A motif in the C-terminal domain of φC31 integrase controls the directionality of recombination

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ABSTRACT

Bacteriophage φC31 encodes an integrase, which acts on the phage and host attachment sites, attP and attB, to form an integrated prophage flanked by attL and attR. In the absence of accessory factors, φC31 integrase cannot catalyse attL x attR recombination to excise the prophage. To understand the mechanism of directionality, mutant integrases were characterized that were active in excision. A hyperactive integrase, Int E449K, gained by attB x attR recombination whilst retaining the ability to recombine attP x attB. A catalytically defective derivative of this mutant, Int S12A, E449K, could form stable complexes with attP/attB, attL/attR, attL/attL and attR/attR under conditions where Int S12A only complexed with attP/attB. Further analysis of the Int E449K-attL/attR synaptic events revealed a preference for one of the two predicted synapse structures with different orientations of the attL/attR sites. Several amino acid substitutions conferring hyperactivity, including E449K, were localized to one face of a predicted coiled-coil motif in the C-terminal domain. This work shows that a motif in the C-terminal domain of φC31 integrase controls the formation of the synaptic interface in both integration and excision, possibly through a direct role in protein–protein interactions.

INTRODUCTION

In order to establish a lysogenic life style, many bacteriophages encode an integrase to integrate the phage genome into the host chromosome (1,2). In association with an accessory factor, the recombination directionality factor (RDF) or Xis, the integrase also excises the phage genome on induction into the lytic pathway (2,3). During integration, the phage integrases act site-specifically by recombining the host and phage attachment sites, attB and attP, to form the hybrid products attL and attR. In excision attL and attR recombine to regenerate attB and attP. Phage integration and excision has to be a highly regulated process to ensure efficient switching between the lytic pathway and lysogeny (1). As some integrases are unidirectional in the absence of accessory factors, phage integrases can be used for site-specific genome manipulations and for the stable integration of transgenes (4). The integrase from the Streptomyces temperate phage φC31 is currently the most widely used phage integrase for genome manipulations (5–14).

The φC31 integrase is a member of a large family of site-specific recombinases, the serine recombinases (15,16). The best understood members of this family are the resolvase/invertases, in particular γδ and Tn3 resolvases and Hin invertase. The resolvase/invertases have an N-terminal catalytic domain (~1–140 aa) and a small (~141–183 aa), C-terminal DNA-binding domain. The φC31 integrase is a member of the large serine recombinases (17). These proteins have similar N-terminal catalytic domains to those of the resolvase/invertases, but they have much larger C-terminal domains of ~300–500 aa. The large serine recombinases also include some transposases such as TnpX from the Clostridium perfringens transposon, Tn4451, and TndX from the C. difficile conjugative transposon CTn5397 (18–20). Tn4451 and CTn5397 confer resistances to chloramphenicol and tetracycline, respectively. TnpX and TndX have been shown to be capable of integrating and excising their respective transposons with no requirement for any additional transposon-encoded accessory factors (20,21).

The first step in a site-specific recombination pathway is recognition and binding of the substrates by the recombinase (22). Protein–protein interactions between the recombinase subunits then bring the two substrates together in a synaptic complex. Ghosh et al. (23) showed that dimers of mycobacteriophage Bxb1 integrase (a large serine recombinase) bind the substrates attB and attP, which are then probably brought together to form a synaptic tetramer. Post-synapsis, the resolvase/invertases and the large serine recombinases appear to use the same
catalytic mechanism (16,22–27). DNA cleavage generates 2 bp staggered breaks in both substrates with the concomitant formation of transient covalent phosphoserine bonds between the recombinases and the recessed S' ends. Strand exchange is thought to occur by 180° rotation of two recombinase subunits bound to half sites relative to the other two subunits (28–30). Joining of the half sites to form recombinant products is inhibited when the two base overhangs at the staggered breaks cannot base pair due to mismatches (31,32). In the serine integrases, the non-palindromic nature of the dinucleotides at the crossover site are critical in determining the left–right polarity of the sites (24,26).

In the absence of any accessory factors φC31, Bxb1 and φRv1 integrases only catalyse the integration reaction attP × attB to form attL and attR (33–35). In binding assays, synaptic complexes can only be detected when integrase is provided with attP and attB and not when provided with attL and attR or any other pairwise combination of att sites (23,25). These observations have led to the idea that integrase has specific conformations when bound to attP and attB that enable the protein–protein interactions required for synopsis and subsequent recombination (23,25,27). A mutation analysis of the φC31 attB site showed that there are indeed sequences in attB that appear to be required for stable synopsis (36). An RDF (gp47) has been reported from Bxb1 that is essential for excisive recombination by Bxb1 integrase, (gpInt) (37). In this system, correct excision is dependent on gp47 to construct a synaptic complex containing gpInt and the DNA sites in a specific alignment that enables attL x attR recombination and prevents aberrant DNA rearrangements (38).

The role of the large C-terminal domain in the recombination mechanism by the phage serine integrases is poorly understood. It seems likely that the C-terminal domains are involved in DNA binding and in the control of directionality. The aim of this investigation was to understand the mechanism of directional control in φC31 integrase through the isolation and characterization of ‘hyperactive integrases’ that could recombine attL x attR sites. This work shows that a motif in the C-terminal domain of φC31 integrase controls the formation of a synaptic interface, possibly through a direct role in protein–protein interactions.

**MATERIALS AND METHODS**

**Bacterial strains, plasmids and recombinant DNA techniques**

*Escherichia coli* DH5α (F- φ80lacZΔM15, ∆(lacZYA-argF), recA1, endA1, hsdR17, phoA, supE44, thi-1, gyrA96, relA1, λ) was used as a general host for plasmid preparation, SDM and for in vivo recombination assays (39). DH5α was grown routinely in 2YT (39). Plasmid DNAs (Table 1 and Supplementary Material) were introduced by transformation or electroporation (39).

In vivo recombination assays were performed by introducing ~50 ng of an integrase expression plasmid into competent DH5α containing either the integration reporter plasmid, pRT504, or the excision reporter plasmid, pPAR1000 (Table 1), and plating the transformants on 2YT plates containing carbenicillin (50 μg/ml), kanamycin (20 μg/ml), IPTG (50 μg/ml) and Xgal (160 μg/ml). The recombination frequency was calculated as the total number of white colonies (including white colonies with blue sectors) divided by the total number of colonies.

**Mutagenesis**

A library of random mutants of the integrase expression plasmid, pHS62, was prepared in the *E. coli* mutator strain, XL1-Red (Stratagene, CA, USA). Over 200 transformants of XL1-Red (pHS62) were pooled and amplified. The extracted plasmids were screened for loss-of-function or gain-of-function phenotypes as described in the Results section.

SDM of the *int* gene was adapted from the method described in the QuikChange® SDM kit (Stratagene, CA, USA), with primers for SDM designed according to the manufacturer’s recommendations. Primers are listed in the Supplementary Material. The DNA polymerase, Pfu (Promega, WI, USA), was used for PCR with pHS62 as the template and the PCR programme set to 98°C denaturation (1 min), 55°C annealing (30s), 72°C extension (12 min) for 30 cycles. PCR reactions were digested with DpnI overnight and the products were introduced into DH5α and plated onto 2YT plates containing carbenicillin. Mutations in the *int* gene were verified by sequencing.

**Expression and purification of integrase**

*E. coli* BL21 DE3 (pLysS) was used as a host for overexpression of integrase. Integrate purifications were carried out as described previously (25). Protein concentrations were determined using the Quickstart Bradford protein reagent (BioRad, CA, USA).

| Plasmid       | att sites | Reference |
|---------------|-----------|-----------|
| pRT504       | attB<sub>273</sub>-lacZ-attP<sub>64</sub> | (25)      |
| pRT508       | attB<sub>273</sub>-lacZ-attP<sub>64</sub> | (25)      |
| pPAR1000     | attL<sub>360</sub>-lacZ-attR<sub>475</sub> | This work |
| pRT600       | attB<sub>30</sub> | (26)      |
| pRT702       | attP<sub>30</sub> | (26)      |
| pMSX18       | attL<sub>360</sub>-lacZa-attP<sub>475</sub> | This work |
| pMSX22       | attP<sub>464</sub>-lacZa-attP<sub>50</sub> | This work |
| pMSX24       | attL<sub>360</sub>-lacZa-attP<sub>50</sub> | This work |
| pMSX26       | attB<sub>30</sub>-lacZa-attL<sub>360</sub> | This work |
| pMSX28       | attB<sub>30</sub>-lacZa-attR<sub>475</sub> | This work |
| pMSX30       | attP<sub>464</sub>-lacZa-attR<sub>475</sub> | This work |
| pMSX32       | attB<sub>30</sub>-lacZa-attB<sub>51</sub> | This work |
| pMSX34       | attL<sub>360</sub>-lacZa-attL<sub>62</sub> | This work |
| pMSX36       | attR<sub>475</sub>-lacZa-attR<sub>475</sub> | This work |
| pPARX18LAA   | attL<sub>360</sub>-A<sub>4</sub> lacZa-attR<sub>475</sub> | This work |
| pPARX18RAA   | attL<sub>360</sub>-A<sub>4</sub> lacZa-attR<sub>475</sub> | This work |
| pMS92        | attL<sub>2</sub>-T<sub>4</sub> | This work |
| pMS93        | attR<sub>9</sub>-T<sub>4</sub> | This work |
| pMS94        | attL<sub>2</sub> | This work |
| pMS95        | attR<sub>9</sub> | This work |

*Numbers in superscript show the length of the att site in base pairs and the sequence of the central dinucleotides if different from wt.
DNA synapsis assays

DNA synapsis assays were performed as described previously (25,27), except that annealed oligonucleotides (Supplementary Material) were used to generate att site probes of identical lengths. Oligonucleotides were annealed (95°C, 10 min, followed by gradual cooling overnight to room temperature), purified by polyacrylamide gel electrophoresis (PAGE) using 5% polyacrylamide and 1 x TBE (Tris–borate–EDTA) buffer. After electrophoresis gels were stained with ethidium bromide and the bands containing dsDNA were excised and eluted from TE (10 mM Tris pH 8, 1 mM EDTA) using the ‘crush and soak’ method (39).

The annealed DNA fragments (~1 pmol) were labelled with α-32P-dCTP by polymerization of the recessed 3’ ends by DNA polymerase I, Klenow Fragment (New England Biolabs) (39). Excess α-32P-dCTP was removed using MicroCLEAN (Microzone Ltd, West Sussex, UK) according to the supplier’s instructions.

DNA binding was performed using 125 nM integrase and 1.5 nM of radio-labelled att site DNA in binding buffer (25). To assay synapsis, 1.5 nM of radio-labelled att site and between 14 and 32 nM of unlabelled PCR fragment encoding a second att site were incubated with 125 nM integrase in binding buffer. Samples were incubated for 2 h at 30°C and the products were separated by PAGE (5% polyacrylamide, 1 x TBE, 4°C, 5 W, 2 h). Gels were dried and exposed to Fuji phosphorimager screens. Images from the phosphorimager (Fuji FLA-3000) were analysed using AIDA software (Raytest).

In vitro recombination assays

Recombination in vitro was assayed either with unlabelled, supercoiled plasmids or with linear DNA fragments, one of which was radio-labelled. In a standard reaction using plasmid substrates containing att sites, ~100 ng of plasmid substrates (final concentration ~1.5 nM) were incubated (30°C, 2 h) in R buffer (10 mM Tris pH 7.5, 1 mM EDTA pH 8, 100 mM, NaCl, 5 mM DTT, 5 mM spermidine, 4.5% glycerol and 0.5 mg/ml bovine serum albumin) with integrase (733 nM) then heat denatured (80°C, 10 min) (25). Direct detection of recombinant molecules was by restriction analysis and agarose gel electrophoresis (25). A quantitative assay for recombination was also used which measured the deletion of lacZ by recombination between flanking att sites. In these assays, the plasmid substrate concentration was lowered to ~0.15 nM but the recombination conditions were otherwise the same as in the standard reaction. Recombinants were detected by introducing the whole reaction mixture into E. coli DH5α by transformation and selecting transformants on media containing carbenicillin, IPTG and XGal. Recombinants (white colonies) were scored amongst a background of non-recombinant (blue colonies).

Recombination of linear fragments was performed using an attP (98 bp) fragment prepared by digestion of pRT702 (HindIII/BgIII) or attL (90 bp) by digestion of pRT600702 (BglII/SphI) (26). Fragments were labelled as described for the synapse assays. Labelled probe (~1.3 nM) and a PCR-generated fragment (~10.6 nM) encoding a 235 bp partner att site and integrase (733 nM) were incubated in R buffer over a time course at 30°C. The reactions were stopped by heat denaturation (80°C, 10 min) and subtilisin (Sigma, Sigma-Aldrich Co. St Louis, MO, USA; 0.016 U, 37°C, 20 min). Subtilisin was inactivated (80°C, 10), the products were separated by PAGE (5% polyacrylamide, 1 x TBE) and analysed as described for the DNA-binding assays.

RESULTS

Integrase mutants that can recombine attL x attR

Plasmids encoding recombination defective or hyperactive integrases were identified using reporter strains of E. coli. The pRT504 encodes attP and attB flanking lacZ (25) and pPAR1000 encodes attL and attR flanking lacZa (Table 1). Transformation of E. coli DH5α (pRT504) or DH5α (pPAR1000) with a plasmid, pHS62 (33), that expresses the wild-type (wt) integrase (Int) yielded 100% white and 100% blue colonies, respectively, when grown on plates containing Xgal, IPTG and antibiotics. Thus, wt Int was fully active on the attP x attB reporter plasmid yielding recombinants that had lost the lacZ gene but was inactive on the attL x attR reporter plasmid. The pH62 was introduced into the mutagenic E. coli strain XL1-Red and plasmid DNA was prepared from the pooled colonies. When the XL1-Red derived pH62 plasmids were introduced into DH5α (pRT504), ~3% of the transformants plated on agar containing Xgal, IPTG and antibiotics were dark blue, light blue or sectored blue suggestive of a defective integrase. When the pH62-derived library was introduced into DH5α (pPAR1000), ~<1% appeared white on the same media and these were candidate gain-of-function mutants. The pH62 derivatives conferring either defective or gain-of-function recombination phenotypes were purified and the inte genes were sequenced to identify the mutations.

Re-transformation assays were used semi-quantitatively to characterize the activities of the mutant integrases. The defective mutants varied between 0% and 90% activity for attB x attP recombination (data not shown). In addition, two mutants, Int L460P and Int Y475H, demonstrated a low level of attL x attR recombination. Int E449G was isolated repeatedly from the gain-of-function screen and was 100% active in both integrative (attP x attB) and excisive (attL x attR) recombination using these in vivo assays.

In vitro recombination with hyperactive integrases

Integrase proteins were purified from E. coli strains expressing Int E449G, Int L460P and Int Y475H. These proteins were used in integration (attB x attP) and excision (attL x attR) assays using plasmid substrates in vitro (Figure 1A). Int E449G and Int L460P were active in both integration and excision, while Int Y475H, in this assay, appeared to be active only in integration.

The properties of the mutant integrases were investigated further using a more sensitive and quantitative recombination assay. Plasmid substrates were constructed, which had every combination of two att sites in head-to-tail orientation flanking lacZ or lacZa (Table 1).
These substrates were used in *in vitro* recombination reactions and, after inactivation of the integrase, the reaction products were introduced into *E. coli* DH5α by transformation. The presence of white colonies on plates containing Xgal, IPTG and carbenicillin indicated loss of transformation. The percentage of white colonies obtained for each integrase is shown. (F) Activities of Int E449K and wt Int on all possible combinations of *att* sites using the following substrates: (B); *attB* x *attP* (pRT508), (C) *attL* x *attR* (pMSX18), (D), *attL* x *attL* (pMSX34) and (E), *attR* x *attR*, (pMSX36). Substrates were incubated with integrase and then introduced into DH5α. The extreme right-hand lane contains molecular weight markers. (B–F) Quantitative *in vitro* recombination assays of Int mutants (grey bars) compared to wt Int (black bars). Recombinants were detected indirectly by excision of *lacZ* or *lacZα* flanked by two *att* sites using the following substrates: (B); *attB* x *attP* (pRT508), (C) *attL* x *attR* (pMSX18), (D), *attL* x *attL* (pMSX34) and (E), *attR* x *attR*, (pMSX36). Substrates were incubated with integrase and then introduced into DH5α by transformation. The percentage of white colonies obtained for each integrase is shown. The activities of Int E449K and wt Int on all possible combinations of *att* sites using the following substrates: P x P (*attP* x *attP*; pMSX22), B x B (*attB* x *attB*; pMSX32), P x R (*attP* x *attR*; pMSX30), P x L (*attP* x *attL*; pMSX24), B x R (*attB* x *attR*; pMSX28), B x L (*attB* x *attL*; pMSX26), L x L (*attL* x *attL*; pMSX34), R x R (*attR* x *attR*; pMSX36), L x R (*attL* x *attR*; pMSX18) and B x P (*attB* x *attP*; pRT504). The data are the average and standard error for two replicates.

The residue at position 449 determines the extent of *attL* x *attR* recombination

Int E449G was the most hyperactive integrase mutant isolated from the XL1-Red mutagenesis experiment. E449 was targeted for site-directed mutagenesis. The mutant proteins were purified and their activities measured by *in vitro* recombination (Figure 1B–E). Except for E449D, all the E449 substitutions tested were active in excision to varying degrees (Figure 1C). This suggests that the negative charge prevented *attL* x *attR* recombination.
Int E449Q and Int E449Y were also relatively poor at attL × attR recombination suggesting that a polar uncharged side chain can also inhibit excision. Generally, the substitutions at E449 were not greatly affected in attP × attB recombination (Figure 1B). The most affected substitution was Int E449Y but as Int E449F was as active as wt Int, steric hindrance alone cannot account for this reduced activity.

Int mutants containing substitutions at E449 that were competent for excision were also able to perform attL × attL and attR × attR recombination (Figure 1D and E). Little or no recombination was observed between other pairwise combinations of att sites (attP × attP, attB × attB, attP × attL, attP × attR, attB × attR, attB × attL; (Figure 1F and data not shown). The Int mutant most active in attL × attR recombination and most similar to wt integrase for attP × attB recombination was E449K (Figure 1B and C).

The hyperactive integrase, Int E449K, synapses attL and attR and other pairs of att sites

Previous studies have shown that wt Int does not synapse sites other than attP with attB (25,27). Synapse assays make use of an active site mutant of integrase, Int S12A, in which the serine nucleophile is altered and this prevents DNA cleavage (25). An integrase double mutant containing both S12A and E449K was purified. The intensities of the synaptic complexes obtained with Int S12A, E449K were highest with attP/attB, attL/attR, attL/attL and attR/attR as substrates (Figure 2). This was consistent with the ability of Int E449K to recombine attP × attB, attL × attR, attLxattL and attR x attR (Figure 1). However, synaptic complexes were also observed with Int S12A, E449K with pairs of att sites which, in recombination assays using Int E449K, did not yield products, e.g. attB × attB (Figures 1F and 2).

A possible explanation for the ability of Int E449K to synapse most pairs of recombination sites is that this protein is tetrameric in solution. The oligomeric state of the hyperactive integrases was analysed by gel filtration. Wt Int, Int L460P, Int Y475H, Int E449G and Int E449K were all shown to have retention times consistent with dimers in solution (data not shown; 27). Thus, like wt integrase, Int E449K probably binds its substrates as a dimer and then forms the synaptic interface by protein–protein interactions within a tetramer.

The double mutants Int S12A,Y475H and Int S12A,L460P were prepared and used in synapsis assays. Neither was able to form a detectable synapse with attP × attB nor attL × attR (data not shown). In addition, DNA-binding affinities by the hyperactive integrases were not significantly different from wt Int (Supplementary Material Figure S1). Int Y475H was defective in recombination compared to wt Int but had higher DNA-binding affinities for all four att sites than wt Int.

The hyperactive integrase, Int E449K, has gained the ability to recombine attL × attR without affecting integration

If the integration and excision reactions are in a reversible equilibrium and the rate of attP × attB recombination...
is much greater than \( attL \times attR \) recombination, then integration by wt Int might appear to be irreversible. The observed \( attL \times attR \) recombination in the hyperactive integrases could therefore be a consequence of a decrease in the overall rate of \( attB \times attP \) recombination (Figure 3).

When the supercoiled plasmid substrates encoding either \( attP \) (pRT702) or \( attB \) (pRT600) were mixed with either wt Int or Int E449K (at 15°C to slow the reaction), the initial rates of recombination appeared similar up to the 8 min time point. After this time wt Int consistently yielded more products. To test this quantitatively, a recombination assay was performed using a radio-labelled DNA fragment encoding \( attP \) and an unlabelled fragment encoding \( attB \) (Figure 3B and C). This assay confirmed that the initial rates of recombination by wt Int and Int E449K were similar in the \( attP \times attB \) assay, but after the first few

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**Figure 3.** Rate of recombination by wt versus E449K integrases. (A) A time course of \( attP \times attB \) recombination and direct detection of recombinants by restriction and agarose gel electrophoresis. The time course was performed at 15°C to slow recombination with 733 nM integrase and reactions stopped by incubation at 80°C, 10 min. Reactions were digested with HindIII. The product of integration, pRT602700, gave two fragments, 5435 bp (attL) and 91 bp (attR; data not shown), resulting from the fusion of the substrates, 3034 bp (pRT700, \( attP \)) and 2513 bp (pRT602, \( attB \)). (B) Time course of \( attP \times attB \) recombination using linear DNA fragments. Purified integrases (733 nM) were incubated with radio-labelled \( attP \) (98 bp; 1.5 nM) and a 235 bp fragment encoding \( attB \) (13 nM) for a time series at 30°C. The reactions were stopped by heating (80°C, 10 min) and then digested with subtilisin before loading onto a 5% polyacrylamide gel. The gel was dried and the radioactivity assayed using a phosphorimager. The positions of free probe (\( attP \)) the products of recombination (\( attL \) and \( attR \)) and the cleaved \( attP \) fragments (\( attP' \) 57 bp and \( attP' \) 41 bp) are indicated. (C) The amount of radioactivity in the products (black lines) versus the cleaved substrates (grey lines) was quantified and plotted against time for (i) \( attB \times attP \) with wt integrase, (ii) \( attB \times attP \) E449K integrase.
minutes the relative rate of product formation in the assay with Int E449K was significantly reduced. The level of cleaved attP present with Int E449K was considerably higher than that seen with wt integrase (Figures 3B and 4C). This is likely to be due to the ability of Int E449K to synapse and cleave the products of attP x attB recombination (attL and attR) upon formation. The same assay used with attL x attR as the initial substrates showed that the rate of excision was much lower than the rate of integration under these conditions and the level of cleaved substrate was about three times the amount of product (Supplementary Figure S2).

This assay revealed that about 40% of the radio-labelled attP remained cleaved after 1 h incubation with wt Int. This suggests that under these conditions, post-cleavage events (i.e. strand exchange and rejoining) were very slow compared to the rates of cleavage. In previous experiments using supercoiled substrates containing attP and attB, no accumulated cleaved intermediates were observed with wt Int (25,27,36). Recombination assays were performed specifically to detect the presence of any DNA cleaved at the crossover site in supercoiled substrates but none was observed (data not shown). The accumulation of cleaved substrates appears to be a feature of using linear DNAs as recombination substrates. When φC31 integrase is used in eukaryotic cell lines there is a significantly high level of micro-deletion that destroys the recombination sites, possibly due to non-homologous end-joining (NHEJ) machinery (7,40). It seems likely that integrase is also inhibited post-cleavage in the eukaryotic nucleus, allowing time for the NHEJ proteins to find and act on the recombination intermediates.

The hyperactive mutations lie within a putative coiled-coil motif

The φC31 Int and the integrases from two other Streptomyces phages, φBT1 and TG1 were subjected to structure prediction algorithms. The φBT1 and TG1 integrases have 23 and 49% identity to φC31 Int, respectively. Sequence alignments between protein homologues can be used to train secondary structure prediction algorithms such as JPRED (41). In the region where the hyperactive mutations mapped in φC31 Int, JPRED predicted two α-helices, H1 and H2, separated by a short linker (Figure 4). The equivalent regions in φBT1 Int and TG1 Int were also predicted to contain two α-helices (not shown). Another structure prediction programme, COILS (42), searches for probable coiled-coil regions in proteins. The φC31 Int, φBT1 Int and TG1 Int were all predicted to contain a region of coiled-coil that in φC31 Int spans the hyperactive region (Figure 4, Table 2). Coiled-coil motifs are normally identified by the presence of a canonically repeating heptad of amino acids, abcdεg, occupying positions (a)–(g) that can be displayed on a helical wheel (Figure 4C). Typically, within a coiled-coil domain amino acids at positions (a) and (d) are hydrophobic and (b), (c), (e), (f) and (g) are polar (43). The predicted coiled-coils in φC31, φBT1 and TG1 all appeared to have a hydrophobic position (d). However, the (a) position in all three of these proteins had two or more positively charged residues that would normally consist of hydrophobic amino acids. Most striking was the abundance of negatively charged amino acids at position (g). The COILS program compares the output from ‘weighted’ and ‘unweighted’ scans. The unweighted scan gives equal importance to the presence of the polar residues as to the hydrophobic residues but this can lead to false positives when the sequence is rich in polar residues. The weighted scan gives the same importance to the two hydrophobic positions as to the five polar positions thus reducing the impact of an abundantly polar sequence. If the difference between the two scans is >20–30% the prediction is likely to be a false positive. In addition, the COILS prediction uses three scanning windows; 14, 21 and 28 aa. A 28 residue window that displays the heptad repeat is highly likely to have a genuine coiled-coil motif, while the smaller scanning windows are more prone to false positives that would arise from a random sequence of amino acids. The φC31 integrase H1-helix maintained a high probability of forming a coiled-coil in both the weighted and unweighted scans using the 28 aa window (Figure 4: ‘Lupas28’ prediction). The H2-helix, however had a >40% drop in the probability of coiled-coil in the weighted versus the unweighted scans with the 28 and 21 residue windows. Only the smaller 14-residue scanning window maintained a >90% probability of forming a coiled-coil (Figure 4). E449 is located at the start of the longest predicted coiled-coil region and so has been included within the schematic representation despite the presence of the probable helix-breaker, P451. Y475 and L460 are located in the predicted coiled-coil domain. φBT1 H1-helix and TG1 H1- and H2-helices were predicted to form coiled-coils using the COIL algorithm (Figure 4, Table 2).

Other large-serine recombinases also showed the presence of predicted α-helix in the C-terminal domain, many of which had a high probability of forming coiled-coils (Table 2). A coiled-coil domain has also been predicted in the large serine transposase TnpX from Tn4451 (44).

Residues that confer hyperactivity lie on one face of the putative coiled-coil motif

A spontaneous mutant of pHS62 was isolated that appeared to be active in attL x attR recombination using the DH5α (pPAR1000) reporter strain. Sequencing of the int gene from this spontaneous mutant revealed a mutation that caused the E463K substitution. E463K was able to give 100% recombinants when introduced into DH5α (pPAR1000) or DH5α (pRT504). E463K, like E449K resides on the (g) face of the predicted canonical coiled-coil (Figure 4C and D). Because of the location and nature of the mutations conferring hyperactivity, SDM was used to change all the acidic residues from E449 to D477 to lysine with the hope of isolating more hyperactive mutants. Two additional mutant integrases were isolated that could recombine attL x attR in vivo; E452K and E456K (Figure 4D). E449K, E456K and E463K all reside at the (g) position of the proposed coiled-coil and E452K lies in the adjacent (c) position (Figure 4C). Mutations of the other residues were either
as wt (D466K, A470K, E473K, E476K and A479K) or inactive (D477K) (Figure 4C and data not shown). These data are consistent with this region presenting an interaction surface, which is altered in the hyperactive proteins.

**Structural asymmetry in the Int E449K synaptic complexes with attL and attR**

In the integration reaction, the polarity of the recombination sites is determined only by the nature of the 2bp at the crossover site (26). In φC31 att sites, this is 5’T and normally the left side of attB (B arm) joins to the right side of attP (P arm) to make attL (BrrP) and the reciprocal product, attR (PrrB). If the crossover sequence is changed to 5’TA in both substrates or 5’AA in one substrate then polarity is either lost (5’TA) or switched (5’AA) (26). We showed previously that with wt Int, attB<sup>Tα</sup> and attP<sup>Tα</sup> recombines to give the normal products, BrP<sup>Tα</sup> (attL<sup>Tα</sup>) and Pt<sup>Tα</sup>B<sup>Tα</sup> (attR<sup>Tα</sup>) and aberrant products, BrP and P<sup>Tα</sup>B, in equal quantities (26). This was interpreted to mean that wt Int forms two synapse structures, differing by the spatial arrangements of P, P<sup>Tα</sup> and B, B<sup>Tα</sup> arms in the complex [Figure 5A(iv)]. Both synapse structures contain
Table 2. Location of putative coiled-coil regions in phage serine integrases

| Origin of integrase | Size (aa) | Predicted coiled-coil region(s)^a | Region size (aa) | Accession number |
|---------------------|----------|----------------------------------|-----------------|-----------------|
| Phage φC31          | 605      | 452–480/489–518                 | 28/29           | NP047974        |
| Phage TG1           | 606      | 463–497/506–533                 | 34/27           | BAF03600        |
| Phage Bcep22        | 518      | 350–387/390–423                 | 37/33           | AAQ54941        |
| Phage U153          | 453      | 337–366/374–424                 | 29/50           | CAD10283        |
| Prophage CP-933X     | 508      | 367–398/400–427                 | 31/27           | NP287356        |
| Phage φFC1          | 464      | 343–370/385–423                 | 27/35           | AAD26564        |
| Phage φ105          | 477      | 358–385/388–415                 | 27/27           | NP690785        |
| Phage TP91          | 398      | 284–314/316–350                 | 30/34           | CAAT2268        |
| Phage SPBc2         | 549      | 398–426/444–475                 | 38/31           | NP046553        |
| Phage 370           | 476      | 385–412                         | 27              | NP606888        |
| Phage TP901-1       | 487      | 405–434                         | 29              | NP112664        |
| Phage φK381         | 552      | 427–455                         | 28              | BAF03598        |
| Phage φBT1          | 594      | 435–469                         | 34              | NP813744        |
| Phage φRV1          | 469      | 346–380                         | 34              | NP216102        |

^aBold values indicates a high probability of forming a coiled-coil structure based on the COILS programme (42).

Int subunits bound to P or P’ arms making a synaptic contact with an Int subunit bound to the B or B’ arm. The premise is that integrase does not dictate a preference for the arrangement of the B and B’ arms with respect to the P or P’ arms in the integrative synapse [Figure 5A (iv)]. Where an Int subunit is bound to a B-type half site and synapses with an Int subunit bound to a P-type half site, we call this interaction ‘complementary’ (Figure 5A). Both synaptic structures in attP x attB recombination involve complementary pairings of Int subunits [Figure 5A (iv)]. The ability of the hyperactive integrases to recombine attL x attR, attL x attL, attR x attR (Figure 1) provided the opportunity to interrogate the nature of the excisive synapse. When each pair of sites is brought together by Int E449K, we expect that two alternative synapse structures are possible that differ markedly from each other [Figure 5A (i) to (iii)]. One type of synapse involves complementary subunit interactions and in the other type of synapse the interactions between the Int subunits are non-complementary, i.e. both Int subunits contact a P-type arm or a B-type arm. If the interaction by an integrase subunit bound to P-type arms is different to that bound to B-type arms then the Int E449K synaptic complexes containing the hybrid sites, attL and attR, may be structurally biased.

Plasmid substrates encoding attL^TA (pMS92) and attR^TA (pMS93) were mixed and incubated with Int E449K at 15°C (Figure 5A–C). The products of recombination reactions containing pMS92 and pMS93 are predicted to include recombinants between three different reactions; attL^TA x attR^TA, attL^TA x attL^TA and attR^TA x attR^TA [Figure 5A (i), (ii) and (iii)]. From each reaction two products (co-integrates 1 and 2) could, theoretically, be made, arising from alternative synapse structures that contain complementary or non-complementary Int interactions (Figure 5A). The two co-integrates from the attL^TA x attR^TA recombination can be detected with XbaI (co-integrate 1; 5458 bp; Figure 5B) or BamHI (co-integrate 2; 5462 bp; Figure 5C). Co-integrate 1 appeared well before co-integrate 2 suggesting that when the sites come together in a synapse, complementary pairing of Int E449K subunits is preferred to the non-complementary pairing [Figure 5A(i), B and C]. As seen previously with wt Int (26), Int E449K did not show any bias towards generating either co-integrate 1 or 2 when the substrates were attP^TA and attB^TA (data not shown).

In these reactions, we also observed attLTA x attLTA and attR^TA x attR^TA recombination. The appearance of co-integrate 2-type products are indicated by the presence of the ~4900 bp band for attL^TA x attL^TA and the ~5980 bp band for the attR^TA x attR^TA in both the BamHI and XbaI digests. These appeared at the same time as the favoured product (co-integrate 1) from the attL^TA x attR^TA reaction [Figure 5A(ii), (iii), B, C] and well before the appearance of the attL^TA x attR^TA unfavoured (co-integrate 2) product. The proposed synapse structures that generated the co-integrate 2-type products from attL^TA x attL^TA or attR^TA x attR^TA recombinations are those where the bound Int E449K subunits interact in complementary pairings, i.e. a subunit bound to a P-type arm interacts with a B-type arm bound subunit (Figure 5A). Thus, in all three reactions, attL^TA x attR^TA, attL^TA x attL^TA and attR^TA x attR^TA products generated from synaptic complexes where the Int E449K interactions are complementary appear at the same time and well before products that arise from synaptic complexes containing non-complementary subunit pairings. It is possible that the unfavoured product of attL^TA x attR^TA recombination (co-integrate 2) could have arisen indirectly through the generation of co-integrate 1 followed by inversion of DNA between the attP^TA and attB^TA sites. To test whether products could be obtained at all via a synapse between attL and attR where the Int E449K subunits interact via non-complementary pairings, it was necessary to invert the polarity of one of the sites by changing the dinucleotides at the crossover site. Two new plasmids, pPARX18L^AA and pPARX18R^AA were generated by SDM of pMSX18 so that the crossover dinucleotides in attL were changed to attL^AA in pPARX18L^AA and those in attR to attR^AA in pPARX18R^AA. Due to the constraint provided by the crossover dinucleotides the only possible recombination products from pPARX18L^AA and pPARX18R^AA are inversion products arising through synaptic complexes in which Int subunits are in non-complementary pairings (Figure 6A). Inversion products were indeed detected, indicating that non-complementary pairings are possible in attL x attR recombination (Figure 6B). The outcome of this experiment was anticipated by the high number of deletion products from E449K Int acting on attL x attL and attR x attR (pMSX34 and pMSX36, respectively; Figure 1), reactions in which the Int subunit interactions were likely to be non-complementary.

These data show that the synaptic complexes between attL/attR and the hyperactive integrase, Int E449K, are structurally biased towards a synapse that proceeds through recombination to generate attP and attB rather than aberrant recombination products (Figure 5). This structural bias between two forms of synapse is not observed for attP x attB recombination (26).
Figure 5. Structural bias in the formation of the Int E449K synaptic complexes with attL and attR sites. (A) Recombination products from plasmids encoding attL<sup>T</sup>A (pMS92), attR<sup>T</sup>A (pMS93), attB<sup>T</sup>A (pRT603) and attP<sup>T</sup>A (pRT701). Products (co-integrates 1 and 2) are derived from reactions between (i) pMS92 (attL<sup>T</sup>A) and pMS93 (attRTA) (ii) pMS92 and pMS92 (iii) pMS93 and pMS93 and (iv) pRT603 (attB<sup>T</sup>A) and pRT701 (attP<sup>T</sup>A). Each attachment site is represented as an arrow. The halves without the arrow heads are the B or P arms, the halves with the arrowheads are the B<sup>0</sup> or P<sup>0</sup> arms. B-type arms are blue and P-type arms are red. The structures of the proposed synaptic complexes are indicated by the alternative alignments of P or P<sup>0</sup> and B or B<sup>0</sup> arms. The putative synaptic tetramer of integrase subunits is shown with each subunit coloured according to whether it is interacting with a B-type arm (blue) or a P-type (red). Int interactions at the synaptic interface are either complementary (Int bound to P-type arms interact with Int bound to B-type arms) or non-complementary (Int subunits that are both bound to P-type or B-type arms). The positions of relevant restriction sites are indicated (B, BamHI; X, XbaI) and the expected restriction fragment sizes of the parental plasmids and the co-integrate products are shown.
predicted coiled-coil (Figure 4). By analogy with other proteins containing coiled-coil domains the putative coiled-coil motif in φC31 integrase may comprise a protein-protein interaction surface (43). In the resolvases, the synaptic interface occurs via the N-terminal domain and hyperactivity (in this case defined as an ability to recombine substrates in the absence of activation by resolvase subunits bound to accessory sites) results from mutations in this region (45,46). If the synaptic interface in φC31 integrase also occurs exclusively through the N-terminal domain, the hyperactive region may enable synapsis indirectly through communication between the C-terminal domain and the N-terminal catalytic domain. Thus, the C-terminal domain may control the formation of the synaptic interface in an analogous way to resolvase responding to the presence of accessory sites or Hin interacting with Fis (16,22). Alternatively, the putative coiled-coil region may act directly through inter-molecular interactions. Possibly, the coiled-coil domain is normally sequestered unless integrase is bound to attP and attB when the coiled-coil region becomes exposed. If the mutations conferring hyperactivity destabilize the sequestration then the coiled-coil domain may be exposed and able to form inter-molecular interactions even when integrase is bound to attL or attR. This is the first indication that synthesis might occur via the C-terminal domain. In both models, synthesis is the step which activates recombination.

Additionally, several mutations within the C-terminal coiled-coil region (Int L460P, Int Y475H and Int D477K) were found to be defective in integration. Derivatives of Int L460P and Int Y475H were unable to form stable synaptic complexes. Clearly the putative coiled-coil domain has an important role in both integration and excision.

It is evident that the integration and excision reactions in the hyperactive mutants are different. Most striking is that Int E449K showed a bias towards one of two alternative synaptic complexes with respect to the spatial positioning of P, P' and B, B' arms in attL x attR recombination, whereas no bias was observed with attP x attB as substrates. The bias in the excision synapse is consistent with an Int subunit bound to a P-type arm preferring to synapse with an Int subunit bound to a B-type arm (Figure 5A). This implies that integrase has a mechanism of transmitting information, probably through protein conformation, from the DNA sequence to which it is bound to the synaptic interface. If this is the case, then Int complexes bound to attP or attB will be conformationally symmetrical but not the same. The symmetry arises from the sequences of attP and attB, which are imperfect inverted repeats. When attP and attB sequences are aligned (with the crossover dinucleotide in the centre), 15 out of 38 bases are identical (36). Most of the sequence divergence lies at the extremities and it is here that mutations in the attB site can have highly detrimental effects on the activity of attB in integration (36). Possibly, it is these unique sequences that specifically affect the conformations of integrase subunits bound to P-type arms or to B-type arms.

The excision reaction catalysed by φC31 Int E449K has similar features to Bxb1 integrase, catalysing attL x attR
recombination in the presence of a recombination directionality factor (RDF; gp47) (38). In the presence of gp47, Bxb1 integrase also has a preference for one of the two possible synaptic complexes with respect to the arrangement of P- or B-type arms (38). Because the \( C31 \) excision reaction with Int E449K occurs without the accessory protein it is possible to infer that the conformational 'readout' from DNA sequence interactions to the structure of the synaptic interface is inherent in the mechanism of the large serine recombinases generally. We emphasize however that Int subunits do not act independently of each other in adopting a conformation that will enable synapsis. No productive synapsis (and almost no recombination) was observed with Int E449K where the numbers of B to P-type arms deviated from 2:2, i.e. reactions such as \( \text{attB x attR}, \text{attB x attL}, \text{attP x attL} \) and \( \text{attP x attR} \) (3:1), as well as \( \text{attP x attP} \) and \( \text{attB x attB} \) (4:0). This selectivity indicates that prior to synapsis, conformations adopted by Int dimers when binding to pairs of \( \text{att} \) sites are still the primary step in controlling against unwanted DNA rearrangements. The discovery of a putative interaction motif in the C-terminal domain of \( C31 \) integrase now provides a focus for understanding how different protein conformations mediate synapsis and the control of directionality in integration versus excision.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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