Characteristics of MuA transposase-catalyzed processing of model transposon end DNA hairpin substrates

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ABSTRACT

Bacteriophage Mu uses non-replicative transposition for integration into the host’s chromosome and replicative transposition for phage propagation. Biochemical and structural comparisons together with evolutionary considerations suggest that the Mu transposition machinery might share functional similarities with machineries of the systems that are known to employ a hairpin intermediate during the catalytic steps of transposition. Model transposon end DNA hairpin substrates were used in a minimal-component in vitro system to study their proficiency to promote Mu transpososome assembly and subsequent MuA-catalyzed chemical reactions leading to the strand transfer product. MuA indeed was able to assemble hairpin substrates into a catalytically competent transpososome, open the hairpin ends and accurately join the opened ends to the target DNA. The hairpin opening and transposon end cleavage reactions had identical metal ion preferences, indicating similar conformations within the catalytic center for these reactions. Hairpin length influenced transpososome assembly as well as catalysis: longer loops were more efficient in these respects. In general, MuA’s proficiency to utilize different types of hairpin substrates indicates a certain degree of flexibility within the transposition machinery core. Overall, the results suggest that non-replicative and replicative transposition systems may structurally and evolutionarily be more closely linked than anticipated previously.

INTRODUCTION

A multitude of mobile genetic elements use the reactions of transpositional DNA recombination for movement within and between the genomes of their host organisms (1). Depending on small differences in the details of the reaction mechanisms, dictated primarily by the structure–function characteristics and regulation of the responsible machineries, the outcome of DNA transposition can be either replicative or non-replicative by nature (2). Of particular importance are the ways the element-encoded transposase or integrase proteins catalyze DNA cleavage and joining reactions (3,4).

Transposases constitute a group of structurally and functionally related proteins with distinct domains responsible for separate subfunctions (5,6). The most important units include a catalytic core and a DNA-binding domain responsible for transposon end recognition. Other domains may provide functions for interactions with auxiliary proteins and accessory DNA sites. Transposases function as a multimer within the context of a specific protein–DNA complex, a transpososome and catalyze two common reactions that unite transposition pathways: (i) a pair of initial cleavages at the transposon ends (typically at the 3′ ends but in some systems at the 5′ ends, e.g. see Hermes transposition below) and (ii) a pair of strand transfer reactions, in which the liberated transposon 3′ ends are joined to the target DNA (7). At least in the two known cases, Mu and Tn5, where assembly of the transpososome is a prerequisite for the chemical steps, the catalysis occurs in trans: i.e. a transposase protomer that is bound to one transposon end via its transposon end-binding domain catalyzes cleavage and strand transfer at the other end of the transposon (8–13).

A majority of transposases belong to a family of DDE transposases and retroviral integrases, referring to an active site that contains a triad of catalytically important amino acids, presumably involved in the coordination of divalent metal ions (6).

Transposing bacteriophage Mu is a widely studied model for DNA transposition in general (14). During the lytic phage propagation, Mu replicates via replicative transposition (15), and this pathway is relatively well characterized (14). Conversely, the outcome of the reaction pathway that leads to the initial Mu integration into the host’s chromosome (following phage infection) is distinctively non-replicative
(16–19), but details of this latter pathway remain to be elucidated.

In general, Mu transposition proceeds within a protein–DNA complex called the Mu transpososome (13,14,20,21). This functional unit forms via an elaborate assembly pathway and synapses the transposon ends. In its core, it contains a tetramer of the critical catalytic protein, MuA (75 kDa monomer), a well-characterized member of the DDE-transposase/integrase family. A number of studies [reviewed by (14)] indicate that the reactions catalyzed by MuA lead to a branched DNA structure, also known as the Shapiro intermediate, that subsequently is processed by host-encoded replication or repair factors, leading to a cointegrate formation or simple insertion, respectively. In both cases, a 5 bp target site duplication is generated. In addition to MuA, a number of cofactors and certain DNA topology are important for Mu propagation in vivo [reviewed by (14)]. The cofactors include specific MuA-binding sites within the phage genome as well as the Mu-encoded target selector protein, MuB. Important host-encoded functions are provided by DNA bending proteins, HU and IHF, as well as certain protein remodeling and DNA replication factors.

Biochemical properties of MuA can be studied effectively by a minimal-component in vitro transposition reaction that includes MuA and a short Mu end-specific DNA segment as the only macromolecular components (22). The reaction faithfully reproduces transpososome assembly, donor cleavage and strand transfer steps; and it has been used for detailed analyses of DNA substrates with variable configuration or sequence (22–27).

Studies of transposon Tn10 and Tn5 systems have revealed a mechanism that uses a transposon end DNA hairpin intermediate during non-replicative transposition (28,29). Between the 3′ end cleavage and strand transfer, the transposases of Tn10 and Tn5 catalyze two additional reaction steps: hairpin formation via the attack of the transposon end 3′-OH group on the opposite strand of the duplex and hairpin opening by hydrolysis that regenerates the 3′-OH residue. Hairpinning of DNA has also been described in related systems such as V(D)J recombination and Hermes transposition, but in these cases the hairpins are formed to seal the flanking DNA ends (30,31).

The evolutionary history of transposable elements must be reflected in their encoded transposition machineries. Yet, although distinct similarities unite a variety of DNA transposition systems, the structural and functional relationships among different elements that use different strategies are unclear. In particular, it is not known when and how different transposases have adapted or lost the catalytic potential for DNA hairpinning. In principle, it might be possible to detect transposases in transition, i.e. retained or potentially emerging activities might be revealed by a detailed analysis. We addressed these issues directly by assembling Mu transpososomes with preformed model hairpin substrates and analyzed the catalytic activities of the ensuing complexes. The Mu transpososome was proficient to accommodate and process a variety of different hairpin substrates, indicating a degree of flexibility within the catalytic machinery. Such flexibility within the active site may be a general determinant that influences or has influenced the choice between the transposition pathways among a variety of transposons.

**MATERIALS AND METHODS**

**Bacteria, proteins and reagents**

*Escherichia coli* strain DH5α (Invitrogen) was used as a cloning host and for plasmid propagation. Bacteria were grown in Luria–Bertani (LB) broth or on LB agar plates (32). To select for plasmid maintenance, the media were supplemented with appropriate antibiotics: ampicillin (Ap, 100 µg/ml) and chloramphenicol (Cm, 10 µg/ml). T4 DNA polymerase was from Promega. T4 polynucleotide kinase (PNK) and T4 DNA ligase were from New England Biolabs. Deoxynucleotides and Dynazyme EXT DNA polymerase preparation were from Finnzymes (Espoo, Finland). BSA was from Sigma, ATP from Roche, and [γ-33P]ATP (1000–3000 Ci/mmol) from GE Healthcare. The commercial enzymes were used under recommended reaction conditions. MuA protein variants were overexpressed and purified as described elsewhere (33). MuB protein was a gift from Dr Kiyoshi Mizuuchi (National Institutes of Health, Bethesda). This protein preparation was made according to Chaconas et al. (34) with the additional step to remove aggregated protein, as described by Adzuma and Mizuuchi (35).

**Oligonucleotides, DNA and DNA techniques**

The oligonucleotides (Supplementary Data) were obtained from commercial sources and purified by urea–PAGE (32) prior to use. When required, the purified oligonucleotides were labeled at their 5′ end with T4 PNK and [γ-33P]ATP (32). Unincorporated nucleotides were removed by sequential phenol and chloroform extractions, passage through a spin column (Bio-Spin 30; Bio-Rad), and ethanol precipitation. To generate double-stranded donor DNA substrates for in vitro transposition reactions, appropriate oligonucleotides were annealed (intraspecifically and/or intermolecularly, see Figure 1) in TEN-buffer (10 mM Tris–HCl, pH 7.5, 0.5 mM EDTA and 50 mM NaCl) as described elsewhere (22). The specific activities of donor DNA substrates were scaled to an equal level of radioactivity by adjustment with corresponding unlabeled substrates. Plasmid DNA was prepared and PCR products were purified using appropriate Qiagen kits. The transposition target plasmid pUC19 (New England Biolabs) was additionally purified by CsCl gradient centrifugation (32) to enrich supercoiled plasmid forms. Standard DNA techniques were performed as described elsewhere (32). DNA sequencing was done at the sequencing service unit of the Institute of Biotechnology, University of Helsinki.

**In vitro transposition reaction and analysis of reaction products**

The standard in vitro transposition reaction was performed in three stages. Initially, transpososomes were assembled for 1 h at 30°C in the absence of target DNA and divalent metal ions (a sample was withdrawn at this stage for the analysis of protein–DNA complexes, see below). Next, target DNA (1 µl of 220 ng/µl stock) was added to the reaction mixture (23 µl) and the incubation was continued for 10 min to allow target capture by transpososomes. To activate catalysis, MgCl₂ (1 µl of 250 mM stock) was then
Figure 1. Donor DNA substrates. (A) Substrate schematic representation. All the Mu-specific substrates contain ~50 bp, starting from the Mu R-end and including the MuA-binding sites R1 and R2 (22). These substrates also include a flanking DNA region, which can be in duplex DNA, linear single strand or in a hairpin loop configuration. Some substrates contain nucleotide substitutions close to the transposon end or within the loop region [see (B)]. The depicted 16 bp flanking sequence has been shown earlier to support efficient catalysis by MuA in an in vitro model substrate assay (22), and the standard loop sequences are derived from that same sequence by joining the endmost Mu-specific 3'-adenine to various nucleotide positions in the opposite strand of the duplex. The Mu end cleavage point is indicated by a solid arrow and the position of the radioactive label by an asterisk. The open arrow shows the position of the nick present in most substrates [see (B)]. (B) Substrate characteristics. The donor substrates were made by annealing one or two oligonucleotides to form double-stranded species as indicated. Most of the hairpin substrates were generated from two oligonucleotides, and these substrates therefore contain a nick [within the R1 MuA-binding site, see (A)]. This nick does not interfere with in vitro transposition reactions as indicated by a direct comparison of reactions with the nicked and corresponding unnicked substrate (compare lanes 4 and 12 in Figure 5). The endmost nucleotides of the transposon DNA are shown in bold, and flanking nucleotides are indicated in regular font. The rHP4 substrate includes a randomly chosen non-Mu sequence with two conserved base pairs mimicking the transposon end.
MgCl₂ was substituted in certain reactions with an equivalent concentration of CaCl₂ or MnCl₂. When indicated, reactions also contained 0.3 μg MuB and 2 mM ATP, both added following a 60 min assembly step, 10 min prior to target addition. In metal ion analysis, MgCl₂ was substituted in certain reactions with an equivalent concentration of CaCl₂ or MnCl₂.

The detection of protein–DNA complexes has been described in detail elsewhere (22). Briefly, competitor DNA to trap loosely bound MuA and 0.2 vol of Ficoll (25%, Pharmacia) were added to the samples prior to gel loading and analysis on a native 2% Metaphor agarose (Cambrex) gel. To detect strand transfer products involving a plasmid target, 0.2 vol of loading dye (0.1% bromophenol blue, 2.5% SDS, 50 mM EDTA and 25% Ficoll 400) was added to the samples prior to gel loading and analysis on a 1.5% LE agarose gel (Promega) in 1× TAE buffer (32) at 5.3 V/cm for 2 h. Agarose gels were dried onto DEAE (Whatman DE-81) paper for autoradiography. Hairpin opening products were analyzed by denaturing 7 M urea–10% PAGE and autoradiography as described elsewhere (36).

Analysis of donor–target junction sequences of transposition reaction products

A 3-fold scaled up transposition reaction was performed with a 3 h final incubation time for catalysis, using HP7 as a donor DNA fragment and pUC19 as a target DNA. The reaction was diluted 1:20 with water, and 1 μl of the diluted mixture was then used as a template for PCR amplification. In the PCR strategy used, the exposed target plasmid 3′-OH groups within the transposition reaction product are initially used for elongation towards the transposon DNA. Following this initial phase, a single transposon-specific primer amplifies a duplex that contains a stretch of transposon DNA attached to each end of the linearized pUC19 target. Thirty cycles of PCR were performed with Dynazyme EXT DNA polymerase as specified by the supplier, using the Mu end-specific primer HSP6. The resulting 2.8 kb PCR product was purified, treated with T4 DNA polymerase in the presence of dNTPs to generate blunt ends (32), and ligated into a DNA fragment containing a selectable marker gene for chloramphenicol resistance. This fragment, the 1.1 kb BamHI fragment of cat-Mu artificial transposon (37), was also treated with T4 DNA polymerase and dNTPs prior to ligation. Ligation products were electrotroduced (38) into DH5α cells, and bacteria were selected for antibiotic resistance on LB-Ap-Cm plates. Plasmid DNA was isolated from 11 clones, and the target sequences flanking transposon DNA were determined using the marker fragment-specific primers HSP350 and HSP349.

RESULTS

MuA catalyzes hairpin processing

DNA hairpin processing by a transposase can be studied in vitro by assembling transpososomes with synthetic DNA model substrates that contain hairpin ends and critical transposase binding sites, as shown for Tn10 and Tn5 (28,29). To elucidate whether the Mu transposition machinery is able to accommodate and process analogous hairpin molecules, we generated a number of ~50 bp Mu R-end model substrates (Figure 1) and used them as a radiolabeled donor DNA in an in vitro transposition assay, the plasmid pUC19 serving as a target DNA (Materials and Methods). Following dissociation of transpososomes by SDS-containing loading buffer, products of this coupled donor cleavage and strand transfer reaction were analyzed by agarose gel electrophoresis and visualized by autoradiography (Figure 2). Two species of reaction products involving the target plasmid are expected in this assay (Figure 2A): the double-ended reaction product (DEP) and the single-ended reaction product (SEP), resulting from the hairpin opening reaction. A third strand transfer product (SEP) was also observed in reactions involving MuA and MuB (lower panel) for 1–6 h. The strand transfer reaction products (SEPs and DEPs) were analyzed by agarose gel electrophoresis and autoradiography. Hairpin opening products were analyzed by denaturing 7 M urea–10% PAGE and autoradiography as described elsewhere (36).

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from utilization of one or two donor molecules, respectively (23–27).

Initially, we analyzed MuA-catalyzed hairpin processing using the substrate HP4, which contains four unpaired hairpin nucleotides (Figure 2B). The precut substrate PC4 served as a positive control for identifiable reaction products. We also included MuB in this experiment due to its stimulatory effect(s) in various types of in vitro assays ([14,23,26,39] and references therein). The hairpin substrate generated the expected reaction product species (SEPs and DEPs) but much less efficiently than the precut control substrate. In addition, the control substrate yielded predominantly DEPs, whereas the hairpin donor produced more SEPs than DEPs. The amount of reaction products did not increase after 1 h incubation, suggesting that the assembly of complexes probably was a limiting factor in the reaction. Addition of MuB to the hairpin substrate reaction increased formation of reaction products to some extent. With the hairpin substrate (data not shown), no reaction products were observed in the absence of MuA, or if the wild-type MuA was replaced by an active-site mutant MuA<sub>ε39Q</sub> (33) defective in catalysis. These data indicate that MuA can catalyze a DNA hairpin cleavage reaction and the subsequent strand transfer step of transposition.

Influence of divalent metal ions
In Mu transposition, donor DNA cleavage and strand transfer require the presence of an appropriate divalent metal ion. While Mg<sup>2+</sup> and Mn<sup>2+</sup> can support both the cleavage and strand transfer reactions, Ca<sup>2+</sup> can support only strand transfer (22). To investigate the divalent metal ion requirements of MuA-catalyzed hairpin processing, we performed transposition reactions with the hairpin donor DNA substrate HP4 in the presence of Mg<sup>2+</sup>, Mn<sup>2+</sup> or Ca<sup>2+</sup> (Figure 3). Control reactions included an uncleaved substrate (UC16) as well as two precleaved substrates with different transposon end configurations: a 4 nt 5'-overhang (PC4) and a blunt end (PC0). All three divalent metal ions supported strand transfer with the precut donors, and the uncleaved donor generated reaction products in the presence of Mg<sup>2+</sup> and Mn<sup>2+</sup>. With the hairpin substrate, detectable amounts of reaction products were generated with Mg<sup>2+</sup> and Mn<sup>2+</sup> but, similar to uncleaved donor, no products were generated with Ca<sup>2+</sup>, suggesting that the hairpin opening stage of hairpin processing probably mimics the donor cleavage step. With the hairpin substrate, Mn<sup>2+</sup> generated more DEPs than SEPs, whereas Mg<sup>2+</sup> produced mainly SEPs (also see Discussion).

Hairpin processing takes place within the Mu transpososome
To study whether hairpin processing takes place within the Mu transpososome, we generated mixed transpososomes (Figure 4A) that contained a labeled hairpin donor DNA (HP4) in combination with unlabeled precut donor DNA (PC4). In these mixed complexes, the precut transposon end is readily available for strand transfer, whereas the hairpin end needs to be opened prior to strand transfer. If both precut and hairpin ends are utilized within a context of a single mixed complex, we should see the relative increase in the generation of labeled DEPs over SEPs in an experiment where increasingly more unlabeled precleaved substrate is used to substitute the labeled hairpin substrate. Analysis of reaction products from such an experiment verified this prediction: the relative amount of labeled DEPs increased upon substitution of increasing amounts of unlabeled precut donor DNA (Figure 4B), indicating that critical reactions involved in hairpin processing must take place within the Mu transpososome.

Effects of hairpin loop length
The length of the transposon flanking DNA within the donor DNA segment, as well as its end configuration, are critical variables determining the assembly, stability and activity characteristics of Mu transpososomes (22). The effect of hairpin loop length and sequence on hairpin processing was therefore investigated (Figure 5). Initially, formation of protein–DNA complexes was analyzed by agarose gel electrophoresis and autoradiography following a 1 h incubation in the absence of divalent metal ions. Under the electrophoretic conditions used, Mu transpososomes can be visualized as stable complexes that withstand a challenge by competitor DNA included in the loading buffer (22). However, depending on the exact reaction conditions and transposon end configuration, certain transpososomes are characteristically less stable and may not be detected by this assay (22). In the latter cases, other types of complexes may be revealed, and they may represent assembly or disassembly intermediates (22). Those hairpin substrates that contained the longest loops primarily yielded complexes that migrated in the gel similarly as the control transpososomes made with precut substrates (Figure 5A, indicated by C1). Upon shortening the loop length, this complex gradually disappeared and there was a simultaneous appearance of another complex having faster gel mobility (indicated by C2). With

Figure 3. Utilization of various metal ions for the catalysis of hairpin processing. In vitro transposition reactions were performed with the indicated substrates by first assembling complexes in the absence of metal ions and then initiating reaction chemistry by adding MgCl<sub>2</sub>, MnCl<sub>2</sub> or CaCl<sub>2</sub> (Materials and Methods). The reaction products, SEPs and DEPs, were analyzed by agarose gel electrophoresis and autoradiography following a 3 h incubation in the presence of metal ion.
the hairpin loop lengths below 4 nt, essentially all of the
observed complexes were of the latter type. Mutated
nucleotides in the transposon end somewhat influenced the
distribution of the complexes detected (compare lanes 4 and
9 as well as lanes 5 and 10), and the non-Mu hairpin substrate
(lane 13) yielded no detectable complexes. The effect of loop
sequence was studied with 7 nt loop substrates, and no
products were detected with those hairpin substrates that contained a relatively large unpaired loop region
(HP7 and HP13A), indicating cleavage at the exact 3’ end of
the transposon. Notably, the same two substrates generated
C1-type protein–DNA complexes (Figure 5A) and yielded rea-
sonable amounts of DEPs (Figure 5B). The cleavage product
was detectable already after 10 min of incubation, and the
yield did not increase appreciably with longer incubation
(Figure 6B). Thus, these data are consistent with the scenario
where the DNA hairpin is opened at the transposon 3’ end
and the end is then used for strand transfer.

To verify that the hairpins are joined to the target DNA,
exactly at the 3’-OH end of the transposon DNA and that
the target site duplication is generated by MuA during the
two reactions involved in hairpin processing, we sequenced
transpon–target junctions from several independent mole-
ules representing transposition reaction products. For this
experiment, we used a 3-fold scaled up transposition reaction
with HP7 donor, pUC19 serving as the target, because such a
reaction generated appreciable amounts of DEPs (target site
duplication can be analyzed from DEPs only). The reaction
products were amplified with a transposon-specific primer
and individual reaction products were cloned for sequence
analysis (Materials and Methods). All 11 clones sequenced
(Table 1) contained a 5 bp target site duplication and, in
each case, the ends of the transposon DNA were accurately
joined to the pUC19 target plasmid. Accordingly, the data
verified genuine two-ended integration upon transposition
as well as utilization of authentic Mu transposition chemistry
with the hallmark 5 bp target site duplication.

**DISCUSSION**

The transposition reaction mechanisms of non-replicative transposons Tn10 and Tn5 involve a DNA hairpin (28,29),
which evidently is reflected on the proficiency of their respective
transposition machineries to accommodate and process model hairpin substrates. A high degree of unity in DNA
transposition reactions in general (3), the fact that Mu can transpose non-replicatively (19), and the previously discov-
ered flexibility in MuA-catalyzed reactions (24,25,27) prompted us to investigate whether the Mu machinery also
could accommodate and process transposon DNA end hair-
pins. We used a minimal-component in vitro assay system
that enabled us to monitor the successful assembly of stable
protein–DNA complexes and detect identifiable hairpin pro-
cessing products.
Figure 5. Effect of hairpin loop length and sequence on stable complex formation and catalysis. (A) Complex formation. Complexes with hairpin (3-fold scaled up reactions) and precleaved (1-fold reactions) substrates were assembled for 1 h in the absence of metal ions, and the reaction products were analyzed by native agarose gel electrophoresis and autoradiography. Primarily, the assay detects transpososomes but may also detect assembly and/or disassembly intermediates. The slower migrating complexes (indicated by C1) include transpososomes. (B) Analysis of strand transfer products. Target DNA (pUC19) and MgCl₂ were added to the assembly reactions to allow catalysis (Materials and Methods), and the reactions were incubated for 6 h prior to analysis of strand transfer products (as in Figure 4). (C) Tabulation of data from (A) and (B). Only a minor portion of the substrates were converted to reaction products (in most cases <1%). The levels of the detected products are based on visual inspection and indicated by a scale of four categories: (−), no products; (+), low level; (++), medium level; (+++), high level. Note that the HP4 substrate did not produce appreciable amounts of DEPs in this experiment, whereas in a similar experiment seen in Figure 2B it did. This apparent discrepancy is due to lower level of radioactivity in the substrate.
Most of the hairpins detected in the Tn10 and Tn5 systems form via an accurate joining of the exact 3’ and 5’ ends of the transposon DNA (28,29). In the Tn5 system, however, a fraction of hairpins were imprecise [i.e. a connection from the 3’ end into the –1 position in the non-transferred strand (26)], indicating that the hairpin configuration within a particular system may vary. As even more extended variability may exist among different systems, it was difficult to make a priori predictions of potentially suitable hairpin substrates for the Mu machinery. Therefore, we generated several types (i.e. variable loop length and sequence) of substrates for the studies. Our initial experiments utilized an imprecise substrate with a 4 nt loop because: (i) such a loop resembles non-complementary nucleotides in a frayed substrate shown to be efficient in promoting the assembly of active transpososomes (22); and (ii) several unpaired loop nucleotides should provide conformational flexibility in DNA close to the active site, a feature that is important for the assembly, stability and activity of the Mu transpososome (22,23,26,40).

Our results show that MuA can open DNA hairpins and that the opened ends are subsequently used for strand transfer. No reaction products were generated without MuA or in the presence of an active-site mutant MuA<sub>E392Q</sub>, indicating that the appearance of SEPs and DEPs is dependent on the presence of a catalytically active MuA and suggesting that the hairpin processing reactions utilize the same active site that is used for the donor DNA cleavage and strand transfer steps. MuB was not required for hairpin processing, but its presence increased the amount of reaction products generated. The reason for this effect is currently unknown but may involve enhancement of transpososome assembly, increased stability of the transpososome, or stimulation of catalysis.

Products of hairpin processing reactions were generated in the presence of Mg<sup>2+</sup> or Mn<sup>2+</sup>, but not with Ca<sup>2+</sup>. As the strand transfer step with precleaved substrates proceeds under all three conditions, the difference must reflect the characteristics of the hairpin opening reaction. At least with respect to ion utilization, the hairpin opening by MuA mimics the donor cleavage reaction. As Mn<sup>2+</sup> rescues certain defects in MuA transposase (33,41,42) and relaxes requirements for donor substrates (39), it was of interest to study whether Mn<sup>2+</sup> would have a similar influence on hairpin processing. In the presence of Mg<sup>2+</sup>, the strand transfer products generated were predominantly SEPs, and a minor fraction of DEPs could only be detected in experiments that utilized substrates with a high specific radioactivity (e.g. Figure 2). Reactions with Mn<sup>2+</sup> yielded mainly DEPs, which suggests that Mn<sup>2+</sup> stimulates coordination between the reactions at each transposon end.

The results with mixed complexes indicate that chemical reactions involved in hairpin processing take place within the Mu transpososome. These data are consistent with the known behavior of MuA: the protein monomers are inert prior to tetramerization into an active transpososome (43), and the active sites within the Mu transpososome are formed by combining structural elements from two different MuA protomers (13). Non-Mu hairpins did not generate stable complexes, and these substrates were not processed, thus providing additional evidence for catalysis within the transpososome. Furthermore, formation of complexes that migrated similarly to known transpososomes in control experiments correlated with the generation of strand transfer products. The length of the hairpin loop influenced the formation of catalytically competent complexes, with longer loops being more productive than those containing only a few unpaired nucleotides. Supposedly, several of the critical contacts required for the transpososome assembly and stability are more optimally positioned with the longer loops. In addition, longer loops may allow more flexibility in the DNA at the vicinity of the endmost transposon nucleotides, thereby facilitating the proper conformation of the active site. The
catalytically productive conformation in the Mu transpososome entails a criss-crossed architecture (13), within which the catalysis of both cleavage and strand transfer occurs in trans (8–10). For structural reasons, and because the strand transfer step is included, it is reasonable to assume that catalysis of the hairpin processing reactions detected in this study also must occur in trans.

The general requirements of the transposase active site for the hairpin formation and opening reactions are not known, but the model of the Tn5 transposase–DNA complex (12) has provided information regarding how the transpososome may engage its substrates and how the active site might function in catalysis. In the Tn5 transpososome, the hairpin forms via dramatic bending of the DNA backbone with concomitant flipping of a thymidine base. Interactions from the phylogenetically conserved motif known as the YREK signature (44) appear to be directly involved in the process (12, 45–47), and base-flipping may be a general feature of hairpinning mechanisms. As we were unable to identify such a motif in MuA, it is difficult to reconcile the mechanism by which the Mu transpososome accommodates hairpins. At least for longer loops, the active site might actually function in a similar fashion and conformation as it does for donor cleavage, and no additional conformational changes in the protein and DNA structures need to be postulated, a scenario also consistent with the metal ion analysis. However, to engage shorter DNA loops in a catalytically competent conformation, structural flexibility in the protein partner within the transpososome may be important.

The Mu transpososome was able to accommodate a wide range of hairpin substrates and engage them in the active site for productive catalysis, although different levels of efficiency were observed with regard to the loop length. MuA appears to process longer hairpins preferentially, probably reflecting more suitable protein–DNA contacts between MuA and a few critical nucleotides in the transposon end. Although shorter loops were still applicable substrates for catalysis, they appeared to uncouple the coordinated action of MuA within the transpososome. Possibly, the accommodation of such suboptimal substrates changes the structure and functioning of the Mu transpososome so that productive reactions in both transposon ends become less likely or impossible to execute. Based on information from the crystal structure of the Tn5 transpososome, in which the active site appears relatively crowded (12,48), the observed degree of flexibility within the Mu transpososome active site is somewhat surprising. However, it is notable that despite the apparent crowding within the active site, the Tn5 transpososome can not only accommodate but also generate two types of hairpins, precise without non-complementary nucleotides and imprecise with one such nucleotide (29).

In vitro studies have shown unequivocally that some transposons such as Tn5 and Tn10 utilize hairpinning to seal transposon ends (28,29), and systems like hAt and V(D)J recombination generate hairpinned flanking DNA ends (30,31). Transposases of some other elements may have a potential to use hairpinning or related strategies but only under certain circumstances. In fact, variation within the hairpinning theme exists, and some mechanisms have been revealed under highly artificial conditions. For example, a mechanism similar to hairpinning is used to form circular transposon intermediates of IS911 (49) and related products have been described for Tn7 (50) as well as Tn10 (51) and suggested for IS903 (52).

MuA was able to process preformed model DNA hairpins in vitro, but can Mu use hairpinning mechanism in vivo? The non-replicative initial integration of Mu has been explained by a model that involves a repair step following the formation of the Shapiro intermediate (53). A very recent study supports this model, as the flanking host DNA appears to remain attached to the Mu DNA upon integration (54). Theoretical hairpinning could provide an alternate route: following the initial single-strand cleavage, MuA-catalyzed hairpinning would liberate the Mu genome from the flanking DNA, and hairpin opening followed by strand transfer would result in simple insertion. Although the above scenario is plausible, it is important to note that our current study does not directly address the question of hairpin formation by MuA, and we do not know whether MuA can form hairpins. Despite our attempts to characterize hairpin intermediates along the Mu transposition reaction pathway in vitro, such intermediates have been elusive, and no conclusive evidence has

Table 1. Transposon-target DNA junctions

| Clone | 5 bp target site duplication |
|-------|-----------------------------|
| 2a    | CAtaac atttgAC              |
| 6a    | CAtcatg agtaAC              |
| 12a   | CAtatcc attggAC             |
| 16a   | CAtaatg attacAC             |
| 17a   | CATgaca actgtAC             |
| 18a   | CACagga gttcAC              |
| 22a   | CActgg gacccAC              |
| 25a   | CAGctgg gacccAC             |
| 27a   | CAtgggt acccaAC             |
| 35    | CATgggc actcgcAC            |
| 45    | CACgccc gcccAC              |

DNA sequences of the transposon–target DNA junctions from 11 insertion events were determined from both transposon ends (Materials and Methods). The DNA regions flanking the transposon DNA are aligned appropriately to highlight in each case the 5 bp target site duplication (standard letters). The endmost nucleotides of the transposon are shown for both transposon ends (bold uppercase letters).
been gained in favor of their existence (unpublished data). Nevertheless, the formal possibility remains that MuA might use hairpinning as an alternate catalytic mechanism, in yet uncharacterized situations in vivo or under special reaction conditions in vitro. Interestingly, MuA can produce DNA hairpins under certain circumstances in vitro [P. A. Rice, H. Savilahti and K. Mizuuchi, unpublished data; (55)]. However, these hairpins differ from the transposon end DNA hairpins, as they are formed at the target DNA and represent reverse reaction products of the strand transfer.

It is possible that MuA might not be able to form hairpins but only process them, perhaps representing an evolutionary remnant of the hairpinning mechanism. Another possibility is that MuA has evolved its current, although imperfect, ability to process hairpins but not to form them. This latter scenario highlights the observation that both donor cleavage and hairpin opening are characteristically very similar reactions. Accordingly, at least with longer hairpin loops, the distinction between those two reactions is arbitrary, and hairpin opening in these cases can be regarded as a form of donor cleavage. The fact that MuA can process short hairpin loops, although relatively poorly, suggests that the evolutionary path from the non-hairpinning to hairpinning mechanism, or vice versa, may be shorter than anticipated previously. Another view is that, similar to what has been observed with protein–DNA complexes of several DNA-modifying enzymes (56–59), distortion of DNA might facilitate the initial cleavage in the Mu system. If this were the case, any hairpin substrate that could accommodate an appropriately distorted structure should be able to promote a productive cleavage reaction. Following cleavage, the catalytic machinery should then be proficient to adopt its normal configuration for the strand transfer step.

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