The Tomato Nucleotide-binding Leucine-rich Repeat (NLR) Immune Receptor I-2 Couples DNA-Binding to Nucleotide-Binding Domain Nucleotide Exchange


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Plant nucleotide-binding leucine-rich repeat (NLR) proteins enable plants to recognize and respond to pathogen attack. Previously, we demonstrated that the Rx1 NLR of potato is able to bind and bend DNA in vitro. DNA binding in situ requires its genuine activation following pathogen perception. However, it is unknown whether other NLR proteins are also able to bind DNA. Nor is it known how DNA binding relates to the ATPase activity intrinsic to NLR switch function required to immune activation. Here we investigate these issues using a recombinant protein corresponding to the N-terminal coiled-coil and nucleotide-binding domain regions of the I-2 NLR of tomato. Wild type I-2 protein bound nucleic acids with a preference of ssDNA ≈ dsDNA > ssRNA, which is distinct from Rx1. I-2 induced bending and melting of DNA. Notably, ATP enhanced DNA binding relative to ADP in the wild type protein, the null P-loop mutant K207R, and the autoactive mutant S233F. DNA binding was found to activate the intrinsic ATPase activity of I-2. Because DNA binding by I-2 was decreased in the presence of ADP when compared with ATP, a cyclic mechanism emerges; activated ATP-associated I-2 binds to DNA, which enhances ATP hydrolysis, releasing ADP-bound I-2 from the DNA. Thus DNA binding is a general property of at least a subset of NLR proteins, and NLR activation is directly linked to its activity at DNA.

Plants rely on an innate immune system to ward off pathogens (1–4). Pathogen perception and recognition specificity is typically controlled by NLR type immune receptors that are capable of perceiving non-self and modified self molecules inside the host cell. NLRs typically detect strain-specific pathogen effectors or the effect these virulence factors exert on host proteins (3, 5, 6). NLR proteins are members of the STAND (signal transduction ATPases with numerous domains) P-loop ATPases of the AAA+ (ATPases associated with diverse cellular activities) superfamily whose multidomain structure allows them to function CC simultaneously as sensor, switch, and response factor (7, 8).

Plant NLRs are named after their central nucleotide-binding (NB) and C-terminal leucine-rich repeat (LRR) domains. The N terminus of a plant NLR is extremely divergent and typically encodes either a coiled-coil (CC) or Toll interleukin receptor domain (3). In addition to these core domains, other domains can be present such as a WRKY, Squamosa promoter binding domain (3). In addition to these core domains, other domains can be present such as a WRKY, Squamosa promoter binding domain (3). In addition to these core domains, other domains can be present such as a WRKY, Squamosa promoter binding domain (3). In addition to these core domains, other domains can be present such as a WRKY, Squamosa promoter binding domain (3). In addition to these core domains, other domains can be present such as a WRKY, Squamosa promoter binding domain (3). In addition to these core domains, other domains can be present such as a WRKY, Squamosa promoter binding domain (3).

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and R proteins, is predicted to function as a nucleotide-operated molecular switch controlling the signaling activity of the protein (8, 13, 14). Biochemical analysis of tomato I-2 and Mi-1, flax M and L6, and barley MLA27 revealed that in the autoinhibited “off” state, the NB-ARC domain is ADP bound (15–17). Upon pathogen recognition, ADP is exchanged for ATP permitting the NB-ARC domain to adopt an activated and structurally distinct “on” state. Hydrolysis of bound ATP into ADP enables the off state to be re-established. There are several lines of supporting evidence for this model. For example, tomato I-2 mutants defective in ATP hydrolysis in vitro are autoactive (16). Further, an autoactive flax M mutant preferentially co-purifies with ATP (17). Interestingly, the NLR NB-ARC domain is not necessarily a strict ATPase. The NB subdomain of rice Os2g_25900, NB-ARC domains of maize P5IP (pollen-signaling protein), and Arabidopsis Rpm1 possesses a nucleotide phosphatase activity (18). The described phosphatase sequentially removes terminal phosphates from the nucleotide to form the nucleoside in a reaction that remains compatible with the switch model.

How activated plant NLRs trigger immune signaling is a crucial and largely unanswered question. Activation of animal NLRs typically induces multimerization, resulting in the formation of a cytoplasmic signaling scaffold on which partners are
activated because of their induced proximity (19). Unlike animal NLRs, no conserved protein binding partners for plant NLRs have been identified to date that could play an analogous role in downstream signaling. The identified NLR interactors are mostly involved in processes such as NLR protein maturation and folding, nucleocytoplasmic shuttling, or effector perception (9, 12, 20). Whereas NLR proteins locate at various subcellular localizations determined by the necessity to intercept effector action at distinct locations, the subcellular localization from which primary immune signaling is activated is not unambiguously resolved. Although some NLRs are confined to a specific subcellular compartment, like the plasma membrane localization of Rpm1 and the nuclear localization of RRS1-R (21, 22), other NLRs have a more dynamic distribution. For instance tobacco N, barley MLA1 and Mla10, other NLRs have a more dynamic distribution. For instance tobacco N, barley MLA1 and Mla10, Arabidopsis RPS4 and SNC1, and potato Rx1 show a nuclear-cytoplasmic distribution (22–29). The functional importance of nuclear localization is emphasized by genetic screens that reveal genes encoding components of the nuclear pore complex to be required for NLR-mediated resistance (30, 31). Furthermore, redistribution of MLA10, N, RPS4, and SNC1 from the nucleus to the cytoplasm compromises their immune signaling (24, 25, 27, 31), suggesting that their signaling target resides in the nucleus. Together with the recent notion that many NLRs work in pairs, suggesting that their signaling target resides in the nucleus.

In line with this hypothesis, we recently identified DNA as a molecular target for an activated NLR. We demonstrated that the potato Rx1 NLR possesses an intrinsic DNA binding and melting activity in vitro (37). In addition, Rx1 was observed to bend and locally melt dsDNA. DNA binding is mediated by its NB-ARC domain in a manner similar to that of the structurally homologous origin of replication binding proteins. An Rx1-DNA interaction in plants was only found upon activation of immune signaling by the Rx-immune elicitor, the CP106 coat virus. No Rx1-DNA interaction was observed in the presence of a PVX virus coat protein variant that is incapable of Rx1 activation, nor was it observed when immune signaling was activated via another NLR protein. This finding raises the exciting possibility that a direct NLR-DNA interaction might be a conserved signaling function of activated NLR proteins. To test this hypothesis, we set out to establish whether other NLR proteins are also able to interact with DNA in an activation-dependent fashion. Demonstration of a direct DNA interaction by an alternative NLR provides support that this phenomenon is of broad biological relevance. Further, these experiments will allow an appraisal of the mode of interaction and could reveal NLR-specific differences in nucleic acid interactions.

The I-2 (immunity to race 2) NLR protein from tomato, conferring resistance to tomato wilt disease caused by the fungus Fusarium oxysporum f.sp. lycopersici, is one of the best studied NLR proteins (16, 38). I-2 is a member of the CC-NB-LRR class of plant NLR proteins that consists of an N-terminal CC domain fused to an NB-LRR domain (39). The CC-NB-ARC domain has been heterologously produced in Escherichia coli and shown to bind specifically to adenosine nucleotides and to possess ATPase activity in vitro (38). Specific amino acid substitutions in its catalytic site that compromise its ATPase activity result in a mutant I-2 protein that confers an autoimmune phenotype when expressed in planta (16). The I-2 protein recognizes the F. oxysporum f.sp. lycopersici produced Avr2 protein inside the plant nucleus (40), making I-2 a prime candidate NLR protein to have a nuclear signaling activity like Rx1, MLA, RPS4, RPS5-RSS1, SNC1, and N (23–27, 29, 41, 42).

Here we demonstrate that the I-2 NLR protein of tomato is able to bind, bend, and melt duplex DNA in vitro. We find important differences between the Rx1-DNA and I-2-DNA interaction; I-2 shows a nucleic acid-binding specificity distinct from Rx1 but similar to the orphan Os02g_25900 NLR of rice. Furthermore, in contrast to Rx1, the I-2 interaction with DNA is coupled to its nucleotide-dependent activation cycle. This directly links DNA interactions with the NLR activation state. We propose that NLR-DNA interactions are a general phenomenon but with NLR-specific differences in the mode of DNA interaction.

**Experimental Procedures**

**Structural Modeling—**Protein fold searches using the Phyre2 protein homology/analogy recognition engine, version 2.0 (43) were undertaken using amino acids 175–519 of I-2 and amino acids 197–334 of Os02g_25900 using both normal and intensive modeling modes. Similar structural homology was also detected using the SAM-T08, HMM-based protein structure prediction server (44). Side chain packing and energy minimization was performed using GalaxyRefine (45). The figures were generated using the PyMOL molecular graphics system (46).

**Protein Expression and Purification—**Proteins corresponding to I-21–519 WT and I-21–519 K207R were generated as described previously (38). The NB subdomain of Os02g_25900 (amino acids 197–334; R1-NB) was generated as previously described (18). Orc1-1 and Orc1-3 of Sulfolobus solfataricus were expressed and purified as previously described (47).

**ATPase Assays—**ATPase assays were typically performed at 37 °C for 30 min with 2.3 μM protein in 50 mM 1,3-bis(tris(hydroxymethyl)methylamino) propane, pH 7.5, 10 mM MgCl2, and 5 μM ATP. Reactions were spiked with 0.5 μCi of 2.8-3H-labeled ATP for quantitation. Reactions were spotted onto a silica thin layer chromatography plate with 1 mM ADP to act as both marker and carrier. The plates were developed in isobutanol:3-methyl-1-butanol:2-ethoxyethanol:ammonia:H2O (9:6:18:9:15). Spots were visualized at 256 nm and quantified using an AR-2000 TLC scanner.

**Electrophoretic Mobility Shift Assays—**The oligonucleotides used for quantitative EMSA are derived from a series of oligonucleotides that enables a comparison of relative DNA binding affinity to varying DNA topologies independent of DNA sequence (48). The oligonucleotide sequences were 5′-TGG GTC AAT GTG GGC AAA GAT GTG GCA TTA TAA TCG TCT ATG AGT TT-3′ (SS1; DNA sense-strand), 5′-AAC GTC ATA GAC GAT AC TAT GCT AGG ACA TCT TTG
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CCC ACG TTG ACC CA-3’ (SS2; DNA antisense-strand), and 5’-UGG GUC AAC GUG GGC AAA GAU GUC CUA GCA AUG UAA UCG UCU AUG ACG UU-3’ (RNA sense-strand) (47). Oligonucleotides were end-labeled with 10 μCi of [γ-32P]ATP using T4 polynucleotide kinase, and unincorporated nucleotides were removed using Micro Bio-Spin columns (Bio-Rad). Protein and 0.15 mM nucleic acids (oligonucleotide 1-ssDNA, annealed oligonucleotide 1 and 2-dsDNA, and ssRNA) were incubated in 20 mM Tris, pH 8.0, 1 mM EDTA before cooling to room temperature. Nucleotides were annealed by heating to 90 °C for 3 min in 10 mM Tris, pH 8.0, 1 mM EDTA before cooling to room temperature. Steady-state FRET was measured using a PerkinElmer LS 55 fluorescence spectrometer at room temperature in a final volume of 50 μl with 50 nM DNA. Measurements used 1.5 μM protein and were incubated for 10 min at room temperature before scanning. A bandwidth of 5 nm was used for both excitation and emission wavelengths. Donor (fluorescein) was excited at 494 nm, and emission spectra were collected from 490 to 650 nm. Acceptor (TAMRA) was directly excited at 558 nm, and emission spectra were collected from 558 to 650 nm. Spectra were collected for both donor and donor-acceptor-labeled double-stranded DNA, and FRET values were calculated from the increase in acceptor emission using the ratio method (50). Ratio was used to calculate energy transfer efficiency (E) using the equation $E = \left( \frac{e_a[558]}{e_a[494]} \right) \times \frac{\Delta \omega - \omega_a}{\omega_a}$, where $e_a[558]$ and $e_a[494]$ are the acceptor and donor extinction coefficients provided by the supplier. Donor-acceptor distances (R) were calculated using the equation $E = \frac{R_0^6}{(R_0^6 + R^6)}$ and a calculated Förster distance ($R_0$) of 49.99 Å. The induced bend angle ($\theta_i$) was calculated using a single-point bend model.

**Time-resolved FRET in Vitro—**Oligonucleotides were as for steady-state FRET in vitro and annealed in the same manner. Strands were annealed by heating to 90 °C for 3 min in 10 mM Tris, pH 8.0, 1 mM EDTA before cooling to room temperature. Measurements used 1.5 μM protein with 50 nM DNA in the presence of 60 mM NaCl and were incubated for 10 min at room temperature before analysis. Time-resolved FRET was assessed using the time-correlated, single photon counting technique. The excitation source was a Pic quant pulsed diode laser LDH-P-C485 (excitation wavelength, 485 nm; 70-ps pulse FWHM at 20 MHz). Fluorescence was detected using an avalanche photodiode (Id Quantique 100-50) linked to a Becker and Hickl SPC 130 time-correlated, single photon counting module. An instrument response function of ~200 ps was measured from Rayleigh scattered light. Fluorescence decays were collected for both donor and donor-acceptor-labeled double-stranded DNA with or without protein using band pass filter detection of the donor emission and at magic angle polarization.

Data were analyzed by the Grinvald-Steinberg method (51) to obtain the fluorescence lifetime for the donor and acceptor-labeled (τDA) and donor only-labeled (τD) oligonucleotides. The data were fitted to a sum of exponentials using an iterative least squares reconvolution procedure with the optical/electrical excitation profile to produce a biexponential decay containing two lifetimes. This profile was obtained from a slide covered with Silica ludox particles, which provides an instant scatter of the excitation pulse. This data-fitting method provided more
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accuracy in the determination of shorter lifetimes than calculating a single average lifetime. Efficiencies of energy transfer were calculated from time-resolved FRET according to $E = 1 - (t_{\text{DA}}/t_{\text{D}})$. Energy transfer efficiencies in the lifetime analysis were higher than those from steady-state experiments as previously observed (52–54). Donor-acceptor distances ($R$) were calculated using the equation $E = R^0_s/(R^0_s + R^0)$ and a calculated Förster distance ($R_0$) of 49.99 Å. The total length of the oligonucleotide with linkers and fluorescent dyes, at maximum extension, was calculated as 81.1 Å. The induced bend angle ($\theta_i$) was calculated using a single-point bend model.

Statistical Analysis—Error bars represent the standard error of the mean with the number of replicates as indicated in the legend. Statistical comparisons (p values) for data that passes a test for normality (D’Agostino and Pearson omnibus normality test and Shapiro-Wilk normality test) were obtained from one-way analysis of variance with the indicated post hoc test. Statistical comparisons (p values) for data that do not pass a test for normality were obtained from a Kruskal-Wallis test with post hoc multiple comparisons test. p values in statistical comparisons are indicated in the figures with letters and indicate compared data sets as described in the figure legends.

Results

I-2 Binds Nucleic Acids in Vitro—The NB-ARC domain of Rx1 (amino acids 143–488) has significant homology with the Cdc6/Orc1 proteins of *Pyrobaculum aerophilum* (Protein Data Bank code 1FNN) and of *Aeropyrum pernix* (Protein Data Bank code 2V1U) in complex with DNA (37). We therefore investigated whether there is a similar structural homology in I-2. Amino acids 175–519 of I-2, encompassing the NB-ARC domain, were analyzed using the Phyre² protein fold recognition engine, and the expected close matches with the pro-apoptotic proteins CED-4 (Protein Data Bank code 2A5Y) and ApaF-1 (Protein Data Bank code 1ZET) were recovered to 100% confidence (55, 56). Like Rx1, high scoring matches (>99% confidence) were also obtained with the Cdc6/Orc1 protein family members. Residues in the N-terminal subdomain and tandem ARC domains were conserved between Cdc6/Orc1 of *A. pernix* and I-2 (29.7% similarity and 9.9% identity). Both the N-terminal NB (amino acids 22–193) and C-terminal ARC domains (amino acids 194–388) in Cdc6/Orc1 contact DNA, inducing deformation of the double helix (47, 49), and the modeled tertiary structure of I-2 closely matched that of Cdc6/Orc1 (Fig. 1A). Structural modeling therefore suggests that the NB-ARC domain of I-2 could contact DNA in a manner similar to that of Cdc6/Orc1 and as hypothesized for Rx1. We therefore investigated whether I-2 is also a DNA-binding protein.

A possible direct I-2-DNA interaction was investigated through in vitro experiments. EMSA using nucleic acid fragments of >5 kb derived from circular bacteriophage dX174 (57) represents a standard methodology to qualitatively assess interactions between a protein and either ssDNA or dsDNA with identical sequences. EMSAs were performed using recombinant wild type I-2 protein (I-2 WT) encompassing the CC-NB-ARC region but lacking the LRR (Fig. 1B). EMSA experiments performed with I-2 WT showed a direct association with both ssDNA and dsDNA, producing an upward shift in the migration of the nucleic acids similar to that of Rx1 and other unrelated DNA-binding proteins (Fig. 1C) (18, 58). I-2 is therefore able to interact with both ssDNA and dsDNA in a sequence-independent manner.

The I-2-DNA interaction was relatively stable because it could be visualized after gel electrophoresis (Fig. 1C). Nevertheless, whereas EMSA using circular bacteriophage dX174 DNA is a useful method to qualitatively assess the interaction, it does not enable robust quantification of the affinity of I-2 for nucleic acids. We therefore shifted to EMSAs using small synthetic oligonucleotides to quantify I-2-nucleic acid interactions (59). EMSA with oligonucleotides provides more robust band shifts on EMSA because of their lower molecular mass.

The affinity of I-2 WT was assessed using 32P-labeled synthetic oligonucleotides whose sequences were unrelated to that of bacteriophage dX174 DNA. I-2 WT showed equivalent affinities for ssDNA and dsDNA but a reduced affinity for ssRNA (Fig. 2A and Table 1). The ordering of affinities...
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of I-2 WT for nucleic acid (ssDNA ≈ dsDNA > ssRNA) is distinct from those of Rx1 (ssDNA > ssRNA > dsDNA), implying that different NLRs can have distinct nucleic acid binding properties in vitro. One caveat to this observation is that I-2 is a refolded recombinant protein. Hence, its altered nucleic acid specificity might be a reflection of its refolded status. We therefore investigated the nucleic acid binding specificity of a natively folded NB-ARC subdomain.

We were unable to produce the individual NB or NB-ARC subdomains of I-2 in E. coli and therefore shifted to the isolated NB subdomain of an I-2 homolog, the orphan NB-LRR protein Os02g_25900 (R1-NB). R1-NB was selected for analysis because it has been previously demonstrated to be readily expressed as a natively folded protein in E. coli (18). R1-NB therefore avoids the criticism that its activity might be affected by the in vitro refolding procedure that could result in a subfraction of potentially mis- or unfolded protein. Furthermore, structural modeling demonstrated a close structural similarity between the NB domains of R1-NB and that of I-2 (Fig. 2B). Like I-2 WT, R1-NB bound both circular bacteriophage φX174 ssDNA and dsDNA in a sequence-independent fashion (Fig. 2C). R1-NB interacted with a variety of oligonucleotides with an ordering of affinities for nucleic acids of ssDNA ≈ dsDNA > ssRNA (Fig. 2D and Table 1). The ranking for R1-NB is similar to that observed for I-2 WT with a clear preference for binding DNA over RNA. We hence conclude that the nucleic acid affinities for I-2 are likely a reflection of its genuine biochemistry and not an artifact of refolding the recombinant protein.

The P-loop mutation K207R of I-2 (I-2 WT) reduces the Kd for ATP in vitro and confers a loss of function phenotype in vivo (38). We therefore set out to test whether the I-2 K207R mutant shows a similar distinct pattern of interaction when compared with the wild type protein. We first compared the affinity of I-2 WT and I-2 K207R for dsDNA. Using an EMSA assay, the affinities of I-2 WT and I-2 K207R for dsDNA were largely indistinguishable (Fig. 3A). However, when the affinity of I-2 WT and I-2 K207R was compared using fluorescence anisotropy, the affinity of I-2 WT for dsDNA was significantly greater than that of I-2 K207R (Fig. 3B and Table 1). Affinities measured by anisotropy depend not only upon the mass of a protein-DNA complex at a given molar ratio but also on its globular structure. EMSA therefore suggests that the binding constants of I-2 WT and I-2 K207R are very similar, but anisotropy reveals that the overall shape of the protein complex on DNA might be different. Together, these data indicate that, like Rx1, I-2 and R1 interact with nucleic acids but that the affinities for different nucleic acids vary for the different NLRs. Further, the structure of the NLR-DNA complex formed in I-2 depends on the presence of an intact P-loop.

I-2 Deforms DNA—The Cdc6/Orc1 family proteins substantially deform origin DNA by bending it with angles of 35 and 20°, respectively, thereby inducing localized melting of the double helix (47, 49, 60). The Rx1 protein is also able to bend

FIGURE 2. The I-2 CC-NB-ARC domain and the R1 NB domain bind nucleic acids in vitro. A, quantitative EMSA analysis giving affinities of I-2 WT for synthetic oligonucleotides corresponding to different nucleic acids (n = 3–6, ±S.E.). B, overlay of a structural homology model for amino acids 197–334 encompassing the NB domain of R1-NB (blue) onto the I-2 NB-ARC domain model of Fig. 1A (yellow), C, EMSA for R1-NB using 100 ng of φX174 DNA (ssDNA) or φX174 RF I DNA (dsDNA). For dsDNA, the upper band represents relaxed circular DNA, whereas the lower band represents supercoiled circular DNA. Note that both bands shift upon incubation with the R1 NB domain. Molecular weight markers are indicated with arrows. D, quantitative EMSA analysis giving affinities of R1 NB for various synthetic oligonucleotides corresponding to different nucleic acids (n = 3–10, ±S.E.).

TABLE 1

Apparent dissociation constants for recombinant NLR domain interactions with nucleic acids

<table>
<thead>
<tr>
<th>Protein Method</th>
<th>ssDNA</th>
<th>dsDNA</th>
<th>ssRNA</th>
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<tbody>
<tr>
<td>1-2, 519 WT EMSA</td>
<td>0.15 ± 0.01</td>
<td>0.22 ± 0.02</td>
<td>&gt;100 μM</td>
</tr>
<tr>
<td>1-2, 519 K207R EMSA</td>
<td>ND</td>
<td>0.23 ± 0.01</td>
<td>ND</td>
</tr>
<tr>
<td>1-2, 519 WT Anisotropy</td>
<td>ND</td>
<td>0.32 ± 0.05</td>
<td>ND</td>
</tr>
<tr>
<td>1-2, 519 K207R Anisotropy</td>
<td>ND</td>
<td>0.94 ± 0.09</td>
<td>ND</td>
</tr>
<tr>
<td>R1-NB EMSA</td>
<td>3.34 ± 0.54</td>
<td>2.75 ± 0.40</td>
<td>12.45 ± 2.34</td>
</tr>
</tbody>
</table>
dsDNA at an angle of 42° (37). We therefore investigated whether I-2 is able to deform dsDNA in a similar fashion. We examined DNA distortion by steady-state FRET using two-sided end-labeled dsDNA (61). DNA distortion was evident by an increase in FRET acceptor emission in the presence of I-21–519WT or I-21–519K207R when compared with the mock (no protein) or to a control protein (BSA) (Fig. 4A). Energy transfer efficiency correlates with the distance between fluorophores and was found to increase in the presence of I-21–519WT or I-21–519K207R. The change in efficiency can be used to calculate their distances and revealed that I-21–519WT and I-21–519K207R induced bend angles (θ) of 21.8–23.9 and 22.1–23.1°, respectively (values represent a range of ± S.E. from the mean). We next measured the fluorescence lifetime, which represents an intrinsic property of the fluorophore that is independent of concentration, photobleaching, and light scattering, as an alternative method to measure DNA distortion through a FRET process. The decrease in fluorescent donor lifetime can only be rationalized because of increased energy transfer from donor to the acceptor. Together with the data from steady-state analysis, these data both show increased energy transfer efficiency from donor to acceptor in the presence of I-21–519WT or I-21–519K207R (Fig. 4B). The recorded fluorescence lifetimes correspond to θ values for I-21–519WT and I-21–519K207R of 29.7–36.7 and 29.6–35.1°, respectively (values represent a range of ± S.E. from the mean). In conclusion, I-21–519 distorts DNA like Cdc6/Orc1 and R1. However, the bend angle induced by I-2, as measured by two independent methods, is substantially lower than that induced by Rx1, indicating a similar yet distinct mode of action.

The Orc1 protein of A. pernix and Rx1 also induces local DNA distortion upon binding. The P1 nuclease can be used to detect distortion specific ssDNA in the presence of the bending proteins (37, 60, 62). We therefore examined the sensitivity of dsDNA oligonucleotides to the ssDNA-specific P1 nuclease following incubation with BSA (negative control), I-21–519, or ORC (positive control). *, p < 0.01 compared with control by one-way analysis of variance with post hoc Dunnett test.
BSA (negative control) was largely resistant to P1 nuclease activity (Fig. 4C). dsDNA was more sensitive to P1 nuclease in the presence of either I-2WT or the mutant I-2K207R, showing that both proteins can melt dsDNA. The P1 sensitivity of dsDNA in the presence of Orc1-1/Orc1-3 was indistinguishable from that of dsDNA in the presence of I-2WT, supporting the interpretation that I-2, similar to Rx1, can cause local dsDNA melting (Fig. 4C). In conclusion, I-2 is able to bend DNA and provoke localized DNA melting. Because the pattern of DNA distortion induced by I-2 is indistinguishable between I-2WT and the mutant I-2K207R, the difference in the nature of the protein-DNA complex is not at the level of melting the DNA.

The I-2-DNA Interaction Is Coupled to the Nucleotide Binding State and Activity of the P-loop—The findings on DNA interaction and distortion seem to indicate that the difference between I-2WT and the mutant I-2K207R is related to the conformational state of the protein (Fig. 3B) and not DNA melting (Fig. 4C). We therefore sought independent evidence to understand the relationship between the nucleotide occupancy and ATPase activity of the P-loop in I-2 (compromised in I-2WT) and DNA binding. In the “switch” model for plant NLR activation, binding of ATP to the NB-ARC domain establishes the on state, whereas hydrolysis of ATP to ADP restores the off state (13). An intact P-loop is essential for nucleotide binding, and mutations in this motif typically result in loss of function alleles (13). We therefore investigated the relationship between P-loop-dependent ATPase activity and DNA binding. The CC-NB-ARC domain of I-2 has a very low intrinsic ATPase activity in vitro. Interestingly, the ATPase activity of I-2WT was increased 2-fold in the presence of DNA (Fig. 5A). Although I-2K207R has a reduced ATPase activity, reflecting its lowered K_m for ATP, its enzymatic activity was also stimulated 2-fold by DNA.

Next, we tested the reciprocal relationship and examined whether nucleotides influence DNA binding by I-2WT. To be able to monitor either increased or reduced binding capacity, an I-2WT protein concentration was used that gives ~50% of maximal DNA binding. The ratio of I-2WT bound to DNA in the presence of 5 μM ATP (giving about 70% nucleotide occupancy for I-2WT (16)) compared with DNA-bound I-2WT in the absence of nucleotides was approximately unity (Fig. 5B). In contrast, the ratio of I-2WT bound to DNA in the presence of 5 μM ADP was ~0.65 as compared with that in the presence of ATP or in the absence of nucleotides, demonstrating a reduced binding of the I-2-ADP complex. Binding of I-2WT to DNA in the presence of the nonhydrolyzable ATP analog, adenosine 5′-(β,γ-imido)triphosphate, was indistinguishable from DNA binding in the presence of ATP, ruling out that the enhanced binding, compared with that measured in the presence of ADP is due to hydrolysis of ATP. The influence of nucleotides on I-2WT DNA binding was the same as that for I-2WT. Because K207R is a K_m mutation that likely affects coordination of the ATP β-PO_4_2^−, it is not surprising that this mutant shows a similar response to the nucleotides. Because the ATP and “no nucleotide” bound states appear equivalent in this assay, it was also to be expected that this mutant behaved similarly to the wild type, despite its lowered K_m for ATP.

To provide further evidence that the difference between the DNA binding ability of the ATP and ADP-bound states is not
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due to changes in nucleotide handling, we investigated an additional I-2 mutant. The S233F mutant is autoactive in vivo because of its reduced ATP hydrolysis rate, whereas it has an affinity for ATP equivalent to wild type (16). Binding of I-2[1–519]S233F to dsDNA was indistinguishable from I-2[1–519]WT (Fig. 5C). We found that for I-2[1–519]S233F, similar to wild type I-2, DNA binding in the presence of 5 μM ADP was only 61 ± 7% (S.D.) of that in the presence of 5 μM ATP. This observation further supports our hypothesis that DNA binding by I-2 is enhanced by ATP relative to ADP and that this change cannot be attributed to differences in nucleotide handling by the different proteins.

Together these results indicate that the intrinsic ATPase activity of I-2 is enhanced in the presence of DNA. This converts the ATP-bound state into the ADP-bound conformation with a consequent increased likelihood for release from DNA.

Discussion

Immune responses induced by NLR immune receptors are broadly conserved (63), suggesting a common signaling network that might be activated via a conserved mechanism. Existing in vitro, in vivo, and bioinformatics data identify the NB-ARC domain as the most conserved domain in NLR proteins. The NB-ARC domain functions as a molecular switch regulating NLR activity. Its conservation makes the NB-ARC domain also a prime candidate to be involved in activating the downstream signaling process. The NB-ARC domain of the potato Rx1 NLR protein was shown to possess intrinsic DNA binding activity that is activated in situ only upon triggering the cellular immune response by admission of the pathogen-derived elicitor (37). Here we demonstrate that both R1 of rice and I-2 of tomato are also able to interact with nucleic acids in vitro. This study therefore establishes nucleic acid interactions as central to the biochemistry for at least a subset of NLR proteins. The observed specificity for ssDNA, dsDNA, and RNA for these proteins revealed subtle differences in their abilities to interact with nucleic acids. The functional relevance of these differences awaits future elucidation.

I-2 shares many biochemical properties with Rx1 and the Cdc6/Orc1 family of DNA-binding proteins. I-2 was observed to bind both ssDNA and dsDNA similar to ORC of Saccharomyces cerevisiae and Rx1 (37, 64). Based upon the Cdc6/Orc1 homology with NLR proteins, and the DNA binding characteristics of the isolated R1 NB domain (Fig. 2B), it is most likely that this domain is also the site of DNA binding in the I-2[1–519] protein, although a role for the CC domain cannot be formally excluded. Unfortunately it was not possible to generate truncated I-2 protein lacking the CC domain to test this hypothesis. Eukaryotic ORCs and Rx1 lack DNA sequence specificity in vitro, and this property is also shared with I-2. Superficially, therefore, I-2 biochemistry and ability to interact with nucleotides appears similar to Rx1. Similar to Rx1, the I-2 protein used for analysis is lacking its LRR domain region. The absence of a domain resembling the LRR in Cdc6/Orc1 proteins validates the use of this truncated protein in assessing DNA binding in NLR proteins. However, we cannot exclude the possibility that the LRR might influence the affinity of the NB-ARC region for DNA through constraints placed on relative domain orientations. These experiments are an important future target that first require the synthesis of significant amounts of full-length NLR protein, currently a considerable technical challenge.

A number of crucial differences, however, between the biochemistry of Rx1 and I-2 are observed. First, Rx1 and I-2 show divergent specificity for different nucleic acid species. Rx1 has a higher affinity for single-stranded nucleic acids (ssDNA and RNA) over double-stranded nucleic acids, whereas I-2 has a higher affinity for dsDNA over RNA. The significance of this for the respective NLRs will await future functional characterization. Both Rx1 and I-2 distort dsDNA but to differing extents. The bend angle introduced into DNA by I-2 (22–33°) is substantially less than that introduced by Rx1 (42°) but of a similar magnitude that as introduced by Orc1-1/Orc1-3 of S. solfatarius (20°) and ORC1 of A. pernix (35°) (37, 47, 49). The most noticeable difference between Rx1 and I-2, however, is that I-2 allows us to make a coupling between its P-loop—required for the switch function—and DNA binding. In contrast to Rx1, I-2 has a measurable ATPase activity, which we found to be stimulated by DNA (Fig. 5A). Furthermore, (nonhydrolyzable) ATP was demonstrated to promote I-2 binding to DNA relative to the ADP-bound state. This observed property is consistent with the switch hypothesis for NB-ARC domain activation stating that the “open” ATP-bound state is the active state triggering immune signaling, whereas the “closed” ADP-bound conformation corresponds to the autoinhibited state (8). Increased DNA binding in the presence of ATP and a nonhydrolyzable ATP analog, when compared with ADP, suggests that DNA binding is stimulated in response to NLR activation upon effector recognition. That ATP-bound and non-nucleotide-bound I-2 show similar binding to DNA suggests that the structure of the empty protein resembles that of the activated state and differs from that of the “closed” ADP bound confirmation. Incubation with DNA stimulates the ATPase activity of I-2, suggesting that DNA binding is a self-limiting process, and it is tempting to speculate that DNA binding and release is a cyclic process triggered by the presence of the effector. A similar cyclic mechanism, allowing multiple rounds of effector recognition providing a means for signal amplification, has been proposed before for Rx and was based on interaction studies of the CC-NB-ARC and LRR domains (65). Further study is required to assess the significance and generality of these findings for other NLRs.

Additional evidence linking coupling of the activation state and switch function of the NB-ARC to DNA binding comes from the analysis of an I-2 P-loop mutant. Wild type I-2 and a variant with a defective P-loop have a similar affinity for double-stranded DNA as assessed by EMSA. However, measurements using fluorescence anisotropy reveal a reduced affinity for the P-loop mutant. These data suggest an overall different shape for the protein-DNA complex for the wild type and P-loop mutant proteins. This difference is not due to alterations in conformation of the bound DNA because the observed pattern of DNA distortion, measured through DNA bending and melting, is identical in both cases. It is reasonable to assume, therefore, that the difference in topology is attributed to the protein. In support of this hypothesis, equivalent mutations in the NB domain of Cdc6 have been shown to affect its ability to
interact with other proteins and form complexes at dsDNA (66, 67). It is formally possible that anisotropy, but not EMSA, detects an I-2 complex that forms for the wild type, but not the mutant protein. Such a complex could result in a higher observed anisotropy. These observations provide insight into the molecular mechanism for how P-loop mutations, commonly used to investigate the role of NLR switch function in immunity, might exert their activity. NLR variants with P-loop modifications, such as the I-2 K207R mutant, showing a decreased $K_m$ [ATP] are defective in immune activation. At cellular [ATP], this defect may not necessarily contribute to decreased ATP binding but may be a reflection of an altered conformational state that could influence trafficking (29) or its activity at DNA (66, 67). Unfortunately, chimeric fusions of I-2 with GFP are nonfunctional, so we cannot use microscopic methods to investigate the effect P-loop mutants have on I-2-mediated DNA binding in planta in relation to immune signaling (37).

In summary, we have identified a DNA binding and distorting activity in the I-2 protein in vitro. This establishes nucleic acid binding as a conserved biochemical feature for at least a subset of plant NLR proteins (1-2, R1, Rx1, and PsIP). A further conserved feature of these NLR proteins in DNA manipulation is through bending and melting of the double helix. We further establish that individual NLR proteins distinctively differ in the interactions with various nucleotide topologies (ssDNA, dsDNA and RNA) that might arise from their functional diversity. DNA binding in I-2 is directly linked to the switch function of its central NB-ARC domain, thus directly linking activity at DNA to known biochemical requirements for NLR activation.

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References

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