

REVIEW ARTICLE

Integrating prokaryotes and eukaryotes: DNA transposases in light of structure

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Abstract

DNA rearrangements are important in genome function and evolution. Genetic material can be rearranged inadvertently during processes such as DNA repair, or can be moved in a controlled manner by enzymes specifically dedicated to the task. DNA transposases comprise one class of such enzymes. These move DNA segments known as transposons to new locations, without the need for sequence homology between transposon and target site. Several biochemically distinct pathways have evolved for DNA transposition, and genetic and biochemical studies have provided valuable insights into many of these. However, structural information on transposases – particularly with DNA substrates – has proven elusive in most cases. On the other hand, large-scale genome sequencing projects have led to an explosion in the number of annotated prokaryotic and eukaryotic mobile elements. Here, we briefly review biochemical and mechanistic aspects of DNA transposition, and propose that integrating sequence information with structural information using bioinformatics tools such as secondary structure prediction and protein threading can lead not only to an additional level of understanding but possibly also to testable hypotheses regarding transposition mechanisms. Detailed understanding of transposition pathways is a prerequisite for the long-term goal of exploiting DNA transposons as genetic tools and as a basis for genetic medical applications.

Keywords: Insertion sequence; integrase; transposition; genome instability; V(D)J recombination

Introduction

DNA transposons are distinct segments of DNA that can move to new locations within a genome without the need for homology between the transposon sequence and the new DNA target site (reviewed in Curcio and Derbyshire, 2003). This distinguishes them from other types of mobile genetic elements such as bacteriophages or genomic islands which use DNA recombinases or resolvases and generally require a short region of sequence identity between the ends of the element and the target site. Another large group of mobile elements, the retrotransposons, are abundant inhabitants of many

eukaryotic genomes but much less frequent in prokaryotes. These copy themselves to a new location using an RNA intermediate that is subsequently reverse transcribed into double-stranded DNA. DNA transposons have no such intervening RNA intermediate, and many proceed through a so-called cut-and-paste mechanism involving a double-strand DNA intermediate. Recently, a completely different mechanism has been discovered in which a single strand of DNA is excised from one genomic location and moved to another (Barabas *et al.*, 2008; Guynet *et al.*, 2008). For the remainder of this review, we will focus on those pathways that proceed using both dsDNA substrates and intermediates.

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(Received 18 September 2009; revised 20 November 2009; accepted 23 November 2009)

ISSN 1040-9238 print/ISSN 1549-7798 online © 2010 Informa UK Ltd
DOI: 10.3109/10409230903505596

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There is an enormous diversity even among the class of transposons which use dsDNA intermediates. Perhaps the simplest of the autonomous transposable elements of this type, and certainly the most numerous in prokaryotes, are the insertion sequences (ISs). Most autonomous DNA elements identified in eukaryotes are also quite similar to prokaryote ISs in mechanism as well as in size. The original distinction between transposons and ISs was that the former carry additional genes to those involved directly in transposition while the latter include only the gene for their cognate transposase. More recently, detailed analysis of ISs in various bacterial genomes has revealed close relatives that include passenger genes encoding antibiotic resistance, methyltransferase activity, transcriptional regulation functions and proteins with unknown function (Siguier *et al.*, 2009), further obscuring the line separating insertion sequences and transposons.

Mechanisms of DNA transposition

Among the dsDNA transposons, the most prevalent use a transposase which carries a conserved triad of amino acids – Asp, Asp and Glu, the DDE motif – which we will discuss in detail in the next section. Retroviruses fall into this category since not only do they encode an enzyme (integrase, IN) closely related to bacterial transposases, but they integrate into the host genome using a double-stranded DNA copy of their RNA genome (reviewed in Craigie, 2001). Moreover, the V(D)J recombination system which generates diversity in immunoglobulins and T cell receptors also uses an enzyme with a DDE motif, RAG1, which promotes transposition-like chemistry and can, under certain circumstances, behave like a DNA transposon (Hiom *et al.*, 1998). In this review we will further restrict ourselves to transposition pathways that proceed through dsDNA intermediates using DDE-type enzymes.

Identification of the DDE triad stemmed from remarkable similarities observed between prokaryotic transposases and retroviral integrases (Fayet *et al.*, 1990; Kulkosky *et al.*, 1992). The crucial role of these residues in catalysis has subsequently been demonstrated by mutagenesis in many transposition systems, and the overall catalytic mechanism of enzymes containing a DDE motif has been well characterized (Mizuuchi, 1992; Mizuuchi and Baker, 2002). They catalyze a single chemical reaction: nucleophilic cleavage of a single phosphodiester bond. The tremendous diversity in transposition mechanisms derives from the combinatorial effect of using different nucleophiles on different DNA substrates in varying combinations, and whether or not replication occurs at some stage during these consecutive recombination events.

In general, transposition occurs first by cleavage of one strand at each transposon end. The nucleophile is generally H_2O (Figure 1), and this reaction produces a 3'OH. For retroviruses, this is called “3' processing”. The two 3'OH groups (one at each end of the mobile element) then serve in a second chemical step as nucleophiles in a concerted attack on both strands of the target DNA (called “strand transfer”). The two strand transfer reactions occur a few base pairs apart on the target DNA, the precise distance between them being characteristic of the particular transposase. The breaks in the target DNA are repaired by host cell enzymes, a process that duplicates the target site on both flanks of the transposon.

There are no covalent enzyme-substrate intermediates formed during this process, distinguishing it from reactions catalyzed by the ubiquitous serine and tyrosine recombinases (Grindley *et al.*, 2006). Instead DNA transposition involves a series of single-step, direct in-line nucleophilic attacks. This mechanism of direct transesterification was firmly established by observations that the phosphate of the target scissile phosphodiester bond undergoes chiral inversion (i.e., an Rp diastereomer undergoes inversion to its Sp form) when chirality is imposed by replacement of a non-bridging oxygen by a sulfur atom. This has been observed to occur during several of the steps catalyzed by various systems including bacteriophage Mu, HIV integrase, Tn10, and V(D)J recombination (Mizuuchi and Adzuma, 1991; van Gent *et al.*, 1996; Mizuuchi, 1997; Gerton *et al.*, 1999; Kennedy *et al.*, 2000).

In spite of their shared transposition chemistry, different DDE-type transposons have developed different variations in the steps leading to formation of a unique insertion intermediate (Figure 1). These differences reflect the way in which the second (non-transferred) strand is processed (Turlan and Chandler, 2000).

A first distinction is whether the second strand is cleaved at all. The simplest scenario is that used in generating the dsDNA intermediate seen during retroviral integration. Retroviral integrase, IN, removes two bases from the transferred strand at each end (Figure 1c). As neither strand was ever joined to anything, this leaves recessed ends with the characteristic 3'OH nucleophile ready for the strand transfer step (Craigie, 2001). In the cases of bacteriophage Mu, transposon Tn3 (and other members of this large family) and the IS6 family of prokaryotic insertion sequence, the non-transferred strand remains intact (Figure 1a) and strand transfer fuses both donor and target DNA molecules to form a so-called Shapiro intermediate (Shapiro, 1979). The branched fusion molecule, in which each transposon end is joined to the donor by one strand and the target by the other, is resolved by replication of the transposon.

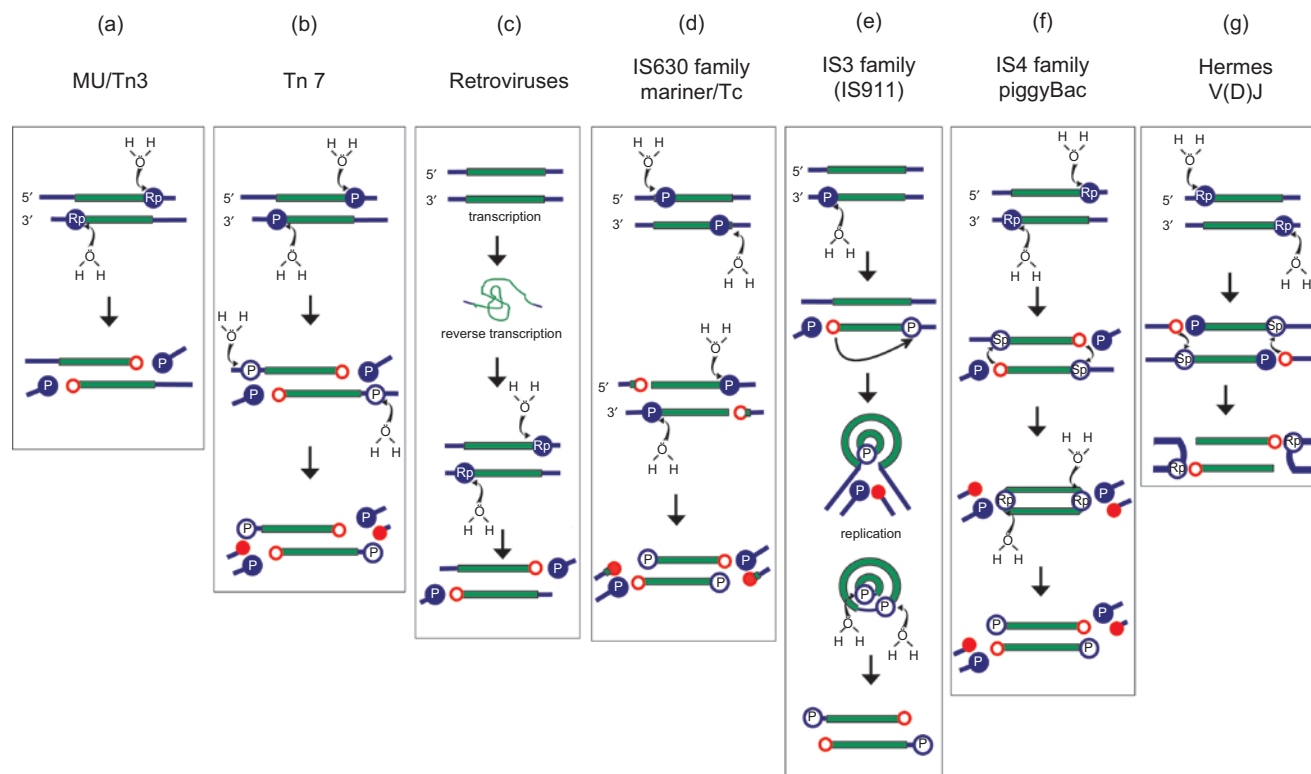


Figure 1. Dealing with the second strand. The color code is as follows: transposon DNA (green); flanking donor DNA (blue); target phosphates destined to be removed from the final liberated transposon (filled blue circles with a white “P”); phosphates destined to remain as 5′ transposon ends (open blue circles); the preferred stereoisomer, Sp or Rp, where known, is indicated within the circles; liberated 3′OH groups involved in strand joining reactions (open red circles); 3′OH destined to be removed from the liberated transposon (filled red circles); H₂O is the attacking nucleophile in the hydrolysis reactions. (a) The Mu and Tn3 cleavage reactions. Note that the preferred stereoisomer has been demonstrated only for Mu and not for Tn3. (b) Tn7 cleavage reactions. Cleavage of the transferred strand (top of panel) is shown occurring prior to cleavage of the non-transferred strand (middle) leading to liberation of the transposon from flanking donor DNA (bottom of panel), although this order of cleavage reactions has not been demonstrated experimentally. The two types of cleavage are catalyzed by different enzymes. (c) Retroviral “processing” reaction, equivalent to cleavage of the transferred strand. An initial transcription step from the integrated provirus is indicated. The RNA genome is then encapsidated with a second copy and undergoes reverse transcription following infection to generate the double strand DNA integration intermediate. The intermediate is flanked by only short fragments of donor material and does not require second strand processing for insertion. (d) Transposition by the members of the IS630 family and the Tc1/Mariner superfamily is initiated by cleavage of the non-transferred strand (top of panel) at several bases within the transposon end (middle) leaving these bases attached to the liberated flanks following cleavage of the transferred strand (bottom). (e) For IS911, IS2, IS3 and other members of the IS3 family, single-end hydrolysis occurs (top). The liberated 3′OH then directs a strand transfer reaction to the same strand several bases 5′ to the other end of the element. This results in the formation of a single-strand circle which is then resolved into a transposon circle by replication from the free 3′OH (filled red circle). Single-strand hydrolysis at each 3′ end within the circle generates a linear transposon which can then undergo integration. (f) The IS4 family and *piggyBac* have similar mechanisms. Following initial nucleophilic attack on the Rp target phosphate, the liberated 3′OH attacks an Sp phosphate in a trans-strand transfer reaction to generate a hairpin intermediate, liberating the transposon from its flanking donor DNA and inverting the target phosphate to its Rp configuration. These then become the substrates for a second hydrolysis. Note that the stereochemistry has been analyzed only in the case of Tn10. (g) *Hermes* and V(D)J transposition occur by initial cleavage of the non-transferred strand (top). The liberated 3′OH on the donor flank then attacks the opposite strand (middle) to generate hairpin structure on the donor flank (bottom). The stereochemistry has been analyzed for V(D)J only. Modified and reprinted from Turlan and Chandler (2000), with permission from Elsevier.

If both donor and target DNA are circular replicons (such as plasmids or chromosomes), this generates a cointegrate where donor and target DNA are joined and delimited at each junction by a single transposon copy. Transposition is then completed by recombination between the directly repeated transposons either using a host recombination system or as, for example with Tn3, using dedicated site-specific recombination

systems encoded by the transposon itself (Grindley, 2002).

For all other known elements, cleavage of the second strand does occur, liberating the transposon from flanking DNA at the donor site. For the transposon Tn7 (Figure 1b), two different enzymes are used to generate an excised intermediate: TnsB catalyzes cleavage of the transferred strand and is predicted to contain a DDE

motif, and TnsA catalyzes cleavage of the non-transferred strand and leaves a three base 5' extension which is presumably removed during the DNA repair steps which must follow integration (Sarnovsky *et al.*, 1996). Tn7 transposition requires the presence of target DNA to activate cleavage and transfer (reviewed in Craig, 2002; Parks and Peters, 2009). For members of the IS630 family and the related Tc1/*mariner* eukaryotic transposons, cleavage of the non-transferred strand occurs prior to cleavage of the transferred strand and at several bases within the transposon (Plasterk *et al.*, 1999; Feng and Colloms, 2007; Figure 1d).

Members of the large IS3 family of prokaryotic ISs have developed yet another way of resolving the second-strand cleavage problem (Figure 1e; reviewed in Rousseau *et al.*, 2002). Here, the transposase catalyzes cleavage at one end only. The resulting 3'OH is then used to attack the same strand three or four nucleotides within the DNA flank at the opposite end, thus creating a single-strand bridge between the two ends and a 3'OH positioned in the flanking donor DNA. This can act as a primer for DNA replication. Replication then produces a free covalently-closed circular IS in which the ends are abutted but separated by three or four base pairs (bp) of the original flanking DNA, and regenerates the original donor molecule. In the integration step, transposase catalyzes appropriate cleavages at each end to yield an integration intermediate with 3'OH ends.

Another way of ensuring second strand cleavage has been adopted by members of the IS4 family, Tn5 (IS50) and Tn10 (IS10) (Figure 1f; reviewed in Haniford, 2006; Reznikoff, 2008) and the related eukaryotic *piggyBac* transposon superfamily. Here the 3'OH generated at the end of the transferred strand is used to attack the opposite (non-transferred) strand to generate a hairpin. The DNA at the hairpin end is then cleaved, again using H₂O as the nucleophile to create a typical insertion intermediate. In the case of *piggyBac*, the geometry of cleavages is such that the excised intermediate carries a 5' tetranucleotide extension consisting of flanking DNA on both non-transferred strands (Mitra *et al.*, 2008).

Finally, to complete our list of known ways by which dsDNA transposons are mobilized, both the V(D)J system (reviewed in Jones and Gellert, 2004) and the members of the *hAT* family of eukaryotic transposons such as *Hermes* (Warren *et al.*, 1994; Zhou *et al.*, 2004) have adopted yet another pathway for second strand cleavage (Figure 1g). Here, cleavage of the non-transferred strand creates a 3'OH on the DNA flank. This then attacks the phosphodiester bond on the transferred strand generating the appropriate 3'OH necessary for strand transfer, and in so doing, creates a hairpin structure in the DNA flanks.

The fundamental building block of DNA transposases: the catalytic core

The common structural features of RNase H-like catalytic domains

The ability of transposases containing a DDE triad to catalyze nucleophilic cleavage of a single phosphodiester bond is a direct result of the role of these three acidic residues. The two Asp residues and a third (which is usually Glu but sometimes Asp, hence the broader DDE/D shorthand that we will use from now on) coordinate two divalent metal ions. These metal ions, most likely Mg²⁺ *in vivo*, are essential cofactors for both the DNA strand cleavage and strand transfer steps. The two metal ion catalytic mechanism was first described by Beese and Steitz (1991) for the exonuclease domain of the Klenow fragment, a domain that is topologically related to the transposase catalytic domains. The validity of the two metal ion mechanism for transposases was later confirmed by the crystal structures of the IS50 transposase (referred to as the "Tn5 transposase"; Davies *et al.* 2000; Steiniger-White *et al.*, 2002), and was further extended to the topologically-related RNase H (Nowotny *et al.*, 2005).

The striking feature of transposases which have catalytic domains containing the DDE/D triad is their topological similarity, despite the often negligible and therefore unrecognizable sequence similarity between them. There is also significant variation in the size of the domains, a consequence of variations in the lengths of the loops connecting the secondary structure elements, the lengths of the secondary structure elements themselves, and – as we shall see – extra domains inserted into the catalytic domain. Figure 2 shows the three-dimensional structures of a representative set of DDE/D catalytic domains that are involved either in transposition or retroviral integration. The conserved core of the domain is a mixed alpha-beta fold, $\beta 1$ - $\beta 2$ - $\beta 3$ - $\alpha 1$ - $\beta 4$ - $\alpha 2/3$ - $\beta 5$ - $\alpha 4$ - $\alpha 5$ (Figure 3), that was first seen for RNase H (Yang *et al.* 1990); therefore, it is probably most accurate to refer to it as the "RNase H-like fold" or "ribonuclease H-like fold" as this is the terminology that has been adopted by the protein fold classification database SCOP (<http://scop.mrc-lmb.cam.ac.uk/scop/>). More precisely, it is a three-layered $\alpha/\beta/\alpha$ domain with a central, mixed five-stranded β -sheet with a characteristic strand order of 32145, where strand 2 is antiparallel to the rest (Figure 2). In the literature, the fold is sometimes referred to as the "retroviral integrase fold".

The DDE/D catalytic triad of carboxylate side chains are always located on topologically equivalent secondary structure elements, with the first D always on $\beta 1$, the second D always on or just after $\beta 4$, and the E/D either on or just before $\alpha 4$. In crystal structures of uncomplexed

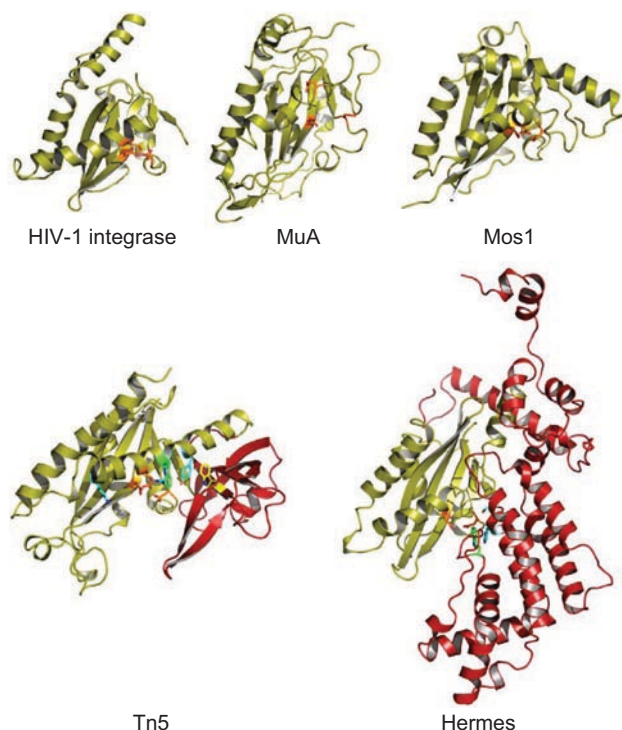


Figure 2. Ribbon diagrams of the aligned catalytic cores of four DNA transposases and of HIV-1 integrase. Residues shown in orange are the carboxylate active site residues, in green are the W residues of the Tn5 transposase and Hermes that are important in the reactions, in blue are the YRK residues of the YREK motif and in yellow is W298 of the Tn5 transposase. The insertion domains of the Tn5 transposase and Hermes are shown in red (note that there is a 15 amino acid loop from residues S481 to K496 that is disordered – and therefore not visible – in Hermes). The proteins are drawn to scale.

(i.e., without DNA bound) transposases, the E/D residue is sometimes on a loop displaying some degree of disorder. For instance, in the crystal structure of the catalytic fragment of the bacteriophage Mu transposase (MuA) (Rice and Mizuuchi, 1995), the E residue (E392) was visible in the electron density but it was also clear that the observed configuration of the catalytic triad would be unable to coordinate two metal ion cofactors, indicating the need for a conformational change. In the first crystal structure of the catalytic domain of HIV-1 IN (Dyda *et al.*, 1994; reviewed in Jaskolski *et al.*, 2009), a 14 amino acid stretch that contains the catalytic E is completely disordered, without detectable electron density. It is quite possible that in many cases these observed disordered loops become ordered upon DNA binding, folding into a regular α -helical structure and becoming an upstream extension of α 4. This means that, at least for some transposases, the active site assembles fully only after the transposon end DNA is captured and is ready for cleavage. This is certainly not always the case as comparison of the uncomplexed Tn5 transposase structure with that bound to transposon-end DNA shows

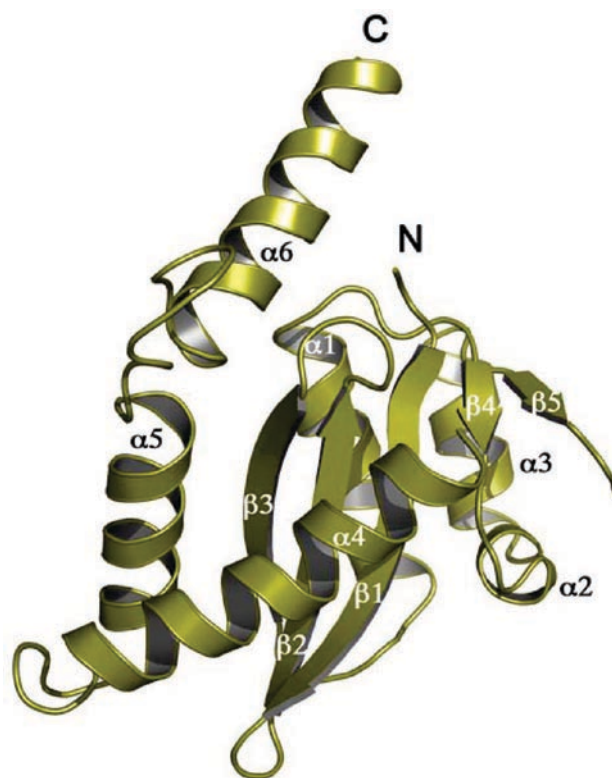


Figure 3. Ribbon diagram of the catalytic core domain of HIV-1 integrase, with the standard secondary structure elements highlighted.

little protein main chain movement, although there are some significant side chain movements (Davies *et al.*, 1999; 2000).

To date, there are only two transposases for which crystal structures have been determined in which both transposon DNA and the transposase catalytic domain are present. These are of the prokaryotic Tn5 transposase dimer bound to two transposon ends (Davies *et al.*, 2000; reviewed in Steiniger-White *et al.*, 2004 and Reznikoff, 2008) and the eukaryotic Mos1 DNA transposase dimer bound to two oligonucleotides representing the transposon Right End (Richardson *et al.*, 2009). In both cases, the bound DNA approximates a version of an intermediate along the transposition pathway in which both strands have already been cleaved. An EM reconstruction is also available for the MuA transposase tetramer bound to two pre-cleaved 50-mer Right Ends (Yuan *et al.*, 2005). Although not directly related to mobile DNA, crystal structures have been determined of RNase H bound to DNA–RNA hybrids (Nowotny *et al.*, 2005), and insights from these structures as they relate to DNA transposition have recently been reviewed (Nowotny, 2009). In addition, an EM study of the RAG1/2 complex bound to recombination signal sequence (RSS) DNA has recently been reported (Grundy *et al.*, 2009).

The catalytic cores of some DNA transposases are interrupted by insertion domains

Interestingly, the region of the catalytic core domain between $\beta 5$ and $\alpha 4$ may have additional significance in the reaction chemistry. As observed in the HIV-1 IN, ASV IN, and MuA transposase structures, the RNase H-like fold generally has a short (7–15 amino acids) loop connecting $\beta 5$ and $\alpha 4$ that is either disordered or only weakly ordered. In contrast, the Tn5 transposase has a 96 amino acid long domain inserted between $\beta 5$ and $\alpha 4$ (Davies *et al.*, 1999). We refer to this domain, or any other domain that is inserted at this topological location in the standard RNase H-like core, as an “insertion domain”. In the case of Tn5, the insertion domain is mostly β -stranded, containing only one short two-turn α -helix and four β -strands that extend the central β -sheet of the catalytic domain. This insertion domain performs a crucial function as it forms a number of intricate interactions with the DNA hairpin that is generated when the 3'OH group of the transferred strand (created at the first cleavage step) attacks the phosphate of the non-transferred strand (Davies *et al.*, 2000; Figure 1f).

A particularly interesting residue, W298, is found within the Tn5 transposase insertion domain. This stacks with the second base of the non-transferred strand which protrudes (is “flipped out”) from the DNA helix (see Figures 6 and 7 in Steiniger-White *et al.*, 2004). This facilitates formation of the tight, short loop hairpin at the transposon end. Importantly, in those crystal structures of transposases where second strand cleavage does not proceed through a hairpin intermediate (MuA, Mos1, and HIV-1 IN) there is no insertion domain between $\beta 5$ and $\alpha 4$. This raises the intriguing possibility that its presence or absence might correlate with the mechanism of second strand processing.

More recently, another type of insertion domain has also been seen – inserted at the same topological location of the standard RNase H-like fold – in the crystal structure of the eukaryotic *Hermes* transposase (Hickman *et al.*, 2005). The long *Hermes* insertion domain (residues 265–552, counted from the end of $\beta 5$ to the beginning of $\alpha 4$) is entirely α -helical and therefore bears no resemblance to the insertion domain of the Tn5 transposase. Indeed, both the Tn5 and the *Hermes* transposase insertion domains are structurally unique. Although, like Tn5, *Hermes* also forms hairpins during transposition, the hairpin is generated on the flanking DNA ends rather than on the transposon ends (Zhou *et al.*, 2004). This is because the *Hermes* transposase first catalyzes cleavage of the non-transferred strand to generate a 3'OH group on the DNA flank; this then attacks the terminal phosphodiester bond on the opposite (transferred) strand to create a flanking DNA

hairpin and a 3'OH group at each end on the transferred strand (Figure 1g).

The *Hermes* insertion domain also contains a conserved Trp residue, W319, shown to be important for one or more steps before strand transfer (Hickman *et al.*, 2005). It presumably acts to facilitate hairpin formation as does the Tn5 transposase W298 residue. Consistent with the interpretation that W319 is therefore likely to be catalytically important, it is located close to the DDE side chains of the active site.

Interestingly, RAG1 is predicted to contain a large insertion domain between the second and third acidic residues of its identified DDE motif (264 residues, counting from D708 to E972), and secondary structure prediction clearly indicates it is likely to be entirely α -helical (Zhou *et al.*, 2004) as is that of *Hermes*. It also contains a conserved Trp, W893, and mutation generates a serious defect in V(D)J recombination; however a role in hairpin formation/stabilization has not been demonstrated (Grundy *et al.*, 2007). Since there is no high resolution three-dimensional structure of RAG1, it is not known if its insertion domain is structurally similar to that of *Hermes*, although it is certainly smaller (264 amino acids separate the second D and the third E in RAG1, whereas the separation is 324 amino acids in *Hermes*).

The *Hermes* insertion domain also contributes to formation of the functional transposase oligomer, another contrast with the insertion domain of Tn5 which plays no apparent role in oligomerization. This additional functionality may partially explain why it is substantially larger than the 96-residue insertion domain of the Tn5 transposase. There is another complication with the reactions catalyzed by *Hermes* and the V(D)J recombination system, relative to that of Tn5, which may account for the substantial difference in sizes of the insertion domains. Whereas for the IS4 family member Tn10 – and therefore also very likely for Tn5 – both hairpin formation and the strand transfer reaction use the same 3'OH group of the transferred strand as a nucleophile (Figure 1f; Kennedy *et al.*, 2000), the first cleavage of the *Hermes* and RAG1 reactions generates an 3'OH nucleophile on the non-transferred strand. This 3'OH group is used to generate the flanking end hairpin yet the nucleophile that is subsequently used for strand transfer is the 3'OH of the transferred strand created during hairpin formation (Figure 1g). The simplest way to imagine how this might happen is if the enzyme active site somehow switches from one DNA strand (that cleaved first) to the other (the transferred strand). This suggests either the need for major conformational rearrangements which are not necessary for the Tn5 transposase, or the use of multiple sites. It is quite possible that these mechanistic differences are reflected in the structural differences displayed by the different insertion domains.

While it is not an insertion domain, a structural motif of 23 amino acids is inserted between strands $\beta 1$ and $\beta 2$ in the standard RNase H-like fold in the *Mos1* transposase. In the uncomplexed three-dimensional structure of *Mos1*, this segment is disordered (Richardson *et al.*, 2009). On the other hand, in the structure of the *Mos1* complex with cleaved transposon Right Ends (Richardson *et al.*, 2009), it becomes ordered and appears to play a crucial role in transposase dimerization, making protein-protein interactions across the dimer interface and important protein-DNA contacts. The transposase dimer seen in the *Mos1*/DNA structure has both 3' ends of two transferred strands in the two active sites, a configuration that appears to be stabilized to a large extent by these protein-protein and protein-DNA interactions. As this configuration captures a state in the reaction where the transferred strands have been cleaved (Figure 1d), the inserted segment therefore contributes to the proper execution of second strand processing, at least indirectly.

Some DNA transposases possess a YREK motif

In addition to the DDE/D motif, another notable sequence motif found in some dsDNA transposases is a Y-(2)-R-(3)-E-(6)-K motif, the so-called "YREK" motif originally identified in the IS4 family (Rezsöházy *et al.*, 1993). This motif is located on $\alpha 4$, where the E of the YREK motif is the same E as that of the catalytic DDE motif. As seen in the structure of the Tn5 transposase bound to DNA, $\alpha 4$ is wedged into the DNA minor groove immediately adjacent to the hairpin (see Figure 4A in Davies *et al.*, 2000). In this location, the Y, R, and K side chains form a number of important contacts with the transposon end. Embedded in the YREK motif, just after R322, is residue W323 which is pushed into the DNA minor groove. It is likely that this interaction is crucial for hairpin formation as it may act to push the base to be flipped out whereas W298 appears to capture this base, probably by base stacking. It appears therefore that hairpin formation during Tn5 transposition is assisted by two Trp residues acting in a "push and pull" configuration (Bischerour and Chalmers, 2007). Interestingly, in the Tn10 transposase (also an IS4 family member) there is instead an M after the R of the YREK motif that may have a similar role in extruding the flipped-out base (Allingham *et al.*, 2001; Bischerour and Chalmers, 2009).

Although it is tempting to generalize the need for two similarly placed Trp residues to all transposition systems that proceed through hairpin intermediates, it may be that the "push and pull" interactions observed for Tn5 system are only needed because the hairpin formed has a very short loop, i.e. the double-stranded break occurs at the very end of the transposon both for Tn5 and for

Tn10. If the resulting hairpin has a longer and therefore presumably more flexible loop (more unpaired bases), flipping out a specific base to make room at the DNA end for the hairpin to form may not be as important as it is for the extremely short loop of Tn5 and Tn10.

While the YREK motif is a feature of the IS4 family and may be indicative of a mechanism that proceeds through a hairpin intermediate on the transposon end (but not on the flanking end), the K/R residue that follows the catalytic E/D by about seven residues is present in many other families that do not possess a YREK motif (Mahillon and Chandler, 1998). This K/R seems likely to be catalytically important, as demonstrated for HIV-1 IN where mutation of K159 (which follows the catalytic E152) leads to a loss of *in vitro* cleavage and strand transfer activities (Jenkins *et al.*, 1997), and for the IS1 transposase, where the similarly located R198A mutation has a serious *in vivo* transposition defect (Ton-Hoang *et al.*, 2004).

RNase H-like DNA transposases are modular proteins

The overall properties of RNase H-like transposases do not rely on the catalytic domain alone. These proteins always carry additional domains such as helix-turn-helix (HTH) or winged-helix domains, zinc-binding domains (mostly with undefined functions), and domains required for multimerization.

While the RNase H-like catalytic domain itself must have some DNA binding activity, this is often non-specific and/or weak. Rather, sequence-specific DNA binding (such as binding transposon or viral DNA ends) is carried out by sequence-specific DNA binding domains of various kinds, typically located upstream of the catalytic domain. In this sense, DDE/D transposases are always modular proteins, with different functions distributed between different domains. The sequence-specific DNA binding function that allows the transposase to locate the ends of its mobile element can reside in a single domain or be distributed among several domains. For example, MuA transposase recognizes its transposon ends using a winged helix DNA binding domain followed by an HTH domain (reviewed in Chaconas and Harshey, 2002; Rice, 2005); members of the *mariner* family use two HTH domains (reviewed in Plasterk *et al.*, 1999; van Pouderooyen *et al.*, 1997; Watkins *et al.*, 2004; Richardson *et al.*, 2009); and OrfAB, the transposase of IS911, uses a single HTH domain (Rousseau *et al.*, 2002).

Among the repertoire of site-specific DNA binding domains, unusual α -helical, non-standard fold domains are sometimes present, as in the case of Tn5 (Davies *et al.*, 2000). The eukaryotic *Hermes* transposase has an

idiosyncratic intertwined helical domain (Hickman *et al.*, 2005), and a similar domain has recently been observed in RAG1 (Yin *et al.*, 2009). Retroviral integrases contain an α -helical Zn²⁺ binding domain upstream from the catalytic domain that is very similar to HTH domains, but its role in site-specific binding of viral ends is not yet clear (Craigie, 2001).

There is even greater diversity in domains downstream of the RNase H-like catalytic domain. Although in some cases, such as many of the *hAT* family eukaryotic DNA transposases, there are no additional domains, when present, C-terminal domains can be involved in a number of functions: non-specific DNA binding, such as binding target DNA (which is generally, but certainly not always, non-specific), or in multimerization or protein-protein interactions with other components of the transpososome. An interesting variation appears with the *IS110* family where the RNase H-like catalytic domain appears at the amino terminus of the transposase and is followed by an all-helical C-terminal domain with no obvious indication as to its function. This unusual arrangement is accompanied by the rather unusual property of strictly sequence-specific integration (Higgins *et al.*, 2009).

RNase H-like DNA transposases need to multimerize

Transposase molecules must form multimeric complexes within the transpososome, and the currently available experimental three-dimensional structures allow us some new insights into how this is achieved. It is remarkable that despite the obvious structural homology of the catalytic domains, the mode of transposase oligomerization is very divergent, as each of the systems for which structural data are available form oligomers in their own unique way. Some DNA transposases such as MuA and Tn5 are monomers when not bound to DNA, and oligomerize upon binding; for instance Tn5 forms a dimer while MuA becomes tetrameric. Others are multimeric on their own, and it is not clear that the multimerization state changes upon DNA binding. For example, there is evidence that the *hAT* superfamily member Hermes is a hexamer even without bound DNA (Hickman *et al.*, 2005) and HIV-1 IN forms tetramers both with and without DNA bound (Bao *et al.*, 2003; Ren *et al.*, 2007; Hare *et al.*, 2009; Michel *et al.*, 2009) as does the purified P element transposase (Tang *et al.*, 2007).

One clear example of a dedicated protein multimerization domain was identified in the *IS3* family where the vast majority of the almost 500 members of this family exhibit a leucine-rich region predicted to form a coiled coil structure or leucine zipper (LZ) (Rousseau *et al.*, 2002). The functionality of this region

was experimentally tested for one family member, *IS911* (Haren *et al.*, 1998, 2000). Site-specific mutagenesis of key LZ residues based on analysis of the *jun/fos* system demonstrated that this region is required for formation of transposition intermediates as judged both by genetic and physical tests *in vivo* and by a standard *in vitro* transposition assay. The inability of these mutants to multimerize was confirmed directly by their inability to undergo co-immunoprecipitation with a tagged wild-type transposase derivative (Haren *et al.*, 1998). These studies also revealed a second region of the protein required for correct multimerization. This region has no particular motifs to suggest how it might function directly in multimerization. Similar leucine-rich regions have been identified in other transposases including the regulatory KP element involved in regulation in P element transposition (Andrews and Gloor, 1995; Lee *et al.*, 1996); *IS1111* (Hoover *et al.*, 1992) and other members of the *IS110* family; the *IS66* family (Gourbeyre *et al.*, personal communication); and Tc1 (Ivics *et al.*, 1996).

An unusual example of a multimerization domain appears in the *Hermes* transposase and RAG1 where an entirely α -helical domain forms a tightly intertwined dimer. This is the same domain that appears to function in site-specific DNA binding (Hickman *et al.*, 2005; Yin *et al.*, 2009). A region of the *hAT* transposases that had been previously implicated in dimerization (Michel *et al.*, 2003), and which has infiltrated the Conserved Domains Database (Marchler-Bauer *et al.*, 2009) as pfam 05699, does not appear to be involved in dimerization; rather, as the structure of *Hermes* revealed, this region meanders through the protein holding adjacent domains together (Hickman *et al.*, 2005), a role that provides an alternative explanation for the original observation that disruption of this region leads to loss of multimerization.

In some cases, the formation of multimers may be the product of protein-protein interactions between several domains, so there may be no single dedicated multimerization domain. For instance, Mos1 uses one of its two N-terminal HTH DNA binding domains and a motif between strands β 1 and β 2 of the catalytic RNase H-like domain to dimerize (Richardson *et al.*, 2009) which also facilitates DNA binding. On the other hand, DNA binding can also contribute to oligomerization, since transposon ends are often bound in "trans" such that a site-specific DNA binding domain of one protomer binds the transposon end that is processed by the catalytic domain of another protomer; in this way, the DNA acts as the "glue" holding (or contributing to holding) the oligomer together. This seems to be the case so far in all three-dimensionally resolved transposase-DNA complexes. For retroviral integrases, the catalytic domain itself is a dimerization domain (Dyda *et al.*, 1994); however, in the formation of the functional tetramer, it is very likely that

other protein–protein and protein–DNA interactions are also involved.

While distinct from transposase multimerization, transposases are sometimes intimately involved through protein–protein interactions with other proteins encoded by the transposon. Well-known examples are bacteriophage Mu and the Tn7 transposon where target immunity, the ability to avoid integrating into a DNA molecule already containing the transposon, is mediated by a transposon-encoded ATPase. This ATPase is a part of the assembled transpososome and interacts directly with the transposase (Baker *et al.*, 1993; Wu and Chaconas, 1994; Craig, 2002; Ronning *et al.*, 2004). V(D)J recombination also relies on a hetero-oligomer between RAG1 and RAG2 for function (Bailin *et al.*, 1999).

Sequence expansion

Over the past few years, the number of prokaryotic and eukaryotic transposons which have been identified has grown tremendously. This is due largely to the extraordinary growth in sequence data deposited in the public databases. For example, there are at present about 1000 completed and publicly deposited prokaryotic genome sequences listed in the GOLD database (Lioliou *et al.*, 2008), the majority of which are poorly annotated for ISs and other mobile genetic elements. Attempts to collate sequences of transposons in a usable form have fallen to a very small number of researchers who have managed to obtain the resources to maintain databases. These include ISfinder (<http://www-is.biotoul.fr>) which provides a list of ISs identified in eubacteria and archaea (Siguier *et al.*, 2006), and Repbase Update (<http://www.girinst.org/repbase/index.html>) which maintains sequences of large families and subfamilies of repetitive elements from eukaryotes (Jurka *et al.*, 2005). A third database, ACLAME (<http://aclame.ulb.ac.be/>) collates a more general collection of prokaryotic mobile genetic elements (Leplae *et al.*, 2004).

The ISfinder database (which is freely consultable and does not require registration) includes over 3000 bacterial ISs. When ISfinder was initiated, ISs were identified from experimental data (e.g. inactivation of a particular gene by insertion) and, while some ISs deposited are still identified in this way, the majority are now defined by homology searches between transposase enzymes and other characteristics of the IS such as genetic organization and the terminal inverted repeats characteristic of transposons with RNase H-like transposases. Thus, in general there is no demonstration that a given IS is active although this can be inferred from those which are present in multiple identical copies in one or several genomes, or by comparison with other closely related members of the family to which they belong. The same

vexing problem with deducing whether an element is active also exists among the collated eukaryotic DNA transposon sequences.

Of prokaryotic ISs, 25 families have been defined in ISfinder (Siguier *et al.*, 2006) and 20 superfamilies of eukaryotic DNA transposons are currently listed on Repbase (Jurka *et al.*, 2005). Recent reviews that provide useful summaries of the properties of prokaryotic and archaea ISs and eukaryotic DNA transposons on a family-by-family basis include those by Mahillon and Chandler (1998) (and its update; see Chandler and Mahillon, 2002), Filée *et al.* (2007), Feschotte and Pritham (2007), and Wicker *et al.* (2007). We note in passing that the designation of “families” (for ISs) as opposed to “superfamilies” (among eukaryotes) appears entirely a matter of convention, rather than a statement regarding “superiority”.

To understand how these families and superfamilies are related to each other – evolutionarily, mechanistically, and structurally – would ideally involve integrating genetic, biochemical, and structural data with results obtained from computational biology, or bioinformatic, approaches. Unfortunately at present, despite the vast amounts of sequence data that are publicly available, only a few transposons and their transposases have been studied experimentally. Nevertheless, the results obtained from classical biochemical and structural approaches serve as the filter through which we are forced to make sense of the sequence data.

Do we now know the folds of the catalytic cores of all DNA transposases?

Of necessity, in the absence of any three-dimensional structural data for most transposition systems, transposase sequence alignments have been the main tool for analyzing these proteins in an effort to determine how they might be related to each other and to transposases of known structure. Using sequence alignments to identify the catalytic DDE/D residues and other conserved amino acids, among the prokaryotic ISs, 17 families have been annotated in ISfinder (Siguier *et al.*, 2006) as containing a transposase with a DDE/D catalytic domain. Among the remaining eight families, the IS607 family possesses a serine transposase, and the IS91 and IS200/IS605 families are members of the HUH superfamily of nucleases, consistent with their known transposition mechanisms which use 5′ phosphotyrosine intermediates. This leaves five families currently unannotated.

Among the 20 superfamilies of eukaryotic transposases identified in Repbase (Jurka *et al.*, 2005), six have been described only in Repbase (*Mirage*, *Rehavkus*, *Nobosib*, *Kolobok*, *ISL2EU*, and *Chapaev* – but see Panchin and Moroz, 2008), making it difficult to objectively assess

their significance. Of these six superfamilies, some are annotated as encoding a transposase with a DDE catalytic core (*Kolobok*, *ISL2EU*, and *Chapaev*) whereas others are annotated as possessing a novel “cut and paste” transposase (for example, *Rehavkus-1_DY*). We eagerly await further description of these transposon superfamilies in order to evaluate the relevance or validity of these assignments.

Of the remaining 14 identified superfamilies, the *Helitrons*, *Cryptons*, and the *Mavericks* can be considered *sui generis*. The *Helitron* transposases (Kapitonov and Jurka, 2001) have Rep domains, suggesting that they are related to the prokaryotic *IS91* family, and thus any active members are expected to transpose using a mechanism related to rolling-circle replication (Koonin and Ilyina, 1993; Mendiola *et al.*, 1994). The few identified *Cryptons* (Goodwin *et al.*, 2003) encode a protein similar to tyrosine recombinases, and therefore they have been proposed to constitute a rather unusual class of LTR retrotransposons (Poulter and Goodwin, 2005). At present it is unclear whether or not these elements are transposons or indeed if they are mobile. The *Mavericks/Polintons* are unusually large elements, generally encoding between four and nine proteins which include a protein-primed DNA polymerase, a retroviral-like integrase, a cysteine protease, and usually an ATPase, suggesting perhaps a novel – and potentially complex – transposition mechanism (Feschotte and Pritham, 2005; Kapitonov and Jurka, 2006; Pritham *et al.*, 2007). Nevertheless, their integrase ORF clearly indicates the presence of an RNase H-like catalytic domain. Although unrelated, a similarly complex organization occurs in the *IS66* (Han *et al.*, 2001) and *Tn7* (Parks and Peters, 2009) families.

Protein sequence alignments have been used in an attempt to place the remaining 11 eukaryotic superfamilies of DNA transposons within the known transposon classification groups. In this way, six superfamilies have been reported to have transposases that are homologous to bacterial IS transposases (summarized in Feschotte and Pritham, 2007; and Table 1): the transposases of the *Tc1/mariners* are related to those of *IS630* (Doak *et al.*, 1994) as are the *Zator* transposases (Bao *et al.*, 2009); *Mutator/MuDr* transposases are related to those of *IS256* (Eisen *et al.*, 1994; Hua-Van and Capy, 2008); *piggyBac* to the *IS4/5* family (Sarkar *et al.*, 2003); *Harbinger/PIF* to *IS5* (Kapitonov and Jurka, 1999; Zhang *et al.*, 2001); *Merlin* to *IS1016* (Feschotte, 2004); leaving only the *En/Spm* (*CACTA*), *hAT*, *P* element, *Transib*, and *Sola* superfamilies without presently identified prokaryotic equivalents. The crystal structures of the catalytic domain of *Hermes*, a representative *hAT* transposase, and of *Mos1*, a *Tc1/mariner* element, established that both possess an RNase H-like catalytic domain (Hickman *et al.*, 2005; Richardson *et al.*, 2006).

How reliable are sequence alignments in predicting structure?

In the absence of structural and biochemical data for the vast majority of IS families and eukaryotic superfamilies, the question arises of how reliably sequence analysis can place all these transposases within the structural universe. Is it reasonable that primary sequence alone has established that the vast majority of dsDNA transposases possess an RNase H-like catalytic domain? Furthermore, where sequence alignments remain ambiguous, how likely is it that all the remaining unannotated ISs and eukaryotic DNA transposases are similarly structurally related?

One concern when relying solely on sequence alignments to identify a catalytic DDE/D motif is that there are many possible reasons why acidic residues might be conserved in proteins and the identification of three “does not necessarily a catalytic site make”. Prior to the deluge of DNA transposon sequences, the limited number of known eukaryotic elements and the lack of information about whether they were active, occasionally proved treacherous in predicting a DDE motif. For example, early efforts to identify the DDE triad in the *hAT* elements (Rubin *et al.*, 2001; Michel *et al.*, 2002) were hampered by the presence of an unrecognized insertion domain, leading investigators to search for relatively closely spaced Ds and Es, rather than allowing for a very large spacing between the D residues and the final E. This led to a brief foray into the possibility of a “DSE” catalytic motif (Bigot *et al.*, 1996) before this provocative proposal was discarded (Zhou *et al.*, 2004).

In other earlier work, efforts to force unexpectedly placed – or simply missing – conserved acidic residues into a DDE motif led to contortions such as the initial “putative noncanonical DDE catalytic site” of the *Transib* transposases (Kapitonov and Jurka, 2003) and the “working alignment of DD(35)E family members” which aligned *TnsA* of the *Tn7* transposon with retroviral integrases and the *MuA* transposase (Sarnovsky *et al.*, 1996). To the credit of the authors of these papers, the clear indications that something was not quite right led quickly to an amended alignment in the first case (Kapitonov and Jurka, 2005) and was the impetus in the second case to experimentally determine the three-dimensional structure of *TnsA* which was unexpectedly found to resemble type II restriction enzymes rather than RNase H, and therefore does not possess a DDE active site (Hickman *et al.*, 2000).

Today, there are many readily available bioinformatic tools that have the potential to bridge the gap between transposase amino acid sequences and protein structure prediction. One level of analysis that can be straightforwardly applied is to determine if the predicted secondary structure is consistent with an RNase

Table 1. List of mobile elements whose transposases have been examined by secondary structure prediction programs.

Family	Element (or protein) analyzed	Active or # copies in genome ¹	From secondary structure, type of DDE/D motif ²	Relevant references ³
IS1	IS1N	>40*	DD(24)E	*Nyman <i>et al.</i> , 1981; Ohta <i>et al.</i> , 2002, 2004;
	ISSto9	5	DD(20)E	Siguier <i>et al.</i> , 2009
IS1595	1. ISPna2	—	DD(36)N	Siguier <i>et al.</i> , 2009
	2. ISH4	—	DD(36)E	Siguier <i>et al.</i> , 2009
	3. IS1016C	—	DD(34)E	Siguier <i>et al.</i> , 2009
	4. IS1595	—	DD(35)N	Siguier <i>et al.</i> , 2009
	5. ISSod11	13	DD(34)H	Siguier <i>et al.</i> , 2009
	6. ISNWi1	—	DD(35)E	Siguier <i>et al.</i> , 2009
	7. ISNha5	—	DD(33)E	Siguier <i>et al.</i> , 2009
	Merlin: MERLIN1_SM	Consensus	DD(36)E	Feschotte, 2004
IS3	IS911	Active	DD(35)E	Polard and Chandler, 1995; Rousseau <i>et al.</i> , 2002
IS481	IS481	~100*	DD(35)E	*Glare <i>et al.</i> , 1990; Chandler and Mahillon, 2002
IS4	IS50R	Active	PDB ID: 1muh DD(β -strand)E	Rezsöházy <i>et al.</i> , 1993; Davies <i>et al.</i> , 2000
IS701	IS701	Active (15*)	DD(β -strand)E	*Mazel <i>et al.</i> , 1991
	ISRso17	7		
ISH3	ISC1359	5	DD(β -strand)E	
	ISC1439A	13		
IS1634	IS1634	Active (~30*)	DD(β -strand)E	*Vilei <i>et al.</i> , 1999
	ISMae5	7		
	ISPlu4	7		
IS5	IS903	Active	DD(65)E	Derbyshire <i>et al.</i> , 1987; Rezsöházy <i>et al.</i> , 1993; Tavakoli <i>et al.</i> , 1997
	PIF/Harbinger: PIFa (<i>Z. mays</i>)	Active	DD(59)E	Zhang <i>et al.</i> , 2001; Kapitonov and Jurka, 2004; Sinzelle <i>et al.</i> , 2008
IS1182	IS660	3	DD(β -strand)E	Takami <i>et al.</i> , 2001
	ISPsy6	14		
IS6	IS6100	Active	DD(34)E	Martin <i>et al.</i> , 1990; Mahillon and Chandler, 1998
IS21	IS21	Active	DD(45)E	Mahillon and Chandler, 1998; Berger and Haas, 2001
IS30	IS30	Active	DD(33)E	Caspers <i>et al.</i> , 1984; Mahillon and Chandler, 1998
IS66	IS679	Active	DD(α -helical?)E	Han <i>et al.</i> , 2001
	ISPsy5	33		
	ISMae8	3		
IS110	IS492	Active	DEDD	Perkins-Balding <i>et al.</i> , 1999;
	IS1111	20	DEDD	Buchner <i>et al.</i> , 2005
IS256	IS256	Active	DD(α -helical)E	Mahillon and Chandler, 1998; Prudhomme <i>et al.</i> , 2002
	MuDr/Foldback (Mutator)	Active	DD(α -helical)E	Eisen <i>et al.</i> , 1994; Babu <i>et al.</i> , 2006; Hua-Van and Capy, 2008
IS630	ISY100	Active	DD(34)E	Doak <i>et al.</i> , 1994; Feng and Colloms, 2007
	Tc1/mariner: Mos1 (<i>D. mauritiana</i>)	Active	PDB ID: 2f7t DD(34)D	Plasterk <i>et al.</i> , 1999; Richardson <i>et al.</i> , 2006
	Zator: Zator-1_HM	36*	DD(43)E	*Bao <i>et al.</i> , 2009
IS982	ISPfu3	5	DD(47)E	Mahillon and Chandler, 1998
IS1380	IS1380A	~100*	DD(β -strand)E	*Takemura <i>et al.</i> , 1991; Chandler and Mahillon, 2002
	piggyBac (<i>T. ni</i>)	Active	DD(β -strand)D	Cary <i>et al.</i> , 1989; Sarkar <i>et al.</i> , 2003; Mitra <i>et al.</i> , 2008
ISAs1	ISAz03	7	DD(β -strand)E/D?	
ISL3	IS31831	Active	DD(α -helical)E	Suzuki <i>et al.</i> , 2006
	IS651	22		
Tn3	Tn3 (<i>E. coli</i>)	Active	DD(α -helical?)E	Grindley, 2002

Table 1. Continued on next page

Table 1. Continued.

Family	Element (or protein) analyzed	Active or # copies in genome ¹	From secondary structure, type of DDE/D motif ²	Relevant references ³
<i>hAT</i>	<i>Hermes</i> (<i>M. domestica</i>)	Active	PDB ID: 2bw3 DD(α -helical)E insertion	Warren <i>et al.</i> , 1994; Rubin <i>et al.</i> , 2001; Hickman <i>et al.</i> , 2005
CACTA	CACTA1 (<i>A. thaliana</i>) En/Spm ZM	Active	DD(α -helical?)E/D?	Miura <i>et al.</i> , 2001; DeMarco <i>et al.</i> , 2006
P	<i>Drosophila</i>	Active	?	Rio, 2002
<i>Transib</i>	<i>Transib1_AG</i> RAG1 (<i>M. musculus</i>)	Consensus Active	DD(α -helical)E DD(α -helical)E	Kapitonov and Jurka, 2005; Chen and Li, 2008 Kim <i>et al.</i> , 1999; Landree <i>et al.</i> , 1999; Lu <i>et al.</i> , 2006
<i>Sola</i>	<i>Sola3-3_HM</i>	Multiple copies*	DD(40)E	*Bao <i>et al.</i> , 2009

¹Information on the number of copies within the host genome was obtained from ISfinder or the reference indicated by the asterisk.

²Where indicated, the secondary structure predicts an insertion domain between $\beta 5$ and $\alpha 4$ with predominantly either β -strands or α -helices.

³Relevant references include reviews or papers that report the results of secondary structure prediction, report sequence alignments or consensus sequences, identify the DDE/D catalytic residues, or demonstrate that the element is active. The association of certain eukaryotic superfamilies to specific IS families is as per Feschotte and Pritham (2005) and references therein.

H-like fold. To ask this question is, in part, to revert to the longstanding problem of protein folding: how well can we predict or calculate the three-dimensional structure of a protein knowing its primary sequence (reviewed in Dill *et al.*, 2007)? Homology modeling is generally not a useful approach to predicting structure if the level of sequence identity is below 20–25% (Cavasotto and Phatak, 2009), and it is clear that most transposases do not have significant sequence identity to those with known structures. At present, the most reliable secondary structure prediction programs are generally considered to be ~80% accurate, and the reliability of protein threading (the prediction of protein structure by a combination of amino acid sequence homology and predicted secondary structure) also continues to improve (Zhang, 2008; Kryshchuk *et al.*, 2009; Qu *et al.*, 2009).

Below, we present an overview of the capacity of these bioinformatic tools to provide information on the structural relationship between different transposases with an identical chemistry but with diverse but related molecular mechanisms. We integrate previously reported results with those using web-based servers that implement secondary structure prediction methods including PSIPRED (Jones, 1999; Bryson *et al.*, 2005) and the Jnet algorithm (Cole *et al.*, 2008), in addition to the pGenTHREADER and pDomTHREADER methods for fold recognition (Lobley *et al.*, 2009). This is applied to representative members of all the IS families either annotated as DDE proteins or unannotated upon ISfinder (Siguier *et al.*, 2006), and nine of the 11 described eukaryotic superfamilies not yet directly demonstrated through structure determination to possess an RNase H-like catalytic core. The representative members were chosen because they are either known to be active or exist in multiple copies (suggesting that they are active); they are shown in Table 1. The analysis

was not performed for the IS4 family, *hAT* transposases, or the Tc1/*mariner* superfamilies as RNase H-like catalytic cores have been demonstrated directly by three-dimensional structure determination; for these, PDB IDs are shown in Table 1. Similarly, the analysis of secondary structure has already been reported for the *Mutator*/*MULE* (“*Mutator*-like elements”) superfamily (Babu *et al.*, 2006; Hua-Van and Capy, 2008); nevertheless, all of these elements are included in Table 1. The assumption is made throughout that, since identifiable sequence homology allows different IS families or eukaryotic superfamilies to be defined, the result for one representative member is highly likely to hold true for all family or superfamily members.

Prokaryotic IS transposases

The predicted secondary structure of representative members of the 17 IS families annotated in ISfinder as DDE/D transposases and of the five currently unannotated families reveals that essentially all have predicted folds consistent with an RNase H-like fold.

The first criterion considered was a conserved order of secondary structure elements with the approximate pattern $\beta 1$ - $\beta 2$ - $\beta 3$ - $\alpha 1$ - $\beta 4$ - $\alpha 2/3$ - $\beta 5$ - $\alpha 4$ - $\alpha 5$. Prediction is, of course, not perfect and not all of these elements are always predicted with high confidence: sometimes they appear unreasonably short, are not predicted at all, or are occasionally interrupted by short bursts of predicted random coil. Nevertheless, the broad outline of the fold can be discerned by visual inspection. This was confirmed by protein threading (Lobley *et al.*, 2009) where, for 19 of the 22 IS families, representative members yielded a match – with a varying degree of certainty – to one or more of the known transposase or retroviral integrase structures in the ranked results (Table 2).

Table 2. Protein threading results for transposases of representative members of IS families and eukaryotic superfamilies. These results are accessible by obtaining the amino acid sequence of the IS transposase from ISfinder (<http://www-is.biotoul.fr>) and submitting it to the PSIPRED server (<http://bioinf.cs.ucl.ac.uk/psipred/>). For the eukaryotic transposases, the sequences were obtained either from RepBase or GenBank. "Rank on List" indicates where in the overall list of threading results the relevant DDE/D protein appeared.

IS/transposon	PDB ID for best relevant threading result	Rank on list	P value (probability of false positive)
IS1 (IS1N)	1exq	#3	1×10^{-3}
IS1595 (IS1595)	1cxq	#1	1×10^{-3}
IS3 (IS911)	1cxq	#1	6×10^{-7}
IS481 (IS481)	1cxq	#1	7×10^{-7}
IS4	Not necessary (see 1mus)	NA	NA
IS701 (IS701)	1mus	#1	2×10^{-7}
ISH3 (ISC1439A)	1mus	#1	2×10^{-4}
IS1634 (IS1634)	1mus	#1	2×10^{-6}
IS5 (IS903)	1cxq	#2	0.002
IS1182 (IS660)	1mus	#3	0.003
IS6 (IS6100)	1cxq	#1	1×10^{-4}
IS21 (IS21)	1exq	#1	1×10^{-4}
IS30 (IS30)	1cxq	#1	1×10^{-5}
IS66 (IS679)	None	NA	NA
IS110 (IS492)	1hjr	#1	3×10^{-4}
IS256 (IS256)	1cxq	#17	0.009
IS630 (ISY100)	1cxq	#3	9×10^{-4}
IS982 (ISPfu3)	1exq	#2	0.003
IS1380 (IS1380A)	1mus	#1	3×10^{-7}
ISAs1 (ISAzo3)	1mus	#1	6×10^{-5}
ISL3 (IS31831)	2bw3	#12	0.007
Tn3 (Tn3; V00613)	None	NA	NA
En/Spm/CACTA (AAC97237)	None	NA	NA
hAT	Not necessary (see 2bw3)	NA	NA
Harbinger/PIF (AF412282)	1bco	#26	0.011
Tc1/mariner	Not necessary (see 2f7t)	NA	NA
Merlin (MERLIN1_SM_1)	1cxq	#1	9×10^{-4}
MuDR/Mutator	Babu <i>et al.</i> , 2006; Hua-Van and Capy, 2008	NA	NA
P element (AAT96009)	None	NA	NA
piggyBac (ABC88680)	1mus	#1	2×10^{-4}
Transib (Transib1_AGp)	None	NA	NA
Sola (Sola3-3_HM)	1cxq	#2	0.002
Zator (Zator-1_HM)	None	NA	NA

NA: not applicable, either because the structure of a representative member has been determined, there were no relevant threading results, or the results have been previously reported. The PDB IDs correspond to the following proteins: 1exq (HIV-1 integrase); 1cxq (ASV integrase); 1mus (Tn5 transposase); 1hjr (RuvC); 2bw3 (Hermes); and 1bco (MuA).

The second criterion required for an RNase H-like fold, dictated by the determined structures, is that the DD of the DDE/D motif must fall on or very close to predicted β 1 and β 4, and the E/D must be on or close to a predicted downstream α -helix. Also supportive was the location of a conserved K/R at a characteristic downstream distance from the catalytic E.

One notable variant in these predictions is whether or not a transposase carries an insertion domain between the second and third conserved acidic residue. On this basis, the IS families can be grouped into those without a predicted insertion domain (IS1, IS3, IS5, IS6, IS21, IS30, IS110, IS481, IS630, IS982, IS1595), those with a largely

α -helical insertion domain (IS256, ISL3, and possibly IS66 and Tn3), and those with a mostly β -strand insertion domain (IS4, IS701, ISH3, IS1634, IS1182, IS1380, ISAs1).

The three IS families that do not yield a straightforward prediction of an RNase H-like catalytic domain are IS110, IS66, and Tn3. For IS110, the predicted fold of the catalytic domain corresponds to that of RuvC, another member of the RNase H superfamily (Ariyoshi *et al.*, 1994), a resemblance has been noted previously (Buchner *et al.*, 2005). Although RuvC has the same topological organization as the RNase H-like transposases, the arrangement of acidic residues, DEDD, is

different with the third and fourth aspartic acid residues appearing one helical turn apart in a C-terminal helix that differs in orientation from either of the C-terminal helices of the RNase H-like fold (reviewed in Yang and Steitz, 1995). For both IS66 and Tn3, the secondary structure prediction results suggest RNase H-like folds with α -helical insertion domains, but this was not supported by the results of protein threading.

A curious observation has recently come to light regarding the IS1595 family whose members can be separated into seven distinct groups (Siguier *et al.*, 2009). Of the seven identified groups (shown in Table 1), only four have the expected DDE triad; members of the other three appear to have either an Asn or His as the conserved third catalytic residue, and two of the three have yet another conserved E further towards the C terminus. For two of the three subfamilies with apparent DDN or DDH motifs, predicted secondary structure places the conserved downstream E on α 5, rather than α 4. As is evident from Figure 2, it seems unlikely that a third catalytic residue located on α 5 would be able to form a two metal ion binding site with residues from β 1 and β 4. It would be extremely revealing to determine the three-dimensional structures of these variant transposases to establish whether they differ from canonical DDE enzymes by assembling an active site using a DDN or DDH motif (discussed in Nowotny, 2009), or if the structure of the catalytic domain changes relative to others to allow a predicted fifth α -helix carrying the canonical E residue to move into the proximity of β 1 and β 4. In this context, it is worth noting that the human SETMAR protein, a fusion between a SET domain with protein methyltransferase activity and a *mariner* transposase (Robertson and Zuppano, 1997), has measurable *in vitro* activity for many of the steps of transposition despite an active site that has a DD(34)N motif (Cordaux *et al.*, 2006; Liu *et al.*, 2007; Miskey *et al.*, 2007).

Eukaryotic DNA transposases

The type of secondary structure analysis outlined above for the prokaryotic IS families has been reported for representative members of several eukaryotic superfamilies including the *hATs* (Zhou *et al.*, 2004) and *piggyBac* (Mitra *et al.*, 2008), and for the *Mutator*/*MULE* superfamily (Babu *et al.*, 2006; Hua-Van and Capy, 2008). For *Hermes* and *piggyBac*, where transposition has been recapitulated *in vitro* with purified transposases, the secondary structure prediction demonstrated that conserved acidic residues that are crucial for activity are located precisely where expected, a result confirmed crystallographically for *Hermes* (Hickman *et al.*, 2005). Analysis of the *Mutator* transposases (Hua-Van and Capy, 2008), which was more extensive than simple

protein secondary structure prediction and protein threading by incorporation of aspects of 3D structure prediction, could serve as a paradigm for further studies on the entire range of eukaryotic transposon superfamilies.

The task of deciding whether a given DNA transposase has a predicted secondary structure consistent with an RNase H-like fold is far less straightforward for eukaryotic elements than for the bacterial ISs. Although in many cases, appropriately placed β -strands relative to a few helices suggest an RNase H-like fold, as shown in Table 2, the results are compelling only for members of the *Harbinger*/*PIF*, *Merlin*, *Mutator*, *piggyBac*, and *Sola* superfamilies.

The predicted secondary structure of *piggyBac* from *Trichoplusia ni* is completely consistent with a fold resembling that of the Tn5 transposase (onto which it can be threaded; Table 2) with several β -strands comprising an insertion domain between β 5 and α 4. The presence of an insertion domain is in accord with the observed large spacing between the second and third identified catalytically essential acidic residues of the DDD motif, D346 and D447 (Mitra *et al.*, 2008). Although the catalytic mechanism of *piggyBac* proceeds through hairpin intermediates on the transposon ends as does the Tn5 system, there is as yet no evidence for two Trp residues partaking in a “push-and-pull” mechanism like Tn5 nor is there a conserved YREK motif. This may well be related to the possibility that a substantially larger loop at the tip of the hairpin is formed on the *piggyBac* ends, requiring less distortion to form than does that of the short loop generated during Tn5 transposition and therefore requiring less assistance from the transposase.

For the *CACTA* transposases, a primary sequence alignment revealed regions of conservation that suggested the possible location of the two aspartate residues of a DDE/D motif (DeMarco *et al.*, 2006). These are indeed located on predicted β -strands that fall in a pattern that strongly suggests a classical RNase H-like fold. Although DeMarco *et al.* (2006) also suggested a possible candidate for the third acidic residue of a DDE/D motif located between the first two Asp residues, it seems far more likely that the *CACTA* transposases possess an α -helical insertion domain and that the final E is located much further towards the C-terminus.

It was proposed some time ago that the V(D)J recombination system, central to the adaptive immune system of jawed vertebrates, likely evolved from an ancient mobile element (Sakano *et al.*, 1979; reviewed in Jones and Gellert, 2004). The obvious conceptual link between DNA transposition and V(D)J recombination was tightened by the observation that the RAG proteins can carry out transpositional strand transfer, leading to 5-bp target site duplications upon insertion

(Hiom *et al.*, 1998). An exciting recent development has been the identification of the *Transib* transposons as the likely ancestors (Kapitonov and Jurka, 2005). Not only is there significant amino acid sequence similarity between the RAG1 core domain and that of the *Transibs*, but the resemblance between *Transib* TIRs and V(D)J recombination signal sequences (RSS) is particularly persuasive.

The secondary structure predicted for the *Transibs* and RAG1 (Kim *et al.*, 1999; Lu *et al.*, 2006) concurs with previous conclusions reached from mutational analysis and sequence alignments (Kim *et al.*, 1999; Landree *et al.*, 1999; Kapitonov and Jurka, 2005) that these proteins likely have catalytic domains with an RNase H-like fold, although this could not be confirmed by protein threading (Table 2). Furthermore, members of the *Transib* superfamily are predicted to have an essentially-all α -helical insertion domain as predicted for RAG1 (Lu *et al.*, 2006). The presence of an insertion domain is consistent with the reported large spacing (206–214 amino acids) among the *Transibs* between the second D and E residues proposed to serve as the catalytic residues (Kapitonov and Jurka, 2005).

One curious conserved sequence feature of the *Transibs* and RAG1 is a CxxC motif (RAG1 residues 727–730 in *Mus musculus*). In one alignment of the insertion domains (Kapitonov and Jurka, 2005), these would occur immediately following $\beta 5$ of the RNase H-like fold. Curiously, a conserved and similarly placed CxxH sequence also occurs in the *hAT* transposases (Zhou *et al.*, 2004); the Hermes structure shows that these residues indeed follow $\beta 5$ and are buried amino acids, at least in the apo-structure (Hickman *et al.*, 2005). A CxxC motif is found in essentially the same place in the predicted RNase H-like fold in *CACTA* transposases (DeMarco *et al.*, 2006) and a CxxH motif in the *MULE* transposases, both of which are predicted to have large α -helical insertion domains (Babu *et al.*, 2006; Hua-Van and Capy, 2008). (In the “rough alignment” of Lu *et al.* (2006), the RAG1 CxxC motif is aligned slightly differently, placing these residues between predicted strands $\beta 4$ and $\beta 5$ of the catalytic core.)

The RAG1 CxxC motif has been proposed to be part of a C2H2 zinc finger (Rodgers *et al.*, 1996), and would represent an intriguing structural insertion-within-an-insertion. For the *Transib* and *hAT* transposases, nearby conserved histidine residues have not been identified, suggesting that their CxxC/H residues might not participate in assembly of a similar Zn^{2+} -binding site.

For transposases of the newly identified *Sola* and *Zator* superfamilies (Bao *et al.*, 2009), the predicted catalytic core secondary structures are consistent with the proposal that these are DDE/D transposases. Among the three groups of *Sola* transposases, the *Sola3* group bears some limited sequence resemblance to

piggyBac transposases (Bao *et al.*, 2009). However, *Sola3* transposases do not have a predicted insertion domain between the identified second and third conserved acidic residue, and the alignment presented in Bao *et al.* (2009) aligns the third D of the *Sola3* DDD motif with a D in *piggyBac* that was not one identified as a catalytic residue by mutational analysis (Mitra *et al.*, 2008). The curious parallel that *Sola3* elements integrate specifically into TTAA target sites as does *piggyBac* (Cary *et al.*, 1989) suggests that the two types of transposases have found different structural solutions to achieve the same outcome, at least as they relate to the catalytic domain.

Finally, the structural aspects of the P element superfamily transposases have been, and remain, a mystery and secondary structure analysis sheds no new light on the issue. The P element transposases have always stood slightly apart from other DNA transposases due to unusual features of transposition such as cuts at the end of the transferred strand and within the non-transferred strand that are staggered by 17 bases, and a requirement for GTP (reviewed in Rio, 2002). Acidic residues that have been determined to be catalytically important are spaced as D(83)D(2)E(13)D, which – regardless of how they are arranged – cannot conform to the expected location of catalytic residues on an RNase H fold.

It appears probable that the eukaryotic DNA transposase superfamilies, like those of the prokaryotic transposons, can be grouped into those without an insertion domain (Tc1/*mariner*, *Merlin*, *Sola*, *Zator*, *Harbinger*/*PIF*), those with a largely α -helical insertion domain (*hAT*, *Mutator*, *Transib*, and probably *CACTA*), and those with a mostly β -strand insertion domain (*piggyBac*); the protein fold of the catalytic domain of the P element transposases remains unknown.

Conclusion

An exciting consequence of studies involving DNA transposons has been their development as tools for manipulating and modifying genomes. The application of prokaryotic transposons to genetically modify prokaryotic organisms is a well-developed technology. Startling advances have recently been made in higher organisms using eukaryotic transposons such as *Sleeping Beauty*, a resurrected salmonid Tc1/*mariner* element (Ivics *et al.*, 1997; 2009); *piggyBac* from the moth *Trichoplusia ni* (Cary *et al.*, 1989; Wu *et al.*, 2006); *Mos1*, a Tc1/*mariner* element from *Drosophila mauritiana* (Medhora *et al.*, 1991); and *Tol2*, a *hAT* transposon from the medaka fish (Koga *et al.*, 1996; Kawakami, 2007). Just a few recent examples of the types of applications include using *piggyBac* to introduce transcription factors into mouse and human stem cells to convert them to a

pluripotent state – and then to subsequently excise the exogenous genes (Woltjen *et al.*, 2009); using *Sleeping Beauty* in a genomic screen to randomly disrupt genes in mice to identify genes and pathways responsible for cancer induction (Dupuy *et al.*, 2005); and cleverly exploiting the tendency of *Sleeping Beauty* to hop locally to investigate regions (within 1.5 Mb of the donor site) that may have cis-regulatory effects on nearby genes (Kokubu *et al.*, 2009).

It is astonishing to realize that, in spite of much effort in obtaining high resolution three-dimensional structures of prokaryotic DNA transposases likely to have RNase H-like folds, that only the Tn5 transposase and the catalytic subdomain of that of bacteriophage MuA have been solved. Similarly, among the eukaryotic transposases, there are only two structures that confirm that representative members of the Tc1/*mariner* family and the *hAT* transposases are members of this larger RNase H-like structural superfamily.

The notion that all the IS families – with the exceptions of IS607, IS91 and IS200/IS605 – are likely on the basis of secondary structure prediction to have RNase H-like folds is not a particularly surprising conclusion. It is, perhaps, precisely what was to be expected. However, it would be fascinating to determine if the predicted presence – and type – of insertion domain correlates with a particular mechanistic aspect of transposition. To date, though, for many of these IS families the necessary biological information (*in vivo* and *in vitro*) is missing.

Among the eukaryotic DNA transposases, secondary structure prediction appears to support the notion that essentially all of the nine superfamilies discussed above do possess catalytic domains with RNase H-like folds. Nevertheless, in the absence of biochemical information on the catalytic relevance of conserved acidic residues – or even data indicating which elements within some superfamilies are active – the evidence is more compelling for some than for others. The jury remains out on the P element. Is it truly the structural outlier, as was eventually established for TnsA of the Tn7 transposon, or does an RNase H-like fold lie somehow encrypted within its primary sequence?

As shown in Figure 1, the known pathways by which dsDNA transposons move from one location to another are wonderfully varied. So are the ever-expanding families and superfamilies of IS and transposon sequences. To conceptually link primary sequence to function will clearly require more experimental structural data; without this, the roles of identifiable transposase features such as insertion domains or multimerization domains or conserved sequence motifs remain a matter of speculation, conjecture, and imagination. In turn, the ability to determine high-resolution crystal structures of transposases and their complexes with DNA rests on the shoulders of *in vitro* and *in vivo* studies on these myriad transposition

systems. In many cases, obtaining experimental results has been difficult due to the often inherently recalcitrant biochemical and biophysical properties of the transposases themselves. One hopes that the availability of vastly expanded sequence information will yield candidates whose experimental study will become possible. The challenges await!

Acknowledgements

We thank Patricia Siguier for supplying alignments of the IS1595 subfamilies and Julia Richardson for graciously providing information on the Mos1/DNA complex prior to publication. We also thank Bob Craigie and Andrea Regier Voth for helpful comments on the manuscript. At the NIH, this work was supported by the Intramural Program of the National Institute of Diabetes and Digestive and Kidney Diseases. In France, this work was supported by intramural funding from the CNRS and by ANR grant MOBIGEN both to M.C.

Declaration of interest

The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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Editor: Michael M. Cox