3.1. Design of Xa27 UPT box variants

The RVDs in the TAL central repeats preferentially recognize one of the four bases [12,13]. The TAL-mediated genome targeting relies on this ‘one RVD to one base’ code. Four RVDs have been extensively used in base recognition: NI for adenine (A), NG for thymine (T), HD for cytosine (C) and NN for guanine (G) [15–18,20–24]. In AvrXa27CRR, there are four NIs (all for A), three NGs (one for T, two for A), two HDs (both for C), five NNs (three for G, two for A), two N’s (one for A, one for C), and one NS (for G) (Fig. 1B). Xa27 UPT box variants were obtained by mutating some of the bases to the less frequently recognized bases (NI A→C, NG T→A, HD C→A, HD

![Fig. 1. AvrXa27CRR purification and design of Xa27 UPT box mutations. (A) SDS-PAGE analysis of the purified AvrXa27CRR. Lane 1, Molecular mass marker (kD); lane 2, supernatant of the cell extract; lane 3, purified AvrXa27CRR (3 μg). (B) Alignment of the Xa27 UPT box and variants. The mutated bases were presented in red and bold font. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image-url)
C→T, NN G→C, N* C→T), and one base to the more frequently recognized base (NS G→A) in light of the cipher deduced previously [12,13] (Fig. 1B). Original bases A in RVD position −1 and T in RVD position 0 preceding Xa27 UPT box were retained in all the tested sequences (Fig. 1B). All the oligonucleotide pairs formed stable duplex DNA substrates except the N* C→T pair, and N* was not a commonly used RVD for targeting, so the N* C→T variant was not studied in the following assays.

3.2. AvrXa27CRR binds to all the tested DNA substrates

AvrXa27CRR protein was expressed and purified to achieve the final purity above 90% (Fig. 1A). The ability of AvrXa27CRR binding to the UPT box DNA substrates was probed by the steady-state fluorescence anisotropy assay. The $K_d$ value for each DNA substrate was determined based on fitting the data to the Michaelis-Menten equation, and could be used as an evaluation of the binding affinity between molecules. Generally, the $K_d$ values of most sequence-specific DNA-binding proteins are in the nanomolar level or lower [31]. In contrast, proteins bind to non-specific DNA sequences with much higher $K_d$ values [32]. Our results suggested that AvrXa27CRR bound to all the tested DNA substrates with $K_d$ values varying from 0.36 ± 0.04 nM to 2.01 ± 0.63 nM (Table 1). Apparently, AvrXa27CRR displayed preference for the wild type Xa27 UPT box with the highest binding affinity ($K_d$ value 0.36 ± 0.04 nM). The Xa27 UPT box variants exhibited an overall 2.0- to 5.5-fold reduction in AvrXa27CRR binding affinity. Among them, the variants NI A→C, NS G→A, HD C→A and NN G→C led to substantial reductions with relatively high $K_d$ values (1.46 ± 0.50 nM to 2.01 ± 0.63 nM), whereas the variants NG T→A and HD C→T presented moderate reductions with intermediate $K_d$ values (0.73 ± 0.08 nM and 1.14 ± 0.18 nM) (Fig. 2, Table 1).

3.3. AvrXa27CRR blocks the unwinding of Xa27 UPT box and variants

Transcription initiation begins with the binding of RNA polymerase and transcription factors to the promoter, followed by unwinding several base pairs of bound DNA region to form an open complex [33]. In order to investigate the effect of AvrXa27CRR on the unwinding of UTP boxes during the TAL-mediated transcriptional activation, a TAL-blocked helicase unwinding assay was introduced. BsPif1 DNA helicase, which unwound partial duplex DNA substrates with a 5′-overhang along the 5′-to-3′ direction, was used to conduct the unwinding assay [34]. In our experiment, each DNA substrate contained a 5′-end poly (dT) tail, a 19-bp UPT box double-stranded region and a 3′-end quenched fluorescence signal. Once the 5′-end quenched fluorophores were uncoupled from each other as a result of BsPif1 catalyzed unwinding, FRET signal could be detected. In the absence of AvrXa27CRR, the characteristic BsPif1 unwinding curves for all the DNA substrates were successfully obtained (Fig. 3). In the presence of AvrXa27CRR, reaction progress curves showed that the binding of AvrXa27CRR to all the tested DNA substrates effectively slowed down the overall unwinding progress (Fig. 3).

Table 1

<table>
<thead>
<tr>
<th>DNA name</th>
<th>$K_d$ (nM)</th>
<th>$K_d$ Ratio (Mutant/WT)</th>
</tr>
</thead>
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<tr>
<td>Xa27 UPT box</td>
<td>0.36 ± 0.04</td>
<td>1.0</td>
</tr>
<tr>
<td>HD C→A</td>
<td>1.52 ± 0.36</td>
<td>4.2</td>
</tr>
<tr>
<td>HD C→T</td>
<td>1.14 ± 0.18</td>
<td>3.1</td>
</tr>
<tr>
<td>NG T→A</td>
<td>0.73 ± 0.08</td>
<td>2.0</td>
</tr>
<tr>
<td>NI A→C</td>
<td>1.46 ± 0.50</td>
<td>4.0</td>
</tr>
<tr>
<td>NN G→C</td>
<td>2.01 ± 0.63</td>
<td>5.5</td>
</tr>
<tr>
<td>NS G→A</td>
<td>1.48 ± 0.42</td>
<td>4.1</td>
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</table>

Discussion

TAL effectors are shrewd tools for pathogens to remodel host plants for their own benefits of infection and propagation [35,36]. The excellent binding specificity of TAL effectors to the UPT box DNA brings hope and inspiration for genome engineering. However, an increasing number of reported off-target occurrences in TALEN-mediated gene editing urge scientists to find out more direct quantitative evaluations of TAL and target DNA interaction. In this study, the affinity of AvrXa27CRR and different UPT boxes was tested by a traditional kinetic analysis. The wild type Xa27 UPT box was the most optimal substrate for AvrXa27CRR, as all variants showed declined binding affinities (Fig. 2, Table 1), which conformed to the one-to-one correspondence rule for RVDs and target nucleotides [12,13]. However, the $K_d$ values for the interactions between AvrXa27CRR and UPT variants in our study were still at a relatively low level compared with those in other related studies (Fig. 2, Table 1) [25,31,32]. These results suggested that besides its favorite DNA sequence, AvrXa27CRR could also bind other dsDNA with similar sequence.

A previously well-studied bacteriophage T4 Dda helicase could displace the E. coli trp repressor from the trpEDCBA operator and unwind the duplex DNA substrate. The binding of trp repressor to the operator decreased the unwinding rate of Dda helicase [37]. Similarly, our results suggested that the binding of AvrXa27CRR to the wild type Xa27 UPT box and variants could postpone the BsPif1 unwinding reaction (Fig. 3, Table 2). Surprisingly, binding of AvrXa27CRR to the wild type Xa27 UPT box was not the strongest obstacle to impede BsPif1 unwinding. On the contrary, binding to some variants (NI A→C, NN G→C, HD C→A and HD C→T) caused more potent constraint on dsDNA separation (Fig. 3, Table 2). We postulated that the interaction between TAL effectors and mutated UPT sequences may significantly inhibit the dsDNA unwinding.
during transcription initiation, leading to transcriptional repression of downstream genes. This could be supported by transient promoter GUS assays in Nicotiana benthamiana, which demonstrated that the GUS reporter would be inactivated once the most favorite bases of RVDs HD (C), NI (A), and NN (G and A) were mutated to other bases [13]. In addition, our result suggested that the effect of AvrXa27 on the separation of variant NS G/A and the wild type Xa27 UPT box was similar (Fig. 3, Table 2), which was also consistent with the observation that mutations of RVD NS’s favorite base did not change the GUS activity [13].

Based on all these results, we propose a TAL-mediated transcriptional activation model, as stated below. TAL effectors

![Fig. 3. The blocking effect of AvrXa27CRR on the BsPif1-mediated DNA unwinding. (A) Time courses of BsPif1 unwinding in the presence and absence of AvrXa27CRR. Curves in black represented BsPif1 alone unwinding reaction progresses, and curves in red represented BsPif1 unwinding reaction progresses in the presence of AvrXa27CRR. (B) A schematic diagram of AvrXa27CRR blocked BsPif1 unwinding. The duplex DNA substrate (blue) was labeled with fluorescent donor fluorescein (green circle) and acceptor hexachlorofluorescein (red circle) at the 3'-end. The 5'-overhang was composed of a 15-nt poly (dT), and a double-strands region denoted the 19-bp Xa27 UPT box or its variants. The yellow triangle and purple oval typified BsPif1 and AvrXa27CRR, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image_url)

Table 2

<table>
<thead>
<tr>
<th>AvrXa27</th>
<th>A1</th>
<th>kobs, 1 (s⁻¹)</th>
<th>Rate (s⁻¹) (A1 × kobs, 1)</th>
<th>Blocking Fold [Rate (-)]/[Rate (+)]</th>
</tr>
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<tbody>
<tr>
<td>Xa27 UPT box</td>
<td>-</td>
<td>0.69 ± 0.03</td>
<td>2.03 ± 0.06</td>
<td>1.40 ± 0.04</td>
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<tr>
<td></td>
<td>+</td>
<td>0.68 ± 0.02</td>
<td>0.73 ± 0.02</td>
<td>0.50 ± 0.01</td>
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<tr>
<td>HD C→A</td>
<td>-</td>
<td>0.83 ± 0.03</td>
<td>2.16 ± 0.07</td>
<td>1.79 ± 0.01</td>
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<tr>
<td></td>
<td>+</td>
<td>0.18 ± 0.01</td>
<td>0.76 ± 0.07</td>
<td>1.14 ± 0.01</td>
</tr>
<tr>
<td>HD C→T</td>
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<td>2.09 ± 0.09</td>
<td>1.58 ± 0.00</td>
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<td></td>
<td>+</td>
<td>0.44 ± 0.02</td>
<td>0.88 ± 0.05</td>
<td>0.38 ± 0.00</td>
</tr>
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<td>NG T→A</td>
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<td>0.47 ± 0.03</td>
<td>2.35 ± 0.11</td>
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<td></td>
<td>+</td>
<td>0.79 ± 0.02</td>
<td>0.91 ± 0.02</td>
<td>0.72 ± 0.00</td>
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<tr>
<td>NI A→C</td>
<td>-</td>
<td>0.65 ± 0.09</td>
<td>1.82 ± 0.17</td>
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</tr>
<tr>
<td></td>
<td>+</td>
<td>0.25 ± 0.01</td>
<td>0.83 ± 0.03</td>
<td>0.20 ± 0.00</td>
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<td>NN G→C</td>
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<td>0.80 ± 0.03</td>
<td>2.34 ± 0.06</td>
<td>1.87 ± 0.02</td>
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<td></td>
<td>+</td>
<td>0.19 ± 0.02</td>
<td>0.85 ± 0.04</td>
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<td>NS G→A</td>
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<td>2.09 ± 0.12</td>
<td>1.39 ± 0.04</td>
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<tr>
<td></td>
<td>+</td>
<td>0.80 ± 0.01</td>
<td>0.79 ± 0.00</td>
<td>0.56 ± 0.00</td>
</tr>
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</table>

*A1 indicated the unwinding amplitude of the fast phase.

kobs, 1 indicated the observed kinetic rate of the fast phase.
preferentially bind to the wild type UPT box with less constraint on the unwinding of dsDNA, thus allowing for the recruitment of RNA polymerase and transcription activation of downstream genes (Fig. 4A). TAL effectors can also bind to non-specific UPT boxes (Fig. 2, Table 1), but in this case dsDNA unwinding will be significantly reduced, consequently resulting in shutdown of downstream genes (Fig. 4B).

The off-target problems of TALENs can be partly illustrated by our model. TALENs can bind to homologous off-target sites, leading to non-specific cleavage, especially when expressed at a high concentration. Besides the one-to-one cipher, therefore, homologous sites should be avoided as much as possible in TALENs design.

In conclusion, our results suggest that most atypical mutations in the Xa27 UPT box decrease the affinity of binding by AvrXa27CRR. However, once AvrXa27CRR binds to mutated Xa27UPT box, their interaction will interfere with dsDNA separation, which is a critical step in transcription initiation, particularly for open complex formation. Our findings thus advance the understanding of the mechanisms underlying TAL-mediated transcriptional activation and the occurrence of TALEN-induced off-target events.

Acknowledgements

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.jbbrc.2017.01.130.

Transparency document

Transparency data related to this article can be found online at http://dx.doi.org/10.1016/j.jbbrc.2017.01.130.

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