Metal Cofactors in the Structure and Activity of the Fowlpox Resolvase

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Poxvirus DNA replication generates linear concatemers containing many copies of the viral genome with inverted repeat sequences at the junctions between monomers. The inverted repeats refold to generate Holliday junctions, which are cleaved by the virus-encoded resolvase enzyme to form unit-length genomes. Here we report studies of the influence of metal cofactors on the activity and structure of the resolvase of fowlpox virus, which provides a tractable model for in vitro studies. Small-molecule inhibitors of related enzymes bind simultaneously to metal cofactors and nearby surface amino acid residues, so understanding enzyme–cofactor interactions is important for the design of antiviral agents. Analysis of inferred active-site residues (D7, E60, K102, D132, and D135) by mutagenesis and metal rescue experiments specified residues that contribute to binding metal ions and that multiple binding sites are probably involved. Differential electrophoretic analysis was used to map the conformation of the DNA junction when bound by resolvase. For the wild-type complex in the presence of EDTA (ethylenediaminetetraacetic acid) or Ca2+, migration was consistent with the DNA arms arranged in near-tetrahedral geometry. However, the D7N active-site mutant resolvase held the arms in a more planar arrangement in EDTA, Ca2+, or Mg2+ conditions, implicating metal-dependent contacts at the active site in the larger architecture of the complex. These data show how divalent metals dictate the conformation of FPV resolvase–DNA complexes and subsequent DNA cleavage.

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Introduction

Poxviruses encode a Holliday junction (HJ) resolving enzyme (designated A22 in vaccinia and fowlpox) that is a potential target for antiviral agents. During poxvirus DNA replication, a concatenated form of the viral DNA is initially produced. The junctions between monomer units within the concatamer form inverted repeat sequences.¹ These sequences can be extruded from the DNA double helix to form a cruciform, which contains a HJ at its base.² The poxvirus resolvase can cleave such structures in vitro³–⁶ and inactivation of resolvase during infection results in accumulation of uncleaved concatemers, indicating that resolvase is required for concatemer resolution in vivo.⁷ After cleavage, ligation of the nicked DNA ends forms the covalently closed hairpin termini characteristic of poxviruses.

The poxvirus resolvase enzymes are members of the RNase H superfamily of DNA-modifying enzymes.³ Each enzyme in the superfamily shares a similar protein fold in the catalytic domain, containing a central five-stranded mixed beta-sheet sandwiched between alpha-helices.⁸ Each active site is thought to be composed of three or four acidic residues. For those family members that have been purified and studied in vitro, each was found to require divalent metal ions for activity. Structural and functional studies have suggested that multiple metal atoms may bind the conserved acidic residues to form the catalytic center, and

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Abbreviations used: FPV, fowlpox virus; HJ, Holliday junction (DNA four-way junction); EDTA, ethylenediaminetetraacetic acid; TBE, Tris–borate–EDTA.
specific models have been proposed based on two metal mechanisms. Members of the RNase H superfamily catalyze nucleophilic substitution at a nucleic acid phosphodiester bond, where a hydroxyl oxygen acts as the nucleophile, attacking the phosphorus atom, and the 3' oxyanion serves as the leaving group. For RNase H, this involves attack of water on the RNA strand of an RNA–DNA hybrid. The RNase H superfamily also includes HJ resolving enzymes, the most thoroughly studied of which is Escherichia coli RuvC. For these enzymes, hydrolysis takes place at a DNA branch point, freeing the individual DNA duplexes from one another. The retroviral integrases, also members of the superfamily, catalyze the attack of a hydroxyl at the 3' end of a donor DNA chain on target DNA strand, thus achieving strand joining by a single-step transesterification.

The HIV integrase enzyme is the target of the FDA-approved drug raltegravir, raising the hope that related drugs might be devised that target poxvirus resolvase and inhibit poxvirus replication. Raltegravir is believed to bind to integrase in part by chelating two metal atoms at the enzyme active site. Pendant groups on raltegravir make additional contacts to nearby positions on the integrase surface that confer specificity. Because we are interested in devising poxvirus resolvase inhibitors, we carried out studies to investigate metal atom binding at the active site by the poxvirus resolvase, the likely binding site for raltegravir-like molecules. Previous work with poxvirus resolvases has focused on the vaccinia virus enzyme and has been limited by the fact that vaccinia resolvase is relatively insoluble and thus difficult to study in vitro. Here we present studies of the role of metal ions in catalysis and the overall geometry of the protein–DNA complex using purified fowlpox resolvase, which is more soluble and therefore a more tractable model.

We first examined metal ion cofactor specificity for catalysis by fowlpox resolvase by testing seven different divalent metal ions in HJ resolution reactions. Next, we used 14 mutant fowlpox resolvase proteins, each with a single-amino-acid substitution at one of five putative active-site residues, to probe for residues required for efficient catalysis. By comparing the catalytic activity of Ala-, Cys-, and amide-substituted proteins under different divalent metal conditions, we were also able to make inferences as to which active-site residues may be involved in direct metal chelation. We then studied the HJ-binding properties of each mutant protein to ensure that the substituted proteins contained no gross defects in DNA binding activity compared to the wild-type protein.

Using the mutants and assembly conditions developed in the above studies, we carried out comparative electrophoresis analysis to determine the geometry of the HJ DNA arms in resolvase complexes. HJ DNAs containing each of the six possible pairs of shortened DNA arms were synthesized, and the mobilities of the free DNA and resolvase complexes were compared by polyacrylamide gel electrophoresis. In the only two previous studies where predicted complex structures deduced from electrophoretic data could later be compared to X-ray crystallographic data, the predicted structures matched the X-ray data. The electrophoretic approach has the advantage that complexes are not subject to possible distortion as a result of crystal packing forces, although the resolution of the electrophoretic method is much lower. We found that the orientation of the arms differed between that in the fowlpox virus (FPV) resolvase–DNA complex and that of free DNA and also differed in the presence of divalent metal ions. The overall complex conformation, although similar to those of other family enzyme members studied previously, contained a measurable degree of asymmetry, a phenomenon not previously observed for a resolvase–HJ complex.

Together, these data show how metal atoms interact at the FPV resolvase active site and also how metals dictate the conformation of the complex. This study provides background and methods for the design of small-molecule inhibitors for the poxvirus resolvase potentially useful as antiviral therapy.

### Results

#### Analysis of the divalent metal requirements for catalysis by FPV resolvase

Previous work has established that the vaccinia virus resolvase requires either Mg$^{2+}$ or Mn$^{2+}$ for its DNA cleavage activity. To investigate the requirements of FPV resolvase, we incubated the enzyme with a fluorescein-tagged DNA HJ (HJ$_{FS}$) substrate under various metal ion conditions and then analyzed the products by polyacrylamide gel electrophoresis (data not shown). We observed product formation in the presence of either Mg$^{2+}$ or Mn$^{2+}$, but not in the presence of ethylenediaminetetraacetic acid (EDTA). Previously we estimated the first-order rate of HJ cleavage to be 0.23 s$^{-1}$ at 25 °C in 15 mM Mg$^{2+}$ under single turnover conditions. We also tested Ca$^{2+}$, Cu$^{2+}$, Fe$^{2+}$, Ni$^{2+}$, and Zn$^{2+}$ at concentrations ranging from 0.4 to 30 mM. Slight cleavage was seen in Ca$^{2+}$, Fe$^{2+}$, Ni$^{2+}$, and Zn$^{2+}$. No cleavage was observed in Ca$^{2+}$. Thus, only Mg$^{2+}$ and Mn$^{2+}$ act efficiently as cofactors, paralleling studies of other RNase H superfamily members.

We also noticed that the titration curve for Mn$^{2+}$ showed a sharp drop in activity between 1.1 and 3.3 mM. This suggested that higher concentrations of Mn$^{2+}$ may be inhibitory, and tests with mixtures of Mn$^{2+}$ and Mg$^{2+}$ showed indeed that concentrations of Mn$^{2+}$ above 3.3 mM inhibited cleavage (data not shown).

#### Catalytic activity of fowlpox resolvase mutants

We wished to identify and analyze the amino acid side chains that constitute the fowlpox resolvase
active site. Figure 1a shows an alignment of conserved residues from known resolvases of the RNase H superfamily, and Fig. 1b shows their positions on a model of the structure inferred from the crystal structure of *E. coli* RuvC. Five conserved residues can be seen—D7, E60, K102, D132, and D135 (fowlpox resolvase numbering scheme). Each of these residues has been implicated as important for function in vaccinia resolvase.5

In order to address whether the conserved acidic residues contribute to catalysis in FPV resolvase by binding directly to metal atoms, we compared substitutions that remove the side-chain functional group (Ala) with those that preserve some potential for metal binding (Asn/Gln or Cys). The Cys substitution is of particular interest because thiolate binds more tightly to Mn2+ than to Mg2+. Thus, if Cys substitutions retain catalytic activity in Mn2+ but not Mg2+, it suggests that the side chain binds directly to the divalent metal ion cofactor.30–33

Each resolvase derivative was incubated with HJ substrate in the presence of 1 mM EDTA, 15 mM Mg2+, 0.5 mM Mn2+, or 5 mM Mn2+. Data for all the amino acid substitution derivatives of FPV resolvase

![Fig. 1. DNA binding and cleavage by FPV resolvase and active-site mutants.](image)

(a) Protein sequence alignment of four conserved motifs of various RNase H-like family HJ resolvases (adapted from Lilley and White12). Shading indicates residues that are 100% conserved within the RNase H-like resolvases. Amino acid numbering refers to fowlpox virus. *S. cere, Saccharomyces cerevisiae*; *C. alb*, *Candida albicans*; *S. pom*, *Schizosaccharomyces pombe*; *E. coli*, *Escherichia coli*; *Myxoma*, myxoma virus; *Mollusc*, molluscum contagiosum virus; *Yaba*, Yaba monkey tumor virus; *Vaccini*, vaccinia virus; *Fowlpox*, fowlpox virus. (b) Model of fowlpox resolvase tertiary structure showing the inferred active site, constructed with SWISS-MODEL29 using the RuvC crystal structure as a template. (c) Native PAGE analysis of a representative metal rescue experiments. DNA cleavage reactions for the indicated amino acid substituted proteins were carried out in the various metal conditions indicated above the gel. EDTA, 1 mM EDTA; Mg2+, 15 mM MgCl2; Mn2+Lo, 0.5 mM MnCl2; Mn2+Hi, 5 mM MnCl2. Reaction products are labeled to the right of the gel. HJ, Holliday junction substrate. (d) Native PAGE analysis of DNA binding by wild-type protein in the presence of duplex or HJ competitor DNA. DNA binding reactions were performed with various amounts of wild-type protein and 2 nM HJ DNA in the presence of either 1 mM EDTA (EDTA) or 1 mM CaCl2 (Ca2+). Reaction products are labeled to the right of the gel. Numbers above the gel lanes indicate the fold excess of competitor DNA to probe DNA (wt/wt). (e) Binding of wild-type FPV resolvase to DNA in the presence of EDTA or Ca2+. DNA binding reactions were performed with various amounts of wild-type protein and 2 nM HJ DNA in the presence of either 1 mM EDTA (EDTA) or 1 mM CaCl2 (Ca2+). –, no protein; filled triangles indicate increasing amounts of protein from left to right: 0.3, 0.9, 2.6, 7.9, 24, 71, 213, 640 ng (the 640-ng amount corresponds to 1.6 μM). Resolvase–DNA complexes are labeled to the right of the gel. (f) Effect of Ca2+ on DNA binding by amino acid substituted resolvases. DNA binding reactions were performed with 600 ng of the indicated protein (1.5 μM) and 2 nM HJ probe DNA in the presence of either 1 mM EDTA (EDTA) or 1 mM CaCl2 (Ca2+).
studied are presented in Table 1. A 4-h incubation time was used to allow comparison of product accumulation for low-activity mutant proteins. Table 2 presents kinetic data comparing estimated initial rates for the metal rescue studies in which FPV resolvase–HJ complexes were preformed, then reactions were initiated by addition of metal cofactor. Reaction progression curves are shown in the Supplementary Report. Results for the different amino acid substitutions are presented below, starting from the N-terminus.

Figure 1c shows a representative gel image for reactions with the wild-type (WT), D7A, D7C, and D7N enzymes. Reactions containing wild-type protein showed no cleavage in the presence of EDTA (Fig. 1c, lane marked “EDTA”), Reactions with Mn2+ or the lower concentration of Mn2+ showed near-complete cleavage of the substrate, whereas reactions with the higher concentration of Mn2+ showed detectable cleavage, but only ~25% of the substrate was consumed. We also examined cleavage of a different branched DNA substrate and obtained similar results (data not shown). Substitution of Ala for the conserved Asp position 7 (D7A) eliminated detectable activity in all the metal conditions. The substitution mutant D7N also was inactive, emphasizing the importance of the acidic side chain. However, D7C showed low but detectable activity in the presence of Mn2+ and barely detectable activity in Mn2+ but lower activity in Mg2+, a ratio of rates of at least 18-fold (Table 2). Thus, the observed Mn2+ rescue is as close to the wild type.

Table 1. Cleavage and DNA binding by FPV resolvase with active-site substitutions

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>15 mM Mg2+</th>
<th>0.5 mM Mn2+</th>
<th>5 mM Mn2+</th>
<th>Kd (nM) ± SE</th>
<th>95% Confidence interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>95.5 ± 2.2, n=13</td>
<td>98.1 ± 3.9, n=9</td>
<td>24.9 ± 2.8, n=9</td>
<td>53 ± 14</td>
<td>39–73</td>
</tr>
<tr>
<td>D7A</td>
<td>0.0 ± 0.3, n=3</td>
<td>0.0 ± 0.0, n=2</td>
<td>0.0 ± 0.3, n=3</td>
<td>128 ± 4</td>
<td>12–1419</td>
</tr>
<tr>
<td>D7C</td>
<td>0.1 ± 0.4, n=2</td>
<td>6.8 ± 0.7, n=2</td>
<td>14.1 ± 1.4, n=3</td>
<td>&lt;22 ± 4</td>
<td>NA</td>
</tr>
<tr>
<td>D7N</td>
<td>0.2 ± 0.1, n=3</td>
<td>0.1 ± 0.5, n=2</td>
<td>0.1 ± 1.2, n=3</td>
<td>36 ± 8</td>
<td>14–94</td>
</tr>
<tr>
<td>E33A</td>
<td>95.9 ± 3.2, n=3</td>
<td>58.1 ± 2.2, n=2</td>
<td>5.0 ± 1.3, n=3</td>
<td>1413 ± 147</td>
<td>39–524</td>
</tr>
<tr>
<td>D55A</td>
<td>98.5 ± 3.3, n=3</td>
<td>57.6 ± 3.4, n=2</td>
<td>4.6 ± 2.7, n=3</td>
<td>760 ± 42</td>
<td>254–2270</td>
</tr>
<tr>
<td>E60A</td>
<td>0.7 ± 0.7, n=3</td>
<td>1.0 ± 0.2, n=2</td>
<td>0.6 ± 1.2, n=2</td>
<td>38 ± 8</td>
<td>7–220</td>
</tr>
<tr>
<td>E60C</td>
<td>0.5 ± 0.1, n=3</td>
<td>8.3 ± 1.7, n=2</td>
<td>11.5 ± 2.4, n=2</td>
<td>5 ± 1</td>
<td>0–254</td>
</tr>
<tr>
<td>E60N</td>
<td>1.1 ± 0.9, n=3</td>
<td>38.1 ± 13.6, n=2</td>
<td>57.5 ± 7.1, n=3</td>
<td>21 ± 5</td>
<td>4–114</td>
</tr>
<tr>
<td>E60Q</td>
<td>0.8 ± 1.2, n=3</td>
<td>1.3 ± 0.7, n=2</td>
<td>0.6 ± 2.6, n=3</td>
<td>421 ± 42</td>
<td>147–1205</td>
</tr>
<tr>
<td>K102A</td>
<td>0.9 ± 0.2, n=3</td>
<td>0.6 ± 0.7, n=2</td>
<td>1.1 ± 10, n=3</td>
<td>384 ± 44</td>
<td>187–789</td>
</tr>
<tr>
<td>K102R</td>
<td>93.8 ± 4.9, n=3</td>
<td>88.0 ± 2.5, n=2</td>
<td>2.9 ± 1.5, n=3</td>
<td>77 ± 7</td>
<td>37–162</td>
</tr>
<tr>
<td>D132A</td>
<td>0.2 ± 0.4, n=3</td>
<td>0.9 ± 0.2, n=2</td>
<td>0.4 ± 3.5, n=2</td>
<td>25 ± 5</td>
<td>16–40</td>
</tr>
<tr>
<td>D132C</td>
<td>0.1 ± 0.8, n=3</td>
<td>7.1 ± 1.1, n=2</td>
<td>12.0 ± 3.6, n=2</td>
<td>144 ± 14</td>
<td>103–201</td>
</tr>
<tr>
<td>D135A</td>
<td>0.1 ± 0.4, n=3</td>
<td>94.3 ± 4.9, n=2</td>
<td>12.5 ± 8.5, n=2</td>
<td>14 ± 3</td>
<td>11–17</td>
</tr>
<tr>
<td>D135C</td>
<td>1.7 ± 1.2, n=3</td>
<td>24.3 ± 9.4, n=2</td>
<td>40.2 ± 9.8, n=3</td>
<td>195 ± 19</td>
<td>76–504</td>
</tr>
<tr>
<td>D135N</td>
<td>1.6 ± 1.0, n=3</td>
<td>4.9 ± 1.2, n=2</td>
<td>31.1 ± 6.1, n=2</td>
<td>144 ± 14</td>
<td>92–226</td>
</tr>
</tbody>
</table>

Cleavage activity: Substrate and product bands from native-PAGE analysis of DNA cleavage reactions were quantified and percent product was calculated. The mean and standard error from n independent reactions are reported. Binding activity: Free probe and shifted bands from native-PAGE analysis of DNA binding reactions over a range of protein concentrations were quantified and the fraction bound was calculated. To compare binding affinities between enzymes, we estimated Kd values using nonlinear regression, with a resolvase dimer as the binding species. The normalized data were plotted as fraction bound versus log[protein] and fit with a sigmoidal dose-response model. Between 3 and 29 independent measurements went into each Kd measurement with three to six different protein concentrations over a 9-to 243-fold concentration range. NA indicates not appropriate (impossible to evaluate).

Substitution of the nonconserved control positions E33 and D55 with Ala had little effect on activity, indicating that these side chains are not important for catalysis. E60A and E60Q were inactive under all metal conditions. E60C showed slight but detectable activity in Mn2+ but lower activity in Mg2+, again a metal rescue phenotype indicating that the E60 side chain likely binds metal to promote catalysis (Table 2).

Unexpectedly, the E60N substitution was also active in Mn2+ but much less active in Mg2+ (Table 1). One explanation of this observation is that the local environment of the N60 amide, being one C–C bond length closer to the protein backbone than the Q60 amide, is sufficiently more basic (perhaps due to closer proximity to K102) to deprotonate the amide bond length closer to the protein backbone than Q60 amide and permit divalent metal ion chelation and catalysis. Another model holds that the E60N mutant yields a less stable enzyme–substrate complex than E60Q, and this assists product formation.

Table 2. Rate analysis for the metal rescue experiment

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Rate of Cleavage (pM/s)</th>
<th>Ratio of rates, 0.5 mM Mn/15 mM Mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>D7C</td>
<td>0.7543 ± 0.0453</td>
<td>0.0248 ± 0.0192</td>
</tr>
<tr>
<td>E60C</td>
<td>0.2072 ± 0.0301</td>
<td>0.0990 ± 0.0276</td>
</tr>
<tr>
<td>D132C</td>
<td>0.5117 ± 0.0653</td>
<td>0.2135 ± 0.0354</td>
</tr>
<tr>
<td>D135C</td>
<td>5.9166 ± 0.3227</td>
<td>0.2541 ± 0.0510</td>
</tr>
</tbody>
</table>

Reactions were started by addition of metal cofactor. Substrate and product bands from native-PAGE analysis of DNA cleavage reactions were quantified and percent product was calculated. Rates were calculated using a single-exponential decay model. From 15 to 18 points were used to determine each rate.

Confidence interval: 95%
Removing the positive charge at the conserved K102 residue via Ala substitution eliminated detectable cleavage activity under all metal conditions. However, preserving the positive charge by substitution of Arg at this position returned activity to wild-type levels (Table 1).

Substitution of Ala at D132 eliminated detectable cleavage activity under all metal conditions. The D132C substituted enzyme showed weak activity in Mn$^{2+}$ and lower activity in Mg$^{2+}$, consistent with a role for the side chain in binding metal during catalysis.

For D135, Asn and Cys substitutions showed slight activity in the presence of Mn$^{2+}$ but much less in Mg$^{2+}$ (Table 1). Substitution with Cys showed a 23-fold higher rate in Mn$^{2+}$, suggesting a role for metal binding (Table 2). Substitution with Ala, a side chain not expected to participate in metal coordination, resulted in loss of activity in Mg$^{2+}$, but unexpectedly retained activity in Mn$^{2+}$, with low Mn$^{2+}$ being more active than high Mn$^{2+}$ as observed for the wild-type enzyme. This complicates the interpretation and suggests that D135 plays a more specialized role in catalysis (we return to this point in the Discussion).

In summary, the metal rescue study implicated D7, E60, D132, and D135 in metal binding during catalysis. Some caution is warranted in interpreting these data because (i) the cleavage rates of the Mn$^{2+}$-rescued Cys substitutions were not restored to wild-type rates, and (ii) the rate-limiting steps in the catalytic cycle were not determined for each mutant and the wild-type enzyme, so it is possible the rate-limiting steps differed. However, one simple interpretation is that the four acidic residues show Mn$^{2+}$ rescue of Cys substitutions because these residues promote catalysis by binding the metal cofactor.

**DNA binding by mutant and wild-type fowlpox resolvases**

An advantage of the fowlpox resolvase model system is that, unlike vaccinia resolvase, simple band shift assays of DNA binding are feasible due to better protein solubility. We previously reported that FPV bound to HJ DNA with a $K_d$ of $2 \times 10^{-8}$ M in the absence of competitor DNA. All of the amino-acid-substituted resolvases studied above were analyzed by band shift assays with the HJ probe in the presence of 300-fold excess of sonicated salmon sperm competitor DNA (Table 1). To prevent cleavage by resolvase during the reaction, binding studies were carried out in the presence of 1 mM EDTA to chelate divalent metals. Several DNA complexes were formed by fowlpox resolvase and are designated as c0, c1, and c2 beside the gels in Fig. 1d–f.

Resolvase binding is specific for branched DNA. To demonstrate the specificity on our fluorescein-tagged HJ substrate, fowlpox resolvase was added to mixtures of the labeled HJ probe and an unlabeled duplex competitor DNA (Fig. 1d). We found that a >2500-fold excess duplex linear competitor DNA was needed to compete for HJ binding (Fig. 1d, duplex). In contrast, only a 3-fold excess HJ competitor DNA was required to observe a reduction in binding (Fig. 1d, lanes marked “HJ”).

Using the HJ substrate in the presence of EDTA (Table 1), we tested each of the fowlpox resolvase mutants for DNA binding activity. Examples of the quantitative binding data are presented in the Supplementary Report. All mutants bound to DNA detectably, but there was considerable variation among them. Errors for some measurements were large because only a narrow concentration range of protein could be interrogated. Two mutant proteins bound more tightly than the wild type (D7C and D135A). These mutants may be useful in efforts to form resolvase–DNA complexes for biophysical and structural analysis. Several showed impaired binding (E33A, D55A, E60Q, K102A, D132C, D135C, and D135N). In summary, all of the FPV resolvase proteins with amino acid substitutions at the four conserved acidic residues showed detectable formation of c1, although some showed altered affinity. Several proteins showed primarily formation of c0 only (E33A, D55A, and K102A), so for these we investigated binding in the presence of metal (instead of EDTA as was used above).

When Ca$^{2+}$ was included in binding assays, the affinity of wild-type resolvase was increased approximately twofold (Fig. 1e). We then studied the effect of Ca$^{2+}$ on binding by K102A and K102R (Fig. 1f) and the controls E33A and D55A (data not shown). All showed increased binding in the presence of Ca$^{2+}$, suggesting that the impairment in DNA binding for these proteins in the presence of EDTA was partially overcome by addition of metal. This helps to explain why E33A and D55A showed little impairment in the cleavage reactions. We also observed some retention of DNA in the gel wells (<5% of total DNA) after electrophoresis in binding experiments with E33A, D55A, and K102A, suggesting these proteins may be more prone to aggregation than the others studied.

**Oriented cleavage on HJ 1 by FPV resolvase**

In the course of a comparative analysis of FPV resolvase interactions with HJ DNA, we noticed that junction 1 from Lilley and coworkers was cleaved in a highly biased fashion, so that cleavage on one pair of strands was strongly favored over the other (Fig. 2). The reason for working with junction 1 was that it was designed to allow probing of resolvase–HJ DNA complexes by comparative electrophoresis—junction 1 is designed to contain a unique restriction enzyme recognition site in each DNA arm, so that pairs of arms can be conveniently shorted by restriction enzyme cleavage. This makes possible analysis of the geometry of the DNA arms by comparative electrophoresis, as is described below. The finding of oriented cleavage by FPV resolvase provided a means of assembling oriented FPV resolvase–HJ DNA complexes for such studies.
Figure 2 documents the oriented cleavage using junction 1S, which is a version of junction 1 synthesized with shorter DNA arms, which we used to improve electrophoretic separation on DNA sequencing-type gels. Incubation of junction 1S with FPV resolvase resulted in cleavage to near-completion on strands H and X (Fig. 2, lanes 6 and 14), whereas only barely detectable cleavage was seen on strands B and R (Fig. 2, lanes 2 and 10). For comparison, the yeast Cce1 enzyme showed weak cleavage on strand R only (Fig. 2, lane 11), indicating that junction 1S lacked sequence features required for efficient cleavage by Cce1. In contrast, T7 endonuclease I cleaved efficiently on all four strands (Fig. 2, lanes 4, 8, 12, and 16). Kinetic analysis showed that cleavage with FPV resolvase was slightly faster on the X strand than on the H strand, consistent with the asymmetry in binding described below, although the difference was slight (data not shown).

We next used order of addition experiments to investigate the role of metal atoms in the formation of the oriented cleavage complex. We compared cleavage in reactions where (i) DNA was mixed with metal, then started with enzyme, or (ii) DNA was mixed with enzyme, then the reaction started with metal. In both cases, oriented cleavage was seen (data not shown). This is consistent with the idea that formation of the oriented complex does not depend on the presence of metal, although we cannot rule out an alternative explanation that the complex is in fast equilibrium, so that the enzyme releases, binds metal, then rebinds in an oriented fashion.

Comparative gel electrophoresis of FPV resolvase–HJ 1 complexes

Using comparative electrophoresis, we took advantage of the ability to assemble oriented FPV resolvase–DNA complexes to analyze the geometrical positions of the DNA arms. In previous studies, this method successfully predicted the global orientation of the DNA arms in a variety of resolvase–HJ complexes, and the predictions have been confirmed by X-ray crystallography for T7 endonuclease I and T4 endonuclease VII bound to HJ DNA. 18–23

To carry out the analysis, six different HJ DNAs were synthesized, all of which contain the same numbers of base pairs, but each of which had one of the six possible combinations of two long DNA arms and two short arms. 18 For any given HJ species, the angle between the two long arms of the junction mainly dictates the mobility of the junction through a polyacrylamide gel matrix, with angles nearer 180° resulting in faster mobility and relatively smaller angles resulting in slower mobility. Assuming that the geometries of the arms are consistent for resolvase–DNA complexes containing all six variants, then electrophoresis of the
six species side by side allows determination of the relative angles between all pairs of arms of the junction. In this study, we used junction 1, a HJ previously analyzed by this method to allow for comparison to previous work.18

We prepared FPV resolvase–DNA complexes with junction 1 using both the wild-type enzyme and the active-site mutant enzyme D7N. As a control, we also studied the six species in the absence of protein and obtained data identical to published results.18 To analyze the effects of metal on the conformation of the DNA, we compared 2 mM EDTA, 200 μM CaCl2, or 200 μM MgCl2. In each case, the complex was incubated in the indicated metal or in EDTA prior to electrophoresis, and the metal or EDTA was added to the electrophoretic buffer as well. Our finding that Ca2+ does not support DNA cleavage allowed us to study the wild-type enzyme under divalent metal ion conditions without DNA cleavage. Use of the active-site mutant D7N allowed us to study FPV resolvase in the presence of Mg2+ without concurrent DNA cleavage. DNA and FPV resolvase–DNA complexes were analyzed by electrophoresis on 8% native polyacrylamide gels.

In the absence of divalent metal ion and resolvase protein (Fig. 3a), we obtained an electrophoretic

![Fig. 3](image)

**Fig. 3.** Conformation of FPV resolvase–HJ complexes analyzed by comparative electrophoresis in the presence of 2 mM EDTA. (a) Comparative gel electrophoresis analysis of fowlpox resolvase–DNA complexes. The long arms of the junctions are designated above the gel lanes by the letters B, H, R, or X. The protein used in each binding reaction is indicated above the gel. Buffer, no protein; WT, wild-type protein; D7N, D7N mutant protein. (b) Structural interpretation of comparative electrophoresis data for the buffer-only control. Each cylinder represents a double-helical arm of the HJ and its identity is labeled by the letters B, H, R, or X. The lack of continuity between adjacent cylinders indicates that the DNA arms are unstacked. More cone-shaped cylinders indicate angling of that DNA arm out of the plane of the paper. (c) Structural interpretation of the data for complex with wild-type FPV. The DNA arms of the complex may be mobile. (d) Structural interpretation of the data for complex with D7N resolvase. More cone-shaped cylinders indicate angling of that DNA arm out of the plane of the paper.
pattern for free junction 1 of slow-fast-slow-slow-fast-slow. This pattern matches previous results and corresponds to an open square planar conformation of junction 1 in which all four arms of the junction lie in the same plane and the angle between each set of adjacent arms is $\sim 90^\circ$ (Fig. 3b). In the comparative electrophoresis experiment with wild-type or D7N resolvase in EDTA, we observed discrete, slower-migrating bands, as expected for formation of FPV resolvase-DNA complexes. In each case, we observed a change in the relative electrophoretic pattern of the six species between the unbound and bound conditions, indicating that HJ binding by FPV resolvase altered the global conformation of the DNA junction.

For the wild-type samples, we observed a relative electrophoretic pattern in which the difference among the six species showed only slight differences (Fig. 3a, “WT”). To the extent that they differed, the samples showed slow-fast-slow-intermediate-intermediate-intermediate mobilities (Fig. 3a). The extreme case of all species having the same mobility would imply either that (i) all pairs of arms have identical angles between them, which can only be satisfied by a tetrahedral geometry, or that (ii) the junction is mobile so that the geometry changes during electrophoresis, resulting in mobilities that reflect multiple conformations. An example of rapid exchange in HJ conformation has been reported previously.\textsuperscript{18,22} If we assume a single geometry, we interpret the observed pattern to arise from a conformation of the DNA lying somewhere between the tetrahedral conformation (all arm–arm angles $\sim 109.5^\circ$) and a conformation in which $\angle BR$ is $\sim 180^\circ$ and the other five arm–arm angles are $\sim 90^\circ$, although it may be more likely that the junction is mobile during electrophoresis.

For the D7N active-site mutant samples analyzed in the absence of divalent metal ion (Fig. 3a and d), we observed an electrophoretic pattern of slow-fast-slow-intermediate-fast-intermediate. We interpret this pattern to arise from a structure close to a 2-fold symmetric open-X conformation, where all four arms lie in the same plane and $\angle BH$ and $\angle RX$ are $<90^\circ$ and $\angle HR$ and $\angle BX$ are $>90^\circ$. However, a true 2-fold symmetric open-X structure would have a pattern of slow-fast-slow-intermediate-fast-slow, as is approximated more closely in Fig. 4c ($\text{Mg}^{2+}$/D7N) and Fig. 5d ($\text{Ca}^{2+}$/D7N). We suggest the deviation from this pattern observed in Fig. 3a and d may be due to the X arm coming slightly out of plane under these conditions, thus deviating from 2-fold symmetry. It is also possible that the other arms have at least modest out-of-plane bending toward a tetrahedral arrangement and that the $\angle XH$ angle is smaller than the others.

In the presence of divalent metal ion (Fig. 4 shows data for $\text{Mg}^{2+}$ and Fig. 5 shows data for $\text{Ca}^{2+}$), we obtained an electrophoretic pattern for free junction 1 of fast-intermediate-slow-slow-intermediate-fast (Figs. 4 and 5, buffer only).

This pattern matches previous analysis of this junction and corresponds to the antiparallel stacked-X conformation in which the arms of the junction are coaxially stacked upon one another with B stacked on H and R stacked on X.

For the samples containing wild-type or active-site mutant FPV resolvase, we observed discrete slower-migrating bands, as expected for formation of protein–DNA complexes. In each case, we observed a sharp change in the relative electrophoretic pattern of the six species between the unbound DNA and the bound DNA, signifying that HJ binding by FPV resolvase also altered the global conformation of the DNA junction in the presence of divalent metal ion.

For the active-site mutant D7N in the presence of divalent metal ion ($\text{Ca}^{2+}$ or $\text{Mg}^{2+}$), we observed an
electrophoretic pattern of slow-fast-intermediate-intermediate-fast-slow (Fig. 4a and c, Mg2+/D7N, and Fig. 5a and d, Ca2+/D7N). We interpret this pattern to arise from an approximately 2-fold symmetric open-X conformation, where all four arms lie in the same plane and \( \angle BH \) and \( \angle RX \) are \( 90^\circ \) and \( \angle HR \) and \( \angle BX \) are \( >90^\circ \).

The wild-type FPV could only be studied under metal conditions in Ca2+, because in Mg2+ the enzyme cleaves DNA. For wild-type FPV resolvase in the presence of Ca2+, we observed an electrophoretic pattern of slow-fast-slow-intermediate-fast, close to the pattern observed for the active-site mutant under EDTA conditions (Fig. 5b, compare EDTA–D7N to Ca2+-WT). We interpret this pattern again to arise from an open-X structure with a slightly out of plane X arm (see above).

**Discussion**

Here we present a study of the role of divalent metal ions in the structure and activity of FPV resolvase. Analysis of metal dependence and amino acid substitutions delineated the requirements for efficient catalysis. The active-site mapping yielded results closely paralleling previous studies of vaccinia virus resolvase and other RNaseH superfamily enzymes. Our use of metal rescue experiments helped establish the roles for some of these active-site constituents in metal binding. In the course of these studies, we found that we could assemble FPV protein onto the junction 1 substrate in a defined orientation. This provided the tools necessary to analyze the geometry of the DNA arms by comparative electrophoresis in oriented complexes, which yielded results that paralleled but were not identical

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**Fig. 5.** Conformation of FPV resolvase–HJ complexes analyzed by comparative electrophoresis in the presence of 200 \( \mu M \) CaCl2. (a) Comparative gel electrophoresis analysis of fowlpox resolvase–DNA complexes. Markings as in Fig. 3. (b) Structural interpretation of comparative electrophoresis data for buffer only, (c) wild-type FPV resolvase, or (d) D7N. Markings as in Fig. 3. The continuity of adjacent cylinders seen in the buffer-only condition indicates that the adjacent arms of the HJ are coaxially stacked.
to previous studies of the RNaseH superfamily resolvase RuvC. Some of the implications of these studies are discussed below.

Residues that were likely components of the FPV resolvase active site could be identified by homology to other RNase H superfamily members. All five of the conserved charged residues in vaccinia resolvase have been substituted with neutral residues, and for each cleavage activity was abolished. However, for the vaccinia protein, due to the poor solubility of the purified proteins it was not possible to assess whether DNA binding was affected by the substitutions, leaving the functions of the residues uncertain. For fowlpox resolvase, substitution of the five conserved charged residues with neutral residues abolished the catalytic activity in Mg$^{2+}$, and all proteins could be shown to retain DNA binding activity. As controls, two nonconserved acidic residues were substituted with Ala, and both of these proteins retained catalytic activity.

Thus, studies of the amino acid substitutions established the importance of these residues for function, but not whether they act by binding metal. Studies of Cys substitutions at the four conserved acidic residues showed detectable restoration of activity in the presence of Mn$^{2+}$ compared to the Ala substitutions. Because Cys binds Mn$^{2+}$ more tightly than Mg$^{2+}$, these findings indicate that residues at this position likely promote catalysis by binding metal. Some caution is warranted for this conclusion, since the rate-limiting step was not determined for the wild-type and mutants, and this may differ, but differential metal binding offers a reasonable model. We also studied substitutions of Asn and/or Gln at three of the conserved acidic residues and some of these proteins also displayed detectable activity in Mn$^{2+}$. Our studies of the conserved residue D135 revealed an unexpected result that differed from the other three conserved acidic residues. Ala substitution at D135 resulted in loss of detectable activity in Mg$^{2+}$, but preservation of activity in Mn$^{2+}$; however, the D135-carboxylate is a requirement for catalysis in Mg$^{2+}$. Given that Mg$^{2+}$ has stricter coordination geometry than Mn$^{2+}$ and is believed to be the physiologic cofactor for this superfamily of enzymes, we speculate the role of D135 is to stabilize a geometry of Mg$^{2+}$ at the enzyme active site that is necessary for catalysis. Also, our finding that the D135A enzyme retains inhibition with high Mn$^{2+}$ suggests D135 does not directly participate in high Mn$^{2+}$ inhibition.

For K102, Ala substitution at the homologous residue in the E. coli RuvC protein suggested this conserved basic residue played a role in DNA binding. However, our finding that Ala and Arg substitution at K102 have similar DNA binding properties, but only the Ala substitution is inactive for cleavage, suggests the function of this residue may be to stabilize the catalytic orientation of the scissile phosphate or nucleophile within the active site and/or to participate directly in the catalytic mechanism as a general base.

The results from our metal rescue experiments are consistent with a cleavage mechanism involving multiple metal atoms bound at the active site, and this suggests approaches to inhibitor development. Using our model (Fig. 1b), we measured the distances between the side-chain carboxylates of D7, E60, and D132, and found the distance between E60 and D132 to be 11.2 Å, between E60 and D7 to be 5.9 Å, and between D7 and D132 to be 7.4 Å. The distance between E60 and D132 is too large to have both residues coordinate a common Mg$^{2+}$ ion simultaneously, and the same conclusion can be made by inspection of the homologous residues in the RuvC crystal structure. Our metal rescue data suggest that both of these residues do in fact contact metal, however, thus providing biochemical evidence supporting the idea of multiple metal-ion binding sites. This finding has implications for targeting of this enzyme with small-molecule inhibitors. HIV-1 integrase inhibitors of the raltegravir class are proposed to chelate two metal atoms at the integrase active site, and so could potentially bind and inhibit poxvirus resolvases. Inhibitors of this class have been identified in a high-throughput screen for molecules active against fowlpox resolvase and are presently under study (Culyba, Hwang, and Bushman, unpublished data).

The diagrams of the geometry of junction 1 in complex with FPV resolvase in Figs. 3–5 summarize our interpretation of the comparative electrophoresis data. Binding of FPV resolvase to junction 1 alters the global conformation of the HJ in both the absence and presence of divalent metal ions. In the presence of divalent metal ions, the protein–DNA complexes adopt structures that allow adjacent pairs of arms to more closely approach one another. For example, for the WT resolvase–HJ complexes, addition of divalent metal switches the conformation of the arms from a more tetrahedral-like configuration to a configuration where three of the arms of the junction lie approximately within the same plane and the fourth arm is slightly deviated out of this plane. Similarly, for the D7N resolvase–HJ complexes, addition of divalent metal ion switches the conformation of the arms from one arm being slightly out of plane to all four arms residing in approximately the same plane. These effects are likely due in part to binding of the divalent metal ions neutralizing charge repulsion between negatively charged phosphates on the adjacent DNA arms. We also observed a similar effect with the active-site mutant—replacing a negatively charged residue at the active site with a neutral residue in D7N also seemed to help the arms of the junction to more closely approach one another. Since we observed this phenomenon under EDTA conditions, it appears to be a largely divalent metal ion independent effect. One explanation may
be that a neutral residue can accommodate the negatively charged scissile phosphodiester at the active site, whereas the wild-type acidic residue cannot due to charge repulsion, and this leads to a tighter mobility constraint on the arms of the HJ by the active-site mutant protein.

We also observed relative electrophoretic patterns of resolvase–HJ complexes consistent with deviation from 2-fold symmetry of the HJ arm positions. This was unexpected, since no significant asymmetry was noted in the patterns obtained in the absence of protein, and FPV resolvase is expected to bind the HJ as a dimer with 2-fold symmetry. Although the asymmetry was detectable in all the FPV resolvase–DNA electrophoretic patterns, it was most prominent for the D7N protein in the presence of EDTA, where we interpreted the pattern to indicate that three of the arms lie approximately in the same plane and the fourth arm deviating out of plane. We speculate that the observed asymmetry in the patterns may be due to an inherent sequence preference of the enzyme, which is evident also in the highly oriented choice of cleavage sites on junction 1.

Comparative electrophoresis studies have been carried out for bacterial RuvC, yeast Cce1, and the phage enzymes T4 endonuclease VII, T7 endonuclease I, and RusA, allowing comparison to FPV resolvase. Each of these five protein–HJ complexes showed a different electrophoretic pattern, likely representing a unique overall structure for each. The Cce1 complex adopted a square planar structure both in the presence and in the absence of divalent metal ion. The RuvC complex showed a pattern consistent with a 2-fold symmetric X-like structure where all four arms of the HJ lie in the same plane both in the presence and in the absence of divalent metal ion. The T4 endonuclease VII complex showed a pattern consistent with a structure similar to that of RuvC, except that two diametrically opposed arms were deviated out of a plane and toward each other. This pattern also did not change in the presence or absence of divalent metal ion. The T7 endonuclease I complex showed a pattern consistent with the arms of the junction remaining coaxially stacked both in the presence and in the absence of divalent metal ion. Lastly, the RusA structure showed a pattern suggesting a tetrahedral positioning of the junction arms whether in the presence or absence of divalent metal ion.

In comparison to the above patterns, our data are most similar to those proposed for the RuvC, Cce1, and T4 endonuclease VII complexes. In each of these models, the arms of the HJ are unstacked, the branch point is open, and the complex has 2-fold rotational symmetry about the dyad axis of the protein dimer, although for FPV resolvase bound to junction 1, our data indicated slight departures from perfect 2-fold symmetry, perhaps related to an inherent sequence preference. Finally, our investigation of the role of divalent metal ion cofactors in fowlpox resolvase catalysis is consistent with a catalytic mechanism involving multiple metal atoms, similar to that proposed for HIV integrase and other RNase H family enzymes. This implies that raltegravir-like inhibitors may also be active against poxvirus resolvase and useful for treating poxvirus infections.

**Experimental Procedures**

**Site-directed mutagenesis**

In their analysis of the crystal structure of T4 endonuclease VII bound to a HJ, Biertumpfel et al. observed that the S-shaped positive electrostatic surface potential of endonuclease VII explains much of the overall binding mode to the HJ studied. They proposed that other HJ resolving enzymes with similar S-shaped positive electrostatic surface potentials, such as RuvC and Cce1, may also be expected to have binding modes similar to that of endonuclease VII. In the endonuclease VII–HJ crystal structure, the protein dimer contacts the HJ predominantly on the minor groove face of the HJ and the S-shaped positive surface facilitates protein–DNA contacts on all four arms of the junction. Of the known HJ resolving enzymes, poxvirus resolvases share the most sequence similarity with that of RuvC and Cce1, and vaccinia virus resolvase has similarly been shown to bind the HJ as a dimer. Our cleavage site data indicate that the active sites of the FPV resolvase contact the minor-groove surface, since the cleavage sites are 3’ to the branch point of the junction. Assuming a similar binding mode for FPV resolvase as with T4 endonuclease VII, we infer that FPV resolvase interacts predominantly with the minor-groove face of the HJ.

In summary, our structural models for the fowlpox resolvase–HJ complex presented here are most similar to those proposed for the RuvC, Cce1, and T4 endonuclease VII complexes. In each of these models, the arms of the HJ are unstacked, the branch point is open, and the complex has 2-fold rotational symmetry about the dyad axis of the protein dimer, although for FPV resolvase bound to junction 1, our data indicated slight departures from perfect 2-fold symmetry, perhaps related to an inherent sequence preference. Finally, our investigation of the role of divalent metal ion cofactors in fowlpox resolvase catalysis is consistent with a catalytic mechanism involving multiple metal atoms, similar to that proposed for HIV integrase and other RNase H family enzymes. This implies that raltegravir-like inhibitors may also be active against poxvirus resolvase and useful for treating poxvirus infections.

**Experimental Procedures**

**Site-directed mutagenesis**

The HJ resolvase open reading frame was PCR-amplified from FPV DNA, cloned into pET29a, expressed in E. coli, and purified as described. Expression plasmids conferring amino acid changes to the resolvase gene product were constructed using the QuikChange site-directed mutagenesis kit (Stratagene) using the pET29a-resolvase vector as template with the appropriate oligonucleotide primer pairs. The desired DNA nucleotide changes were confirmed by DNA sequencing. Enzyme storage buffer was 300 mM NaCl, 20 mM Hepes (pH 8.0), 1 mM β-mercaptoethanol, 0.1 mM EDTA, and 40% glycerol. In some cases 50 mM arginine plus 50 mM glutamate was added as a solubilizing agent. Cce1 was purified as described.
Construction of synthetic HJs

Sequences of the DNA oligonucleotides used to construct the synthetic HJs are compiled in Supplementary Table 1. HJFS was used for activity assays. Oligonucleotide 1 contains a 6-carboxyfluorescein (F) 3’-end label and was purified by high-performance liquid chromatography (HPLC). Oligonucleotides 2–5 were purified by PAGE. DNA concentrations were determined by UV spectrophotometry. The HJ was assembled by annealing together oligonucleotides 1–4. The 40-bp duplex marker DNA was constructed by annealing together oligonucleotides 1 and 5. Duplex competitor DNA was constructed by annealing together an unlabeled version of oligonucleotide 1 with oligonucleotide 5.Annexing reactions contained 10 μM labeled DNA and 1.1-fold excess unlabeled DNA and were carried out in the presence of 100 mM NaCl by heating to 95 °C and allowing the solutions to cool slowly to room temperature over a period of 90 min. Junction 1 and junction 1* were annealed as described under the section Comparative electrophoresis analysis. The oligonucleotides that compose these DNA junctions were purified by PAGE.

DNA cleavage assays

For tests of mutants in the metal rescue studies, except where noted, 240 nM enzyme was used with 2 nM HJFS (experiments in Table 1) or 20 nM HJFS (experiments in Table 2) in a final volume of 20 μl. Final solution conditions were 25 mM Tris–HCl (pH 8.0), 100 mM NaCl, 1 mM DTT, and 1% glycerol. Metal concentrations used for each reaction are specified in the tables and figure legends. Proteins were diluted in 10 mM Hepes (pH 8.0), 1 mM β-mercaptoethanol, 0.1 mM EDTA, and 10% glycerol prior to addition to cleavage reactions. Reactions were initiated by addition of resolvase (Table 1) or metal cofactor (Table 2). Reaction mixtures were incubated at 37 °C for the indicated times and then stopped with a 5× solution containing 100 mM EDTA, 5% SDS, 30% glycerol, and 0.1% bromophenol blue. Products of the reaction were separat- ed on 10% polyacrylamide gels by electrophoresis under native conditions in 1× TBE (Tris–borate–EDTA) buffer. The fluorescein label was visualized with a Typhoon instrument (GE Healthcare). Gels were scanned for native conditions in 1× TBE buffer.

Radioactivity was quantified with a Typhoon instrument (GE Healthcare). Oligonucleotide 2–5 were 25 mM Tris–HCl (pH 7.9), 100 mM NaCl, and 0.1 mM EDTA, and 1–5% glycerol. Binding reactions were incubated at 37 °C for 15 min and then a 6× loading dye consisting of 30% glycerol and 0.1% bromo- phenol blue was added. Ten microliters of the resultant solution was loaded on a 1.5-mm-thick, 20 cm polyacryl- amide gel (4%) and electrophoresed at ~85 V at 4 °C until the bromophenol blue reached the bottom of the gel. Products of the binding reaction were visualized as above. To compare binding affinities between enzymes, we estimated KD values using nonlinear regression, taking a dimer as the binding species. The normalized data were plotted as fraction bound versus log[protein] and fit with a sigmoid dose–response model. Between 3 and 29 measurements went into each KD measurement using three to six different protein concentrations over a 9- to 243-fold concentration range (we note that due to the narrow protein concentration range studied the error was high for a few samples). The Supplementary Report shows some examples of the quantitative binding data. Curves were fit using Prism software.

Comparative electrophoresis analysis

To study FPV resolvase binding during comparative electrophoresis, we first analyzed the stability of the wild- type enzyme–DNA complex in the presence of EDTA. Competitor challenge experiments showed that dissociation was slow (~1 h for 50% dissociation, data not shown). In addition, FPV resolvase–DNA complexes showed discrete mobilities in gels even during prolonged electrophoresis, further substantiating their stability for both D7N and wild type in both the presence and absence of metal cofactors, all indicating that complexes were sufficiently stable for analysis.

The six long-arm/short-arm versions of junction 1 were prepared by first annealing stoichiometric proportions of the oligonucleotides that compose junction 1 together (see Table S1). DNA concentrations were determined by UV spectrophotometry. Before the annealing reaction, one of the DNA strands was 5’-end-labeled with [32P]ATP, T4 polynucleotide kinase and [γ-32P]ATP, and the excess [γ-32P]ATP was removed with a G50 spin column (GE Healthcare). Annealing reactions contained 200 nM of each oligonucleotide, 100 mM NaCl, and 5 mM MgCl2. Annealing reactions were heated to 95 °C and then slow cooled to room temperature over 5 h. After annealing, junction 1 was digested with the six pairwise combinations of four restriction enzymes (BamHI [B], HindIII [H], EcoRI [R], and XbaI [X]) in reaction buffer containing 10 mM MgCl2, as recom- mended by the manufacturer (New England BioLabs). Each arm of junction 1 contains a unique restriction site (B, H, R, or X, as above). The nomenclature of restriction-digested junction 1 products used in the text denotes the “long arms” (i.e., the nondigested arms), so that junction “BR” refers to junction 1 digested with HindIII and XbaI.

For the MgCl2 or EDTA binding reactions, the digested products were used directly without further purification. For binding reactions carried out in CaCl2, the residual MgCl2 in the DNA solution was first removed by dialysis against 1× TAE buffer [10 mM Tris–HCl (pH 7.9), 1 mM EDTA], so as to prevent DNA cleavage in binding reactions using wild-type fowlpox resolvase.
Binding reactions were carried out by adding 10 pmol (0.2 μg, 1 μM) of FPV resolvase protein to a solution containing 10 fmol (1 nM) of 32P-labeled junction 1 and 30 μg/ml poly(dl-dC)-poly(dl-dC) competitor DNA in a final volume of 10 μl. Solution conditions were 10 mM Tris–HCl (pH 7.9), 100 mM NaCl, 1 mM DTT, 5% glycerol, and either 10 mM EDTA, 1 mM MgCl2, or 1 mM CaCl2. Binding reactions were incubated at 37 °C for 10 min and then one-sixth volume of a loading dye consisting of 15% Ficoll–400 and 0.1% bromophenol blue was added. Binding reactions were loaded directly onto 8.4 % acrylamide gels (width = 30 cm × 45.0 cm × 0.15 cm or 32.5 cm × 37.5 cm × 0.15 cm, 6 mm teeth) running at a constant 500 V. Electrophoresis was carried out at room temperature and stopped after 16 h. The electrophoresis buffer used for the EDTA binding reactions was 1 × TBE buffer [92 mM Tris–borate (pH 8.4), 2 mM EDTA]. The electrophoresis buffers used for the MgCl2 and CaCl2 binding reactions were the same except, instead of 2 mM EDTA, the buffers contained 200 μM MgCl2 and 200 μM CaCl2, respectively. After electrophoresis, the wet gel was exposed to a phosphor screen and the DNA was visualized using a Typhoon instrument (GE Healthcare).

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2010.03.054

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