Guanosine triphosphate acts as a cofactor to promote assembly of initial P-element transposase–DNA synaptic complexes

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P transposable elements in Drosophila are members of a larger class of mobile elements that move using a cut-and-paste mechanism. P-element transposase uses guanosine triphosphate (GTP) as a cofactor for transposition. Here, we use atomic force microscopy (AFM) to visualize protein–DNA complexes formed during the initial stages of P-element transposition. These studies reveal that GTP acts to promote assembly of the first detectable noncovalent precleavage synaptic complex. This initial complex then randomly and independently cleaves each P-element end. These data show that GTP acts to promote protein–DNA assembly, and may explain why P-element excision often leads to unidirectional deletions.

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Transposable elements are ubiquitous mobile DNA segments found in both prokaryotic and eukaryotic organisms (Craig et al. 2002). Mobile elements are now known to make up major portions of eukaryotic genomes, including the human genome (Lander et al. 2001; Venter et al. 2005). While all known transposase proteins use a metal cofactor (Rice and Baker 2001; Craig et al. 2002; Gelbert 2002), the use of GTP in the P-element reaction is unique among this family of polynucleotidyl transferases. GTP is known to serve as a cofactor in many diverse biochemical processes, such as the ras superfamily of small GTPases (Bourne et al. 1990, 1991), the assembly of the dynamin family of vesicle transport proteins (Margolin 2000; Song and Schmid 2003; Praefcke and McMahon 2004), the assembly of tubulin into microtubules (Nogales et al. 2003), protein translation on the ribosome (Spirin 2002), and the self-splicing of group I introns (Cech 1990, Doudna and Cech 2002). Thus, it has been of interest to understand the role of GTP in P-element transposition.

Like other transposition reactions, the P-element transposes via a two-step reaction (Craig 1995) in which, initially, the two ends of the transposon are cleaved from the flanking DNA (donor cleavage) and, subsequently, the excised transposon intermediate captures and integrates into a target DNA site (strand transfer) [Fig. 1A]. Attempts to detect P-element transposase–DNA complexes using native gel electrophoresis proved unsuccessful, and we turned to the method of atomic force microscopy (AFM) to visualize and quantitate P-element reaction products and intermediates formed during transposition. This method has been used to analyze protein–DNA complexes involved in transcription and DNA repair (Wyman et al. 1997; de Jager et al. 2001). Here, we show that the role of GTP is to promote formation of synaptic complexes with both P-element ends prior to DNA cleavage. Cleavage at the P-element ends is not coordinated, with each end being randomly and independently cleaved. These findings illuminate the role GTP plays in P-element transposition and may explain why unidirectional deletions are observed following P-element excision.

Results and Discussion

Using tapping-mode AFM, initial experiments were performed using purified P-element transposase [Fig. 1B] and a linear DNA fragment carrying a 0.6-kb P element located asymmetrically in the DNA with flanking regions of 0.3 and 0.6 kb [Fig. 2A, topl. This natural 0.6-kb P element was active in vivo for transposition into the white locus (O’Hare and Rubin 1983). However, it should be noted that the 0.6-kb P element used here is much smaller than the 2.9-kb full-length autonomous element, but that in natural P-strain flies, P elements from 0.5 to 2.9 kb exist and can be mobilized (Engels 1989; Rio 2002). These reactions were performed in the presence of the cofactors GTP and Mg2+ at levels known to be optimal for both excision and strand transfer in vitro. In these reactions, four types of molecules were observed. First, we observed loops with tails, held together by a protein knob [Fig. 2B]. Measurement of these
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Figure 1. The P-element transposition reaction. (A) P-element transposition takes place in two stages, donor cleavage and target joining or integration. Following DNA repair, a new P-element insertion is created. (B) SDS-polyacrylamide gel electrophoresis analysis of purified P-element transposase after elution indicated by arrow (lane 1), and the resin after elution (lane 2), stained with Coomassie blue. Lane M is protein molecular weight standards, indicated in kilodaltons at left. (C) LM-PCR assay for donor DNA cleavage at the P-element ends analyzed by polyacrylamide gel electrophoresis using untagged transposase (H0.1) (lanes 1, 2) or the 3' Py-tagged transposase (lanes 3, 4), either with (+) or without (−) added GTP. Lane M is a 100-bp DNA marker, indicated in base pairs at left.

tail lengths indicated that these molecules were synaptic complexes in which P-element transposase had brought the two P-element ends together. Second, we observed loops with one tail, either the long or short one, consistent with cleavage of one P-element end without coordinated cleavage of the other end [Fig. 2C–D]. Third, there were circular molecules held together by a protein knob that were measured to be 0.6 kb, the size of the excised P element [Fig. 2E]. Molecules of these types were not observed in the absence of P-element transposase or when a non-P-element DNA fragment of similar length was used indicating the specificity of protein–DNA complex assembly and the dependence on terminal P-element DNA sequences [data not shown].

In order to investigate the role of GTP in the P-element transposition reaction and the relationships among the various structures observed, we performed time-course experiments in the presence and absence of GTP and quantitated the number of each type of molecule at time points from 0.5 to 26 h (Figs. 3, 4). These experiments show that GTP acts as a cofactor to stimulate the assembly of the initial noncovalent synaptic complex between the P-element transposase and the P-element DNA ends. In the absence of GTP, loops with two tails were observed with a low frequency (<1%) at early time points, but accumulated to some extent (~7%) after 26 h [Fig. 3A]. We believe that the activity responsible for the slow accumulation of this product is due to residual GTP bound to a fraction of the purified P-element transposase, as indicated from activity assays [see Fig. 1C]. In contrast, when reactions were carried out in the presence of 2 mM GTP, there was a rapid accumulation of P-element synaptic complexes [Fig. 3B]. At the 0.5-h time point, there are ~7% synaptic complexes [loops with tails], and these complexes appear to go away with time, consistent with them being the initial complexes that subsequently react to cleave the P-element ends away from the flanking donor DNA. These observations indicate that GTP acts as an allosteric effector to promote specific DNA–protein synaptic complex formation. When GTP is added to transposase, in the absence of P-element DNA, no change in the oligomerization state or apparent size of the protein was observed [data not shown].

Next, we measured the abundance of partially cleaved products with either a long tail or a short tail [Fig. 4A–D]. In the absence of GTP, both types of molecules accumulated slowly in time and to a similar extent over the time course of the reaction [Fig. 4A,C]. In contrast, in the presence of GTP, these molecules accumulated more rapidly, peaking at 3 h, and then decaying at the longest time point sampled, 26 h [Fig. 4B,D]. These observations are consistent with the single-ended cleavage products being intermediates in the reaction, which then give rise to the doubly cleaved circular molecules [Fig. 2E]. Interestingly, because both the long- and short-tailed molecules follow similar kinetics and are very similar in abundance [Fig. 4B,D], each P-element end appears to be cleaved independently and randomly during the donor cleavage reaction with release of the flanking donor DNA duplex. Thus, there is not a concerted reaction in which cleavage of both P-element ends is coupled. These findings may also explain why in vivo, following P-element excision, deletions of DNA sequences flanking the P-element insertion site are often unidirectional [Engels 1989; Adams and Sekelsky 2002, Rio 2002]. The released donor DNA end might be subject to nucleolytic degradation prior to entering the DNA repair pathway(s), thus giving rise to the observed flanking deletions.

Figure 2. Visualization of P-element transposase protein–DNA complexes. (A) Diagram of the P element containing substrate DNA fragment. A 629-bp P element is flanked by 300 and 600 bp, respectively, at each end. P-element 31-bp terminal inverted repeats and the transposase-binding sites are indicated, as is the transposase protein [spheres]. (B) Pre-cleavage synaptic complex [loop with two tails]. (C, D) Intermediate single-cleavage products, loop with short tail [C] or loop with long tail [D]. (E) Cleaved donor complex, circles with a protein “knob”. The Y-axis values were generated by using the indicated percentage of complexes [loops with tails] compared with the total of measured full-length DNA molecules [with or without protein bound] visualized under AFM. The experiment was repeated three times, and the experimental variations were shown with standard deviation error bars.
The circular cleaved donor DNA complexes (circles without tails) were also quantitated as a function of time in the presence or absence of GTP [Fig. 4E,F]. In the absence of GTP, very few circular complexes accumulated with time [Fig. 4E]. In contrast, in the presence of GTP, these doubly cleaved excised transposon complexes accumulated as a function of time and appeared to be stable, because at the last 26-h time point, they represent ~13% of the total DNA molecules in the population. The stability of these excised complexes with the transposase may explain why in vivo, the P-element excision frequency is greater than the excision and reinsertion frequency [Engels 1989; Rio 2002]. At the last 26-h time point, several molecules in which the circular complexes have apparently captured a target DNA were observed [data not shown]. Thus, the capture of target DNA by the stable cleaved donor complexes may be slow, possibly requiring a conformational change to reposition the P-element DNA termini or to generate a new DNA-binding site into which the target DNA would dock. It is also possible that in these reactions, P-element DNA is a poor target for insertion.

P elements are unusual as the only member of the transposase/integrase family to use GTP as a cofactor [Craig et al. 2002]. Previous studies showed that GTP or any of three commonly used nonhydrolyzable GTP analogs supported both the donor cleavage and strand transfer reactions carried out by P-element transposase in vitro [Kaufman and Rio 1992]. Here, we show that GTP promotes assembly of the earliest noncovalent complex of transposase with P-element DNA, in which the ends of the transposon DNA are synapsed. GTP is also known to modulate assembly of other proteins, such as dynamin [Margolin 2000; Song and Schmid 2003; Praefcke and McMahon 2004], tubulin [Nogales et al. 2003], and the bacterial restriction endonuclease McrBC [Panne et al. 2001]. Interestingly, GTP was recently implicated in preventing a transposition reaction by a potential reaction with the RAG1 subunit of the V(D)J recombinase [Tsai and Schatz 2003]. While the role of GTP in the V(D)J reaction is not clear, the C terminus of RAG2 appears to be capable of performing the same, or a similar, function [Elkin et al. 2003; Tsai and Schatz 2003]. Thus, unlike any of the other members of the transposase/integrase family, P elements use GTP to modulate the assembly of catalytically active nucleoprotein complexes. Finally, we do not know whether P-element transposase can hydrolyze GTP. This activity has not been detected using standard assays with or without DNA present. However, it might be the case that hydrolysis could be used to disassemble strand transfer complexes and recycle the protein. This type of mechanism is commonly used by GTP and ATP hydrolytic enzymes, for instance, in membrane vesicle fusion [Song and Schmid 2003; Praefcke and McMahon 2004], microtubule disassembly [Nogales et al. 2003], and DNA topoisomerase recycling [Corbett and Berger 2004]. Future AFM studies should provide insight into the role played by GTP and its possible hydrolysis in the later stages of the P-element transposition reaction cycle.

Materials and methods

**Protein purification and excision activity assay**

P-element transposase tagged at its C terminus was purified using peptide elution from a polyoma g17-g11 monoclonal antibody column as described [Porfiri et al. 1995; Rubinfeld and Polakis 1995; Mul and Rio 1997]. Untagged transposase was purified using heparin-agarose and DNA affinity chromatography as described [Kaufman and Rio 1992; Beall and Rio 1997, 1998]. LM-PCR assays to detect donor DNA cleavage at the P-element ends were performed as described [Beall and Rio 1997].

**DNA preparation**

Linear DNA excision substrate was amplified by PCR from the plasmid pSlp2-Km using Vent DNA polymerase (New England Biolabs). The DNA substrate, containing a 0.6-kb P-element flanked by either 0.3 or 0.6 kb of adjacent non-P-element DNA, was then purified from an agarose gel using a Qiagen purification kit.

**Excision reaction**

P-element donor cleavage reactions were carried out by mixing 1 µL of purified 3 Py-TNP (~50 µg/mL) in HCKED buffer [25 mM Hepes-KOH at
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