Zinc is essential for high-affinity DNA binding and recombinase activity of \(\phi C31\) integrase

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ABSTRACT

The mechanism through which the large serine recombinases bind DNA is poorly understood. Alignments of \(\phi C31\) integrase (Int) and its relatives indicate the presence of a conserved motif containing four cysteines resembling a zinc finger. Inductively coupled plasma–mass spectrometry (ICP–MS) confirmed that an Int monomer contains one atom of zinc. Pre-incubation of Int with ethylenediaminetetraacetic acid (EDTA) was detrimental for both recombination activity and DNA binding affinities but full activity could be restored by adding back Zn\(^{2+}\). Mutations in the cysteines and other highly conserved residues yielded proteins that were hypersensitive to proteases, suggesting that without zinc the domain is unfolded. Substitutions in the highly charged region between the conserved cysteines led to lowered DNA binding affinities while circular dichroism revealed that these variant Ints were not greatly affected in overall folding. Int was protected from inhibition by EDTA when DNA containing an attachment site was present suggesting that the zinc finger and the DNA are in close proximity. A truncated mutant of Int, \(\text{hInt}^\text{V371SUGA}\), lacking the putative zinc finger could bind DNA with low affinity. The data are consistent with there being at least two DNA binding motifs in Int one of which is the zinc finger-like motif.

INTRODUCTION

Temperate bacteriophages often encode an integrase (Int) that mediates site specific recombination between phage and host attachment sites, \(\text{attP}\) and \(\text{attB}\), respectively, to produce the integrated prophage flanked by the recombinant sites, \(\text{attL}\) and \(\text{attR}\). Excisive recombination between \(\text{attL}\) and \(\text{attR}\) regenerates the free circular phage genome containing an intact \(\text{attP}\) site and the \(\text{attB}\) site on the chromosome. Phage Ints are either tyrosine Ints and related to \(\lambda\) Int or they are serine Ints of which \(\phi C31\) Int and Bxb1 gpInt are the best studied (1,2). The serine Ints and other large serine recombinases such as some transposases (e.g. TnpX from Tn\(^{4451}\)) have N-terminal domains (NTDs) that are homologous in sequence, structure and function to the catalytic NTDs of the resolvase/invertases family of serine recombinases (1–6). The large serine recombinases however have much larger C-terminal domains (CTDs) of 300–500 amino acids compared with the small, ~40 residue CTDs of the resolvase/invertases. The CTDs of the serine Ints are required for substrate recognition and for controlling the directionality of recombination i.e. integration versus excision (7–9).

\(\phi C31\) Int requires only small (\(<50\) bp) \(\text{attP}\) and \(\text{attB}\) sites for integration (10). \(\text{attP}\) and \(\text{attB}\) share only 32% sequence identity and they each contain imperfect inverted repeats that flank the dinucleotide where the crossover occurs (11,12). \(\text{attL}\) and \(\text{attR}\) are the products of reciprocal DNA exchange between \(\text{attP}\) and \(\text{attB}\) and therefore contain a half site of \(\text{attP}\) and a half site from \(\text{attB}\). Excision (\(\text{attL} \times \text{attR}\) recombination) requires Int but only occurs in the presence of the recombination directionality factor (RDF) or Xis, an accessory protein that controls the directionality of Int (T. Khaleel, E. Younger, A.R. McEwan, M.C.M. Smith, unpublished). The DNA substrates themselves also have a crucial role in controlling Int activity. In the absence of Xis, Int only recombines \(\text{attP}\) and \(\text{attB}\) and has no activity on any other pair of attachment sites (13). Moreover, without Xis, Int only brings \(\text{attP}\) and \(\text{attB}\) together to form a stable synaptic complex, an early stage in the recombination reaction pathway in which Int dimers are thought to bring two attachment sites together by the formation of Int tetramers (14). As no other pair of attachment sites will form stable synaptic complexes, it is clear that substrate binding is an allosteric event where \(\text{attP}\), \(\text{attB}\) and...
**MATERIALS AND METHODS**

**Strains, cultures and plasmids**

*Escherichia coli* DH5α [fhuA2 Δ(argF-lacZ)U169 phoA glnV44 Φ80 Δ(lacZ)M15 gyrA96 recA1 relA1 endA1 thi-1 hsdR17] and *E. coli* BW25113/pIJ790 (18). The DNA-encoding apramycin resistance cassette was encoded by pJ773. *Escherichia coli* BL21 F−ompT gal dcm lon hsdS8 (F− mB−), λ (DE3), pLysS was used as a host for overexpression of *Int* and its derivatives.

**Protein expression and purification**

Int, hInt and Int variants including the 6× histidine tagged CTD fragment (hCTD), were purified as described previously (8,14). Western blots were performed on proteins taken from three stages of the expression and purification procedure of hInt mutants i.e. whole cell extracts, the crude cell lysates and a fraction following Ni2+ affinity chromatography. Aliquots were then loaded onto a 4–12% PAGE gel, blotted onto nylon membranes and probed with a rabbit polyclonal antibody to Int followed by a secondary horse raddish peroxidise (HRP) conjugate goat anti-rabbit Ig antibody or an anti-His tag antibody-HRP conjugate (Qiagen). Detection was performed using an enhanced chemiluminescent (ECL) substrate (Pierce). A Superose-6 and a Superdex S-200 column were used for analytical size exclusion chromatography (SEC) on an AKTA FPLC (GE Healthcare) pre-equilibrated with GF buffer (20 mM Tris-HCl pH 8.0, 500 mM NaCl, 10 mM β-mercaptoethanol). The apparent molecular weights were determined by comparison of peaks obtained at A280 with those obtained with molecular weight standards (Bio-Rad).

**Recombination assays**

*In vivo* recombination assays were performed by the introduction of a plasmid expressing int into *E. coli* DH5α (pARM101) and selection on agar containing IPTG, X-gal, apramycin and kanamycin (9,14). A plasmid expressing wild-type Int gives 100% white colonies in this assay. Subculture from blue colonies on the same medium sequenced for validation. Residues are numbered according to their position in wild-type Int (1–605) as expressed by pH562 (13).

Plasmid pARM101 was used as a reporter for the *in vivo* activity assays and was constructed by exchanging the kanamycin resistance gene from pRT504 (14) with an apramycin resistance gene by recombination in *E. coli* BW25113/pIJ790 (18). The DNA-encoding apramycin resistance was amplified by PCR using primers MS400 and MS401 (Supplementary Table S1) from the REDIRECT cassette encoded by pJ773. *Escherichia coli* BL21 F−ompT gal dcm lon hsdS8(F− mB−), λ (DE3), pLysS was used as a host for overexpression of *Int* and its derivatives.
was used to detect any white colonies that might segregate and would thus be indicative of very low levels of Int activity.

*In vitro* recombination assays were performed as described previously except that modifications were made to buffers (19). The effect of EDTA upon Int activity was assayed by supplementing RxE buffer [10 mM Tris–PO₄, 100 mM NaCl, 5 mM DTT, 5 mM spermidine, 4.5% glycerol and 0.5 mg/ml bovine serum albumin (BSA)] with EDTA as indicated. Int (final concentration 1 μM) was added to the modified recombination buffer and pre-incubated (30°C, 1 h) prior to the addition of pRT702 and pRT600 containing *attP* and *attB*, respectively (final concentrations ~3 nM). Reactions were further incubated (30°C, 2 h) and the reaction products were then cut with HindIII and separated by electrophoresis (0.8% agarose/TBE gel). *In vitro* assays to measure the protection against EDTA by *att* sites were performed by pre-incubation (1 h, 30°C) of Int in RS buffer (10 mM Tris–PO₄, 100 mM Na₂SO₄, 5 mM DTT, 5 mM spermidine, 4.5% glycerol and 0.5 mg/ml BSA) with or without 2 mM EDTA and with or without linear 50-mer *att* sites, made from annealed DNA oligonucleotides, at a 2:1 molar ratio of Int monomer: 50-mer *att* site (400 nM Int and 200 nM *att* DNA). Recombination was then assayed as above by addition of a partner *att* site located on a supercoiled plasmid and in reactions that did not contain an *att* site during pre-incubation, a 50-mer *att* site. Reaction mixtures were digested with SacI and separated by agarose gel electrophoresis to detect recombinant products.

*In vitro* recombination was also assayed using the reporter plasmid pRT508 that contains *attP* and *attB* flanking lacZ (9). Approximately 0.15 nM pRT508 was used in an *in vitro* recombination assay; the reaction mixture was introduced into DH5α and with the proportion of blue versus white colonies growing on agar containing ampicillin, IPTG and XGal indicated the recombination activity.

*In vitro* DNA binding assays

*In vitro* DNA binding affinities were determined as described previously using radiolabelled annealed oligonucleotides as probes in electrophoretic mobility shift assays (EMSAs) (8). For standard affinity assays 1.0 ng labelled probe in binding buffer (20 mM Tris–HCl, pH 8.0, 50 mM KCl, 5% glycerol, 79 870 M⁻¹/C0₁ cm, 1 h) prior to the addition of pRT702 and pRT600 containing *attP* and *attB*, respectively (final concentrations ~3 nM). Reactions were further incubated (30°C, 2 h) and the complexes separated and detected as for the standard assay.

**ICP–MS**

Protein-bound sulphur and zinc were assayed by coupling SEC to ICP–MS. Sample (5–40 μl) of ~18 μM Int was injected onto a HiTrap SM 5 ml desalting column (SEC) and eluted using buffer [10 mM Tris–HCl (normalpur, VWR), 50 mM KCl (analR, BDH) pH 7.40] and a flow rate of 0.7 ml/min. The mobile phase pumped through the SEC column by a high performance liquid chromatography (HPLC) system (Accela, Thermo Scientific) was coupled to the Meinhard-nebuliser of an Element-II ICP–MS (Thermo Scientific) via a T-piece which allowed the introduction of 74Ge as internal standard. Data were recorded at medium resolution with a dwell time per isotope ratio of 10 ms/20 peaks (250 ms total/isotope). No evidence of interference between the 60Zn and 32S²⁺ signals was observed. To determine sulphur to zinc ratios 33S, 64Zn isotopes were measured alongside 74Ge (as internal standard, BDH). For calibration stock solutions of 100 mM methionine (Sigma, 98%), 10 mM ZnSO₄ (1000 p.p.m., BDH) and 10 mM EDTA (normalpur, VWR) were prepared (and further dilutions as required) as standards. To verify sensitivity and accuracy of the Zn/S ratios carbonic anhydrase from bovine erythrocytes (2500 U/mg protein, Sigma) and superoxide dismutase from bovine erythrocytes (7000 U/mg protein Sigma) were used. Quantification of isotope was determined via integrated peak area.

**Circular dichroism spectroscopy**

Circular dichroism (CD) Spectra of the protein solutions were obtained using a Jasco J-810 spectropolarimeter. Far UV CD measurements (185–260 nm) were collected in quartz cells of 0.02 cm path length using the following parameters: temp 25°C; scan speed 50 nm/min; bandwidth 1.0 nm; response 0.5 s and data pitch 0.2 nm. Int and derivatives were exchanged into a buffer compatible with CD measurements [25 mM Tris–PO₄ (pH 7.5), 200 mM Na₂SO₄] with a desalting column (5 ml HiTrap, GE Healthcare). Chloride ions present in the normal purification/storage buffer absorb strongly below 200 nm which would have affected the reliability of the data. The activity of the protein in RS buffer containing sodium sulphate instead of sodium chloride was tested and showed no loss of activity.

Protein concentrations were determined using a spectrophotometer at 280 nm using an estimated extinction coefficient, ε₂₈₀, for Int, hist and hCTD of 78 380, 85 830 and 79 870 M⁻¹/cm⁻¹, respectively. In all measurements, the CD spectrum of the buffer was subtracted from each protein spectrum. To obtain final CD spectra, five CD scans were accumulated and averaged for each sample. The spectra were corrected for protein concentration and cell path length before being analysed by DICHROWEB (20,21) an online server which hosts the various algorithms used to estimate protein secondary structures. Near UV CD measurements (250–320 nm) were obtained in a 0.5 cm quartz cuvette using the
following parameters: temp 25°C; scan speed 10 nm/min; bandwidth 1.0 nm; response 2 s and data pitch 0.2 nm.

RESULTS

The CTDs of the large serine recombinases contain a conserved motif that resembles a zinc finger (Figure 1), but does not conform to known classes (16). Four conserved cysteines present in most large serine recombinases could coordinate a zinc atom and, if so, this motif is likely to be a new class of zinc finger.

φC31 Int contains zinc

To determine whether zinc is present, Int was analysed by HPLC–ICP–MS. Concomitant analysis of sulphur and zinc showed that the Int containing fraction eluting from the HPLC column contained zinc (Figure 2). No other metals were detected by ICP–MS in Int. Known amounts of the S and Zn were used as standards and peak integration was used to obtain a ratio of sulphur to zinc. At an injection volume of 40 μl a reproducible ratio of 15:1 S:Zn was obtained or just under one atom of zinc bound to one monomer of Int (Table 1). The discrepancy from the expected ratio of 17:1 may be explained by zinc attrition in the column or the result of contamination with sulphur. Zinc was also present in the isolated large CTD of Int, hCTD, containing amino acid 155–605 fused to an N-terminal histidine tag (8) (Table 1), but was undetectable in a truncated Int, hIntV371SGA containing a stop codon within the zinc finger-like motif (data not shown). These data localize the zinc binding motif to within the CTD.

Zinc binding by Int is specific and essential for recombination activity

To assess the biological importance of metal binding by the zinc finger-like motif, Int was treated with metal chelators or a cysteine modifier and used in an activity assay. Int was pre-incubated with increasing concentrations of EDTA prior to the addition of substrates attP and attB and further incubation to allow recombination to occur. Pre-incubation with EDTA had an inhibitory effect on the integration reaction, even at the lowest EDTA concentration (Figure 3, top panel, lanes 1–6). Int activity was sometimes reduced after pre-incubation, even in the absence of added chelator (e.g. Figure 3, top panel, lanes 1 and 7). It was possible, however, to completely reconstitute Int activity by addition of ZnSO₄ (final concentration 0.5 mM) at all concentrations of EDTA tested, including the control sample that contained no EDTA (Figure 3, middle panel, lanes 13–18). This restoration of full activity suggests that Int can lose zinc, either by leaching in the absence of EDTA or due to the chelation of zinc by EDTA (Note that addition of zinc to Int without pre-incubation had no effect on activity). Addition of exogenous zinc after treatment with chelator restored Int activity, suggesting that zinc is rapidly rebound by Int and is sufficiently stable, even when the chelator is in excess, to drive recombination.

The ability of other cations (Mn²⁺, Cu²⁺, Co²⁺, Mg²⁺, Ca²⁺, Fe²⁺, Ni²⁺) to restore activity was tested but only Ca²⁺ and Mg²⁺ gave any observable recovery of activity after EDTA treatment (Figure 3). ICP–MS tests on all the salt solutions showed that they all contained small amounts of zinc that in the case of the calcium and magnesium salts may have restored some Int activity without poisoning. Overall the data show that Int specifically requires zinc for activity. In a time course of inhibition of Int activity by EDTA a 20 min pre-incubation was sufficient to abolish activity (Supplementary Figure S1). Despite several attempts we were not able to purify the apo-Int after treatment with EDTA. EDTA treated Int was used in SEC–ICP–MS analysis and although the signal was low due to poor protein solubility, no zinc was detected in the fractions containing Int (data not shown). The cysteine modifying compound iodoacetate also strongly inhibited the activity of Int, consistent with cysteines having an important role in recombination activity (Supplementary Figure S2). Taken together these data indicate that Int monomers coordinate a zinc atom, probably via the conserved cysteines, and that zinc is essential for recombination activity.

Zinc is required for high-affinity DNA binding by Int

A possible role for the zinc finger-like motif in Int is in substrate recognition and DNA binding. We, therefore,

Figure 2. Quantitation of zinc (light grey) and sulphur (dark grey) in φC31 Int eluting from SEC and elements detected simultaneously by ICP-MS.

Table 1. ICP-MS analysis of sulphur and zinc in Int and hCTD

| Injection volume (μl) | Sulphur (μmol/ml) | Zinc (μmol/ml) | S/Zn |
|----------------------|------------------|----------------|------|
| Integrase            |                  |                |      |
| 5                    | 0.0666           | 0.00461        | 14.5 |
| 10                   | 0.0645           | 0.00426        | 15.1 |
| 20                   | 0.108            | 0.00646        | 16.6 |
| 40                   | 0.121            | 0.00791        | 15.3 |
| Mean (SD)            |                  |                | 15.4 (0.9) |
| hCTD                 |                  |                |      |
| 5                    | 0.00237          | 0.000157       | 15.3 |
| 10                   | 0.00381          | 0.000236       | 15.1 |
| 20                   | 0.00515          | 0.000315       | 16.6 |
| Mean (SD)            |                  |                | 15.9 (0.5) |
tested the effect of EDTA on Int DNA binding activity in EMSA experiments (Figure 4A). Pre-incubation of Int in the presence of 2 mM EDTA led to greatly reduced affinities (50% binding of substrates was ≥1333 nM) for all four attachment sites. Affinities could be partly restored on addition of 0.5 mM ZnSO₄ (50% binding of probe was between 21 and 83 nM compared to 5 to 21 nM for the non-EDTA treated, non-pre-incubated Int control). Pre-incubation in the absence of EDTA also partly inhibited Int binding activity but this too could be restored by addition of ZnSO₄ (Figure 4B). Other metal ions were tested for their ability to restore DNA binding after pre-incubation and but only zinc ions could restore full activity to Int pre-incubated with EDTA (Figure 4B). The EDTA sensitive binding activity of Int is therefore dependent specifically on zinc ions. These data indicate that EDTA inhibits DNA binding by Int, presumably through chelating zinc that is lost from Int during pre-incubation. However, even at EDTA concentrations that completely inhibit Int recombination activity, Int can still bind the attachment sites although with greatly reduced affinities (Figure 4A). Binding by Int to its substrates is, therefore, comprised of EDTA resistant and EDTA sensitive binding. The EDTA sensitivity implies that high-affinity DNA binding requires zinc.

Cysteines and other conserved residues in the zinc finger-like motif are required for protein stability

Four conserved cysteines, C366, C368, C387 and C405, could coordinate the zinc atom and substitution of these residues might enable purification of an apo-protein (Figure 1). These residues, and two other conserved amino acids, Y385 and M372, were substituted in hInt, an N-terminally His-tagged derivative of Int. All of the mutated proteins hIntC366A, hIntC368A, hIntM372A, hIntM372I, hIntY385F, hIntC387A, hIntC405A and a double mutant, hIntC366A, C368A, gave rise to the same breakdown products (~44 and ~22 kDa) after expression in E. coli (Supplementary Figure S3 and data not shown). Western blot analysis and peptide mass fingerprinting confirmed that the ~44 kDa band was an hInt fragment containing the His-tag and the NTD but truncated at the zinc finger-like motif (data not shown). Coordination of zinc during expression is, therefore, vital for correct folding of the protein and without this the unfolded protein is liable to attack from E. coli proteases. Purified wild-type Int treated with EDTA does not breakdown indicating that the protein is stable in the absence of proteases (data not shown). These data indicate the importance of these residues to the stable folding of this motif. Unsurprisingly these mutated proteins showed no recombination activity in an in vivo recombination assay (data not shown).

Substitutions in the zinc finger-like motif affect DNA binding affinities

Amino acid substitutions were generated between S374 and I381 (Table 2). These residues were chosen as they are in a putative loop between two zinc coordinating cysteines. In addition they flank a partially conserved glycine, a residue that allows for sharp turns in the peptide backbone (Figure 1). The polar and charged amino acids could potentially make contacts with the DNA. The mutated proteins were purified, tested for in vitro recombination activity and DNA binding activity (Table 2 and Figure 5). While almost all the mutated proteins had changes in affinity compared with wild-type Int, the most severely affected proteins for integration activity were those with substitutions in four consecutive residues S374 to G377. Two types of DNA binding affinity defects were observed with this set of Int variants; those that had reduced affinities for all the att sites (hIntS374L, hIntK375A, hIntR376E, hIntG377F, hIntE378A, hIntE379A and hIntS380A) and those whose affinities to attB were affected more than to other att sites (for e.g. hIntK375Q, hIntK375E, hIntR376A and hIntG377K; Table 2 and Figure 5). In some proteins (hIntK375E, hIntR376A, hIntG377K), attL binding also appeared to be more affected that attP or attR binding. attB and attL both contain a B half site that has been shown previously to have a greater contribution than the B’ half site to the activity of the attB site (15). One substitution, hIntR376E,
abolished nearly all DNA binding (Table 2 and Figure 5). Thus an intact zinc finger-like motif is required for high-affinity DNA binding and may have a role in distinguishing between an attB and an attP site.

CD spectroscopy and SEC to evaluate secondary structure of binding defective mutants

The DNA binding defects in the Int variants could be due to loss of direct contacts between protein and DNA or due to gross structural changes that indirectly perturb DNA binding affinity. Int variants with some of the more severe binding defects were, therefore, analysed by CD and SEC to try to detect structural changes that might be correlated with poor DNA binding (Figure 6; Supplementary Tables S1 and S2).

The near UV CD spectra showed that the environments of the aromatic residues in wild-type Int and most of the variant proteins were essentially the same and strongly implied that there were no gross structural changes in the variant proteins (Figure 6A). The proteins with the most deviant fingerprints in the near UV region were hCTD (expected as this protein has lost the entire NTD).
Table 2. Integration activities and DNA binding affinities of Int mutants

| Plasmid   | Int variant | In vivo activity: % white/+ or – in subculture | In vitro activity: transformation assay (% white) | Int concentrations (nM) for 50% binding of probe |
|-----------|-------------|------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
| pHS62     | Int         | 100/+                                          | 91                                              | 21–83                                           |
| pARM010   | hInt        | 100/+                                          | 87                                              | 21–83                                           |
| pADD002   | hIntS374A   | 42/+                                           | 78                                              | 21–83, 1.3–5                                    |
| pARM024   | hIntS374L   | 0/–                                            | 2                                               | >1333                                           |
| pARM070   | hIntK375A   | 2/+                                            | 20                                              | 1333, 83, 21–83                                 |
| pARM103   | hIntK375Q   | 0/–                                            | <1                                              | 83–333, 83                                     |
| pARM104   | hIntK375E   | 0/–                                            | <1                                              | 83–333, 83                                     |
| pARM071   | hIntR376A   | 1/+                                            | <1                                              | 83–333, 83                                     |
| pARM105   | hIntR376E   | 0/–                                            | <1                                              | >1333, >1333                                   |
| pARM096   | hIntR376P   | 0/–                                            | <1                                              | >1333, 333                                     |
| pADD001   | hIntG377L   | 61/+                                           | 78                                              | 333                                             |
| pADD002   | hIntG377A   | 100/+                                          | 84                                              | 83–333, 83                                     |
| pARM020   | hIntG377D   | 40/+                                           | 58                                              | 83–333, 83                                     |
| pARM021   | hIntG377F   | 4/+                                            | <1                                              | 83–333, 83                                     |
| pARM022   | hIntG377K   | 0/–                                            | 1                                               | 333                                             |
| pARM072   | hIntE379A   | 100/+                                          | 94                                              | 83–333, 83                                     |
| pARM073   | hIntE379A   | 100/+                                          | 93                                              | 83–333, 83                                     |
| pARM074   | hIntS380A   | 100/+                                          | 95                                              | 83–333, 83                                     |
| pARM107   | hIntS380D   | 32/+                                           | 29                                              | 333                                             |
| pARM108   | hIntS380N   | 12/+                                           | 70                                              | 83–333, 83                                     |
| pARM109   | hIntS380T   | 100/+                                          | 94                                              | 83–333, 83                                     |
| pARM100   | hIntE381M   | 100/+                                          | 95                                              | 83–333, 83                                     |
| pARM106   | hIntV371SUGA| 0/–                                            | <1                                              | >1333, 333                                     |

Bold indicates a lower affinity for a substrate than the wild-type Int. ‘ND’ indicates that the data were not determined.

and hIntK375Q, an Int variant that is recombinationally inactive and defective in attB binding but able to bind attP, attL and attR with near normal affinities (Table 2). Most of the Int mutants showed a slight increase in the intensity of their ellipticity values in the far UV region which implies (assuming accurate protein estimations) that the amino acid substitutions in all these mutants have all undergone small, similar changes to overall secondary structural conformation, such as stiffening or removal of structural constraints that affect the tertiary conformation (Supplementary Table S2). The major outliers in the far UV were hCTD and hIntS380D, a partially active protein with very high binding affinities to all the attachment sites (Figure 6B, Table 2). Overall the differences between the secondary structures of the Int mutants compared with hInt or Int were small. In particular, there was little change in the estimated proportions of turn or disorder although the mutant proteins appeared reproducibly to have marginally more predicted alpha helix compared with beta-sheet (Supplementary Table S2). Crucially none of the observed differences in CD spectra could be correlated with changes in DNA binding affinities or recombination activities.

Analysis of the Int variants by SEC indicated no changes in oligomeric structures; all the Int proteins with amino acid substitutions behaved as dimers in solution (Supplementary Table S3).

A truncated Int lacking an intact zinc finger-like motif has reduced binding affinities

A by product of the site-directed mutagenesis experiments was an int allele encoding a V371S substitution followed by a stop codon (hIntV371SUGA). This protein, truncated within the zinc finger-like motif, binds the att sites with much lower affinities than hInt but still generates the two complexes (I and II) typically observed when Int binds its substrates (Figure 5B). The residual binding activity by hIntV371SUGA shows that there is another, as yet uncharacterized, DNA recognition and binding motif in Int. Binding to the substrates by hIntV371SUGA resembled binding by EDTA-treated Int as Int–DNA complexes were only observed at the highest Int concentrations. Thus an intact zinc finger-like motif is required for high-affinity DNA binding.

Inhibition of Int activity by EDTA can be prevented by the presence of attachment sites

If the zinc finger is indeed a DNA recognition and binding motif and comes into close contact with the attachment sites, we hypothesized that Int in complex with the att sites should be less sensitive to EDTA than Int in free solution. Annealed oligonucleotides containing the sequences of attP or attB were used to attain stoichiometric ratios of att site 50-mer to a dimer of Int. Int was pre-incubated in the presence or absence of the 50-mer att site and in the presence or absence of EDTA for 1 h. Recombination activity was then assayed by addition of a partner att site and in the presence of attachment sites encoded on a supercoiled plasmid and, in those reactions that did not contain an att site at the start of pre-incubation, a 50-mer att site (Figure 7). Recombination yielded linearized plasmid which could be further digested to give restriction fragments diagnostic of recombination products. As before, pre-incubation of Int with EDTA inhibited attP × attB recombination and
activity could be mostly restored by addition of ZnSO₄. However, if the attP or attB 50-mers were added at the start of pre-incubation with EDTA, no inhibition of recombination activity was observed suggesting that these substrates could protect Int from the zinc chelating effects of EDTA (Figure 7).

No protection was observed when a non-specific oligonucleotide or if poly dI-dC were added to the pre-incubation mix (data not shown). These data show that Int specifically bound to its attachments sites is protected from the inhibitory activity of EDTA suggesting that the zinc finger-like motif is in direct contact with the DNA.

DISCUSSION

Zinc fingers are a large family of small protein domains with diverse sequence, size and range of functions. Mostly zinc fingers are structural domains involved in interactions with other molecules, particularly nucleic acids and proteins (16,22). Zinc binding can often be predicted from sequence as typical zinc binding is through interactions with cysteine, histidine, glutamic acid or aspartic acid side chains, but most frequently through tetra-coordination via cysteines (16). Here we present the evidence for a putative zinc finger in φC31 Int. An Int monomer contains a Zn²⁺ atom and both recombinase and DNA binding activities of Int were sensitive to EDTA. Mutant Ints containing substitutions in the cysteines that are thought to be coordinating the Zn²⁺ atom are unstable indicating that this motif requires zinc for structural integrity.

Evidence is also presented that the putative zinc finger is directly involved in recognition and binding to the attachment sites. Amino acid substitutions can have dramatic effects on DNA binding affinity by zinc fingers (23,24). Similarly amino acid substitutions in the zinc finger-like motif in φC31 Int also reduced the DNA binding affinities to variable extents, the most severe of which could be due to an ionic clash (hIntR376E) and appears to abolish DNA binding to a greater extent than either treatment of Int with EDTA or truncation of Int at V371 (hIntV371UGA). Mutant Ints with lesser reductions in affinity could be due to loss of contacts (substitutions with alanine) or steric blocks (e.g. G377K, G377F; Table 2) (25,26). We showed that the att sites could protect Int against inhibition by EDTA; this is probably because the DNA and the zinc finger-like motif are in close proximity and the DNA prevents EDTA from accessing the Zn²⁺. High-affinity DNA binding by Int is EDTA sensitive but that there is also an EDTA resistant, low affinity activity. Indeed the truncated protein, hIntV371UGA, indicates that Int has at least one other DNA binding region. In Bxb1 gpInt a similar construct to φC31 hIntV371UGA also binds DNA with low affinity and substitutions in both Bxb1 gpInt and TnpX in regions equivalent to the truncated CTD in hIntV371UGA affect DNA binding (7,27). Despite this strong evidence for direct contact by the zinc finger-like motif to DNA, we cannot yet rule out an indirect effect in which this motif influences either this second DNA binding region or an as yet unidentified DNA binding motif elsewhere in the protein.
In the absence of Xis Int forms a synaptic complex only when attP and attB are substrates indicating that binding to these substrates evokes different Int conformations as a result of different protein:DNA recognition events (1,7,10). A fundamental question to address given that attP and attB are such different sequences is whether Int has two DNA binding regions; one for P type arms and one for B arms. This seems unlikely given that mutants described here can be generally reduced in all att site binding. However, many of the Int variants were more affected in attB than to attL/R and attP, indicating that the interaction with attB, and possibly B type arms, might involve a different set of contacts to those made with attP. Moreover recombination activity is more likely to be lost by amino acid substitutions in S374 to G377 than in E378 to I381, although DNA binding can be affected by substitutions throughout the zinc finger-like motif (Table 2, Figure 5). Thus contacts made to the attB and attP sites via S374, K375 and R376 could be critical for inducing conformation changes required for activation of recombination. In our experiments we noted a distinct reduction in the amount of complex I formed in the EMSA experiments with hIntS374L, hIntR376A and hIntG377K compared with wild-type Int. Int binds to all the substrates to generate two complexes (I and II) that we have previously interpreted as containing either a monomer or a dimer of Int, respectively (10). Possibly the dimer complex is more stable in some of the Int variants than in the wild-type Int complexes and that destabilizing the dimer interface may therefore be one of the changes that occurs to generate the active synaptic tetramer.

In this article, we have demonstrated for the first time the importance of zinc in a phage-encoded Int that is likely a part of a DNA binding motif. The serine Ints are widely used as tools for genome engineering and gene therapy, [for example, see Ref. (28)]. The mechanism through which these recombinases recognize their DNA substrates

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**Figure 6.** CD of φC31 Int and derivatives. Superimposed spectra of all the protein analysed are shown (panel A; near UV, panel B; far UV). The outliers in the near UV are hCTD (dotted line) and the wild-type Int (dashed line). All the remaining spectra in the near UV superimpose with the hInt protein. In the far UV the outliers are hCTD (crosses) and hIntS380D (circles); hInt and Int are the dashed and dotted lines, respectively. In the far UV spectra, all the hInt derivatives with an amino acid substitution therefore have slightly enhanced ellipticity compared with the hInt.
and activate recombination is of interest for rational engineering and improvement of recombination efficiency in heterologous environments.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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REFERENCES
1. Smith, M.C.M., Brown, W.R., McEwan, A.R. and Rowley, P.A. (2010) Site-specific recombination by φC31 integrase and other large serine recombinases. Biochem. Soc. Trans., 38, 388–394.
2. Grindley, N.D.F., Whiteson, K.L. and Rice, P.A. (2006) Mechanisms of site-specific recombination. Annu. Rev. Biochem., 75, 567–605.
3. Smith, M.C.M. and Thorpe, H.M. (2002) Diversity in the serine recombinases. Mol. Microbiol., 44, 299–307.
4. Yuan, P., Gupta, K. and Van Duyne, G.D. (2008) Tetrameric structure of a serine integrate catalytic domain. Structure, 16, 1275–1286.
5. Sanderson, M.R., Freemont, P.S., Rice, P.A., Goldman, A., Hatfull, G.F., Grindley, N.D.F. and Steitz, T.A. (1990) The crystal structure of the catalytic domain of the site-specific recombination enzyme gamma delta resolvase at 2.7 Å resolution. Cell, 63, 1323–1329.
6. Ghosh, P., Kim, A.I. and Hatfull, G.F. (2003) The orientation of mycobacteriophage Bxb1 integration is solely dependent on the central dinucleotide of attP and attB. Mol. Cell, 12, 1101–1111.
7. Ghosh, P., Pannunzio, N.R. and Hatfull, G.F. (2005) Synapsis in phage Bxb1 integration: selection mechanism for the correct pair of recombination sites. J. Mol. Biol., 349, 331–348.
8. McEwan, A.R., Rowley, P.A. and Smith, M.C.M. (2009) DNA binding and synapsis by the large C-terminal domain of φC31 integrase. Nucleic Acids Res., 37, 4764–4773.
9. Rowley, P.A., Smith, M.C.A., Younger, E. and Smith, M.C.M. (2008) A motif in the C-terminal domain of φC31 integrase controls the directionality of recombination. Nucleic Acids Res., 36, 3879–3891.
10. Thorpe, H.M., Wilson, S.E. and Smith, M.C.M. (2000) Control of directionality in the site-specific recombination system of the streptomyces phage φC31. Mol. Microbiol., 38, 232–241.
11. Kuhlbrandt, W. and Iqbal, K. (1991) In vitro recombination of the streptomyces bacteriophage φC31. J. Mol. Biol., 222, 897–908.
12. Combes, P., Till, R., Bee, S. and Smith, M.C.M. (2002) The streptomyces genome contains multiple pseudo-attB sites for the φC31-encoded site-specific recombination system. J. Bacteriol., 184, 5746–5752.
13. Thorpe, H.M. and Smith, M.C.M. (1998) In vitro site-specific integration of bacteriophage DNA catalyzed by a recombinase of the resolvase/invertase family. Proc. Natl Acad. Sci. USA, 95, 5505–5510.
14. Smith, M.C.A., Till, R., Brady, K., Soultanas, P., Thorpe, H. and Smith, M.C.M. (2004) Synapsis and DNA cleavage in φC31 integrase-mediated site-specific recombination. Nucleic Acids Res., 32, 2607–2617.
15. Gupta, M., Till, R. and Smith, M.C.M. (2007) Sequences in attB that affect the ability of φC31 integrase to synapse and to activate DNA cleavage. Nucleic Acids Res., 35, 3407–3419.
16. Krishna, S.S., Majumdar, I. and Grishin, N.V. (2003) Structural classification of zinc fingers: survey and summary. Nucleic Acids Res., 31, 532–550.
17. Liu, H. and Naismith, J.H. (2009) A simple and efficient expression and purification system using two newly constructed vectors. Protein expression and purification, 63, 102–111.
18. Gust, B., Kieser, T. and Chater, K.F. (2002) REDIRECT Technology: PCR-targeting System in Streptomyces coelicolor. John Innes Centre, Norwich.
19. Smith, M.C.A., Till, R. and Smith, M.C.M. (2004) Switching the polarity of a bacteriophage integration system. Mol. Microbiol., 51, 1719–1728.
20. Whitmore, L. and Wallace, B.A. (2004) DICHROWEB, an online server for protein secondary structure analysis from circular dichroism spectroscopic data. Nucleic Acids Res., 32, W668–W673.
21. Whitmore, L. and Wallace, B.A. (2008) Protein secondary structure analyses from circular dichroism spectroscopy: methods and reference databases. Biopolymers, 89, 392–400.
22. Gamsjaeger, R., Liew, C.K., Loughlin, F.E., Crossley, M. and Mackay, J.P. (2007) Sticky fingers: zinc-fingers as protein-recognition motifs. TIBS, 32, 63–70.
23. Nardelli, J., Gibson, T. and Charnay, P. (1992) Zinc finger-DNA recognition: analysis of base specificity by site-directed mutagenesis. *Nucleic Acids Res.*, **20**, 4137–4144.

24. Clouaire, T., Roussigne, M., Ecochard, V., Mathe, C., Amalric, F. and Girard, J.P. (2005) The THAP domain of THAP1 is a large C2CH module with zinc-dependent sequence-specific DNA-binding activity. *Proc. Natl Acad. Sci. USA*, **102**, 6907–6912.

25. Luscombe, N.M., Laskowski, R.A. and Thornton, J.M. (2001) Amino acid-base interactions: a three-dimensional analysis of protein-DNA interactions at an atomic level. *Nucleic Acids Res.*, **29**, 2860–2874.

26. Rohs, R., Jin, X., West, S.M., Joshi, R., Honig, B. and Mann, R.S. (2010) Origins of specificity in protein-DNA recognition. *Ann. Rev. Biochem.*, **79**, 233–269.

27. Adams, V., Lucet, I.S., Tynan, F.E., Chiarezza, M., Howarth, P.M., Kim, J., Rossjohn, J., Lyra, D. and Rood, J.I. (2006) Two distinct regions of the large serine recombinase Tnpx are required for DNA binding and biological function. *Mol. Microbiol.*, **60**, 591–601.

28. Calos, M.P. (2006) The φC31 integrase system for gene therapy. *Curr. Gene Therapy*, **6**, 633–645.

29. Poirot, O., O’Toole, E. and Notredame, C. (2003) Tcoffee@igs: a web server for computing, evaluating and combining multiple sequence alignments. *Nucleic Acids Res.*, **31**, 3503–3506.