A specific terminal structure is required for Ty1 transposition

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Yeast retrotransposon Ty1 directs the synthesis of virus-like particles (VLPs) consisting of Ty1-encoded proteins, RNA, and reverse transcripts. Ty1 reverse transcripts, tagged with a selectable marker and found within VLPs, are capable of transposing into naked target DNA in vitro. Cassettes consisting of a Ty long terminal repeat (LTR) or δ, marked with supF, and flanked by appropriate restriction sites were constructed. These artificial substrates, whose termini resemble those of linear, full-length Ty1 reverse transcripts, can be coincubated with VLPs (containing unmarked reverse transcripts), resulting in the very efficient integration of the artificial substrate. The results suggest that Ty DNA is limiting for transposition in vivo, suggesting that inefficient reverse transcription regulates Ty1 transposition. Analysis of the transposition of these model substrates, which resemble in vivo Ty1 transposition intermediates or differ from them in subtle ways, shows that Ty transposition proceeds by the linkage of the 3' hydroxyl residue of the reverse transcript to target DNA.

[Key Words: Saccharomyces cerevisiae; in vitro integration; reverse transcription; transposition]

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The Ty1 element of Saccharomyces cerevisiae transposes to new sites using a mechanism akin to retroviral reverse transcription [Garfinkel et al. 1985; Mellor et al. 1985] and integration [Eichinger and Boeke 1988] processes. Ty1-encoded gene products and Ty1 RNA assemble into a virus-like particle (VLP) resembling retroviral core particles [Garfinkel et al. 1985; Mellor et al. 1985]. Small amounts of reverse transcriptase activity and free linear, double-stranded Ty1 DNAs [referred to as reverse transcripts in this paper] are also found in Ty1 VLPs [Eichinger and Boeke 1988]. An efficient tool for the study of Ty element transposition in vivo and in vitro is a family of recombinant plasmids called pGTy plasmids. These consist of the regulated yeast GAL1 promoter fused to a Ty element at its transcription initiation site, cloned in a high-copy-number plasmid [Boeke et al. 1985]. The Ty element in one such plasmid, pGTy/H3, can be marked by inserting foreign DNA into a nonessential site; such marked pGTy elements can transpose [Boeke et al. 1985, 1988].

Insertion of plasmid supF, which carries the bacterial supF gene, into pGTy/H3 results in a marked element whose transposition can be monitored both in vivo and in vitro. Transposition in vivo is monitored by determining the frequency of transposition into host chromosomal DNA per unit time [Boeke et al. 1985, 1988]. In vitro transposition (i.e., the in vitro integration step of the transposition process) is measured as follows. Ty1-H3-supF VLPs are prepared from cells containing the pGTy1-H3-supF plasmid. The VLPs contain a small amount of preformed Ty1-H3-supF reverse transcript, primarily in the form of double-stranded linear DNA molecules, although other forms of Ty DNA [such as one long terminal repeat (LTR) circles] are also detected [Eichinger and Boeke 1988]. The supF-marked VLPs are incubated with target agt− WES/AB DNA and Mg²⁺. This A strain has amber mutations in three essential genes. The reaction mixture is then deproteinized, and the nucleic acid mixture is packaged into λ phage heads. Transposition of Ty1-H3-supF into nonessential portions of the λ genome results in recombinant phage λ derivatives able to form plaques on nonsuppressing bacterial host strains. Transposition is quantified by comparing the number of recombinant (supF-containing) phage [number growing on the nonsuppressing host] to the total number of phage in the final packaging reaction [number growing on a suppressor-containing host]. The in vitro reaction requires only the particulate Ty VLP fraction; there is no direct evidence that soluble macromolecular factors participate in the reaction [Eichinger and Boeke 1988]. [It is formally possible that non-Ty products contaminating the VLPs participate in the integration reaction.] Ty VLPs have structural and functional properties that are very similar to those of particulate Moloney murine leukemia virus nucleoprotein complexes isolated from infected cells [Brown et al. 1987, 1989; Fujiwara and Mizuuchi 1988, Bowerman et al. 1989, Fujiwara and Craigie 1989].
Results

Artificial substrates for in vitro Ty1 transposition

We are studying the Ty1 integration process with defined components by developing a set of cloned substrates consisting only of terminal Ty LTR (or B) sequences flanking a plasmid, \( \pi \alpha n7 \), bearing the selectable marker \( \text{supF} \). These substrates are generated by cutting with various restriction enzymes whose recognition sites flank the LTR sequences and then modifying the restriction enzyme-generated termini further with other enzymes. Two such substrates have been constructed, using polymerase chain reaction (PCR), and cloned in M13 vectors (for details of the constructions, see Materials and methods). The structures of M13 inserts \( \delta \pi 1 \) (flanked by FokI and NsiI sites) and \( \delta \pi 2 \) (flanked by HindIII sites), from which a family of substrates can be generated, are shown in Figure 1. The \( \delta \pi 1 \) insert can be converted to a blunt-ended substrate molecule \( \delta \pi 1-0 \), whose terminal structure corresponds to that of a full-length Ty DNA sequence, by cutting M13\( \pi \alpha n7 \) DNA with FokI and filling in the ends (with DNA polymerase I large fragment). A molecule of identical structure can be generated by cutting M13\( \pi \alpha n7 \) with NsiI (or M13\( \pi \alpha n2 \) with HindIII) and treating with S1 nuclease.

\( \delta \pi 1-0 \) molecules, when mixed with Ty1 VLPs (which lack a selectable marker, themselves) and target \( \lambda \) DNA, are efficiently transposed into the \( \lambda \) DNA in vitro, presumably by a reaction requiring the transposon-encoded IN (integrase) protein. The VLP-mediated reaction using this exogenous substrate has the same very simple biochemical requirements of the reaction based on marked Ty VLPs with internal \( \pi \alpha n7 \) substrates. Under optimal conditions, at least 0.09% of this exogenous \( \delta \pi 1-0 \) substrate is converted into recombinant molecules (calculated as indicated in Materials and methods). It is also instructive to consider the efficiency of the reaction in terms of how many cell-equivalents of VLPs are required to give rise to each in vitro transposition event. When 50 ng of \( \delta \pi 1-0 \) substrate (a saturating amount) is mixed

Figure 1. Structure and transposition of recombinant M13 phage DNA and derived \( \delta \pi \) substrates. [a] The diagram indicates the structure of M13\( \pi \alpha n1 \) and M13\( \pi \alpha n2 \) DNA. The design of M13\( \pi \alpha n1 \) takes advantage of the unusual properties of FokI, whose recognition and cut sites are different. The boxed triangle indicates LTR or \( \delta \) sequence. The sequences above and below the diagram show the relevant sequences surrounding the \( \pi \alpha n7 \) sequence. \( \delta \) sequences are shaded. Restriction enzyme recognition sites are boxed; cleavage sites are indicated by arrows. [b] Structures of substrates derived from M13\( \pi \alpha n7 \) phage DNAs (for detailed methods of synthesis, see Materials and methods). Terminal functional groups are indicated in boldface when different in structure or position from those in \( \pi \alpha n1-0 \). \( \pi \alpha n1-0 \) is produced by cleaving M13\( \pi \alpha n1 \) with FokI and filling in the ends. \( \pi \alpha n1-1 \) is produced by FokI cleavage of M13\( \pi \alpha n1 \). \( \pi \alpha n1-2 \) differs from \( \pi \alpha n1-0 \) in that it lacks a 3'-hydroxyl group (the terminal adenosines were incorporated in the form of 2',3'-ddATP). \( \pi \alpha n1-3 \) lacks 5' phosphate groups as the result of enzymatic dephosphorylation. \( \pi \alpha n1-4 \) is produced by NsiI cleavage of M13\( \pi \alpha n1 \). Note that the terminal dinucleotide of this substrate, CA, is identical to the functional ends of \( \pi \alpha n1-0 \). \( \pi \alpha n2-1 \) is produced by cleaving M13\( \pi \alpha n2 \) with HindIII. [c] The effectiveness of these substrates in the in vitro transposition reaction (relative to \( \pi \alpha n1-0 \)) is indicated.
with VLPs derived from \( \sim 3.5 \times 10^7 \) cells under ideal conditions, then \( 3.4 \times 10^7 \) transposition events result. Thus, a one-cell equivalent of VLPs gives rise to one transposition event. This efficiency, however, because the efficiency varies from preparation to preparation and presumably reflects the ability to recover is presumably a minimum estimate of the "active" VLPs.

**In vitro transposition events have the expected structure**

Restriction mapping of 10 \( \lambda / \delta \pi \) recombinant phage was performed; the digests were all consistent with simple insertion of \( \delta \pi \) into \( \lambda \). Nine of these were unambiguously mapped within the \( \lambda \) genome, and three were sequenced on both ends. The expected 5-bp duplication of \( \lambda \) sequence was found at the termini of each \( \delta \pi \) insert [Fig. 2].

**Ty DNA is limiting for transposition**

Unlike the reaction requiring internal Ty-\( \pi \text{-an7} \) substrate, the reaction utilizing exogenous \( \delta \pi \) substrates can be sustained over long time periods [Fig. 3]. Whereas the reaction utilizing internal Ty-\( \pi \text{-an7} \) DNA reaches a maximum within several minutes, the \( \delta \pi \text{-1-0} \)-dependent reaction is nearly linear for at least 1 hr. This result suggests that the amount of DNA with appropriate terminal structures inside the Ty-VLPs is limited. This interpretation is also supported by our inability to observe conversion of a significant fraction of the Ty DNA found inside particles into recombinant molecules. [To detect such recombinants, we used a Southern blotting approach similar to that used by others studying retrovirus integration [Fujiiwara and Mizuuchi 1988; Brown et al. 1989; data not shown]]. Such high-efficiency conversion of endogenous reverse transcripts to recombinant form has been reported for the in vitro reaction of Moloney murine leukemia virus [MuLV] nucleoprotein particles, in which 10–25% of the reverse transcripts integrate into target DNA. We estimate that the Ty integration reaction utilizing internal substrate incorporates only 0.1–0.3% of the reverse transcripts in the VLPs into target DNA [Eichinger and Boeke 1988]. We tested whether the in vitro integration reaction utilizing endogenous reverse transcripts is limited by non-nucleic acid components required for integration by adding \( \delta \pi \text{-1-0} \) DNA to such a reaction. Addition of exogenous artificial substrate results in \( \sim 100\)-fold more in vitro transposition events (Table 1).

We also made direct measurements of the total amount of Ty DNA in Ty1-VLP preparations [from cells bearing induced pGTy1-H3 plasmids]. Remarkably, the VLPs in these cells contain only \( \sim 10 \) full-length molecules of unintegrated Ty DNA per cell (Table 2).

**Structural requirements for the in vitro integration reaction**

Because the integration reaction occurs under conditions that prevent DNA synthesis, it is presumed that the immediate product of integration is a recombinant molecule with a 5-bp gap in the target DNA at each end of the \( \delta \pi \) sequences; these gaps could be at either the 5' or 3' ends of \( \delta \pi \). Several variant forms of the \( \delta \pi \) substrate have been constructed to evaluate the reaction requirements at the level of the Ty1 substrate.

We generated a family of analogs of the efficient trans-

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**Figure 2.** Structure of termini of in vitro integration events. The 5-bp duplication of Ty sequences is indicated by shading. Sites of insertion into \( \lambda \) are \( \lambda \text{dp-1}, 23376; \lambda \text{dp-3}, 47218; \lambda \text{dp-9}, 23677 \) [Sanger et al. 1982].

**Figure 3.** Time courses of in vitro transposition reactions. Aliquots were withdrawn from the reaction, as indicated. (□) Reaction using VLPs containing \( \pi \text{-an7} \)-marked reverse transcripts, prepared from strain JB619. (●) Reaction using VLPs containing unmarked reverse transcripts, prepared from strain JB224, and \( \delta \pi \text{-1-0} \) DNA.

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**Table 1.** Transposition of endogenous reverse transcripts is limited by the amount of available substrate

| Source of Ty VLPs | \( \delta \pi \text{-1-0} \) added [ng] | Total marked transposition events* |
|-------------------|-----------|----------------------------------|
| JB224b            | 0         | \(<6.0 \times 10^6\)             |
| JB224b            | 50        | \(3.4 \times 10^7\)             |
| JB619c            | 0         | \(1.4 \times 10^7\)             |
| JB619c            | 50        | \(1.3 \times 10^7\)             |

*Transposition of both endogenous marked reverse transcript (in the case of JB619) and of \( \delta \pi \) DNA are detected.

bContains unmarked Ty reverse transcripts [Boeke et al. 1985; Eichinger and Boeke 1988].

cContains \( \pi \text{-an7} \)-marked Ty reverse transcripts [Eichinger and Boeke 1988].
position substrate δπ1-0 by a variety of enzymatic methods (for details, see Materials and methods). These analogs were then compared to δπ1-0 for in vitro integration efficiency. The substrates tested fell into two classes: (1) inactive as a transposition substrate (transposing at frequencies at least 100-fold lower than δπ1-0 itself), or (2) about as effective as δπ1-0 itself (Fig. 1). Substrate δπ1-2, which is identical to δπ1-0, except that it lacks a 3'-hydroxyl group, is inactive for transposition. This substrate is prepared by filling in the last adenylate residues using 2',3'-dideoxyadenosine triphosphate rather than 2'-deoxyadenosine triphosphate. Substrate δπ1-4, produced by digesting M13δπ1 with NsiI, has four extra nucleotides on the 3' end of the substrate; it is inactive as a substrate for transposition. Furthermore, removal of the 5' phosphate of substrate δπ1-0 with alkaline phosphatase produces a substrate, δπ1-3, fully active in transposition, indicating that the 5' phosphate groups of the transposon are dispensable for transposition. These results strongly support the idea that the 3'-hydroxyl end of the δπ substrate is joined to the 5' end of a staggered break in target DNA and that the 5' ends of δπ and the 3' ends of target DNA flank the gaps. Substrate δπ2-1, produced by digestion of M13δπ2 with HindIII, is inactive as a substrate. Apparently, although no 5' phosphate is required for δπ transposition, the presence of four extra deoxyribonucleosides at the 5' end of the substrate blocks δπ transposition effectively. This is surprising because in natural Ty1 reverse transcripts, the 5' end (of the "bottom" strand) is thought to be formed by extension of RNA primers such as initiator–methionine tRNA (Eibel et al. 1981; K. Chapman, A. Bystrom, and J.D. Boeke, unpubl.). The hindrance of the in vitro reaction by extra DNA bases suggests that such RNA primers must be removed prior to formation of ends active in transposition.

A covalently closed circular substrate is inactive in transposition

We generated a covalently closed circular form of δπ1-0, as well as concatemers of δπ1-0, by ligation of δπ1-0 monomers with T4 DNA ligase (Fig. 4). These derivatives of δπ1-0, as well as unreacted δπ1-0, were eluted from an agarose gel and used as substrates in the in vitro integration reaction. Whereas δπ1-0 (and its concatemerized forms; data not shown) readily gave rise to recombinants, none were detected with the covalently closed circular material, indicating that it is at least 50-fold less active in transposition than the corresponding monomer linear substrate.

Only terminal Ty1 sequences are required for efficient integration

We constructed two deletion derivatives of M13δπ1, M13δπ1-U3Δ12 and M13δπ1-U5Δ12 (Fig. 5), which contain only 12 bp of Ty1 sequence at the 5' and 3' ends of the LTR, respectively. FokI treatment, followed by filling in, converts both of these mutant DNAs into active substrates. The activities of these substrates are within twofold of the activity of δπ1-0. Similarly, the activity of the double deletion mutant, M13δπ1U3Δ12/U5Δ12, containing only 24 bp of Ty1 sequence, is within twofold of δπ1-0 (Table 3). Thus, the transposition machinery recognizes a region of <13 bp at each terminus of the δπ substrate molecule.

Table 2. DNA content of Ty VLPs

| Strain | Number of cells | Ty1 DNA* (ng) | Copies per cell |
|--------|----------------|---------------|----------------|
| JB619  | 1.7 x 10^9       | 100           | 7.8            |
| JB224  | 1.4 x 10^9       | 130           | 14.7           |

*Amount of DNA recovered from VLP preparation, determined by quantitative Southern blotting with plasmid DNA standards.

Figure 4. Ligation products of δπ1-0 and their transposition. (A) Preparative ethidium-bromide-stained agarose gel. (Lane 1) Molecular weight markers; (lane 2) unligated δπ1-0; (lane 3) ligated δπ1-0. Indicated bands were purified by electroelution and used for in vitro transposition. The relative efficiency of transposition, per molecule of input δπ substrate, normalized to δπ1-0, is indicated for bands 1 and 2. (B) DNA eluted from bands 1 and 2 was re-electrophoresed with (lanes 4–6) and without (lanes 1–3) prior DdeI digestion. There is one DdeI site in δπ1-0. The gel was transferred to nitrocellulose and hybridized with a Ty-specific probe. Based on their behavior following DdeI digestion, the bands were identified as follows: (band 1) Monomeric circles; (band 2) monomeric linears (may also contain some dimeric circles); (band 3) dimeric linears; (bands 4 and up) multimeric linears. (Lanes 1 and 4) Unligated δπ1-0; (lanes 2 and 5) band 1; (lanes 3 and 6) band 2. Band 1 material could only be detected after cleavage (cf. lanes 2 and 5).
Discussion

Ty1 reverse transcripts are found in the form of a nucleoprotein complex, the VLP [Eichinger and Boeke 1988]. A similar structure can be isolated from cells infected with the Moloney MuLV [Brown et al. 1987, Bowman et al. 1989]. These structures contain all of the macromolecules needed to effect in vitro integration. Our studies and related studies of Moloney virus integration [Fujiiwara and Craigie 1989] show that the reverse transcript can be provided as a cloned molecule whose termini resemble those of native reverse transcripts; apparently, the integration machinery can recognize these soluble molecules and, in the case of Ty1 at least, can integrate them very efficiently. Whether the joining reaction takes place within the VLP [where the IN protein presumably resides] or outside the VLP structure is uncertain. However, we note that nucleases and substrates for reverse transcriptase are able to access the Ty-encoded reverse transcriptase without detergent treatment or other obvious insult to the VLPs [Garfinkel et al. 1985; Mellor et al. 1985]. Thus, it is not unlikely that δr1-0 DNA can enter Ty1 VLPs. By manipulating the functional groups and DNA sequences at the termini of these molecules, we have established that the Ty1 integration reaction [1] works efficiently using a blunt-ended substrate, [2] requires the 3'-hydroxyl group of the terminal adenylate residues, [3] does not require 5' phosphates, and [4] recognizes sequences within the terminal 12 bp of the reverse transcript.

That the Ty1 integration reaction uses a blunt-ended substrate differs from the situation in retroviruses, where a substrate with a 2-bp gap at the 3' end, which is generated by IN protein [Brown et al. 1989; Roth et al. 1989], is used. This difference is perhaps not too surprising when the slight differences in sequence organization between Ty1 and retroviruses is considered. The position of the putative primer binding site for Ty reverse transcription of the [-] strand [relative to the LTR sequence] is offset from that of retroviruses by 2 bp [Fig. 6]. Because the exact nature of + strand priming in Ty1 is unknown, a similar sequence comparison at this end cannot be made. However, if the reverse transcript is to be terminally symmetrical, as it is in retroviruses, then a similar juxtaposition of priming site and LTR is expected for the + strand in Ty1 reverse transcripts.

The terminal functional group requirements for the reverse transcript indicate that the recombinant intermediate molecule, generated when a Ty1 reverse transcript integrates into a target DNA, is identical in overall topology to that formed during bacteriophage Mu and Tn10 transposition [Craigie and Mizuuchi 1985; Benjamin and Kleckner 1989] and by Moloney MuLV integration [Fujiiwara and Mizuuchi 1988, Brown et al. 1989]. Thus, all transposons for which this structure has been determined share this property. The small number of nucleotides required for the integration reaction [presumably reflecting the recognition site of IN] is also in line with that determined in retroviral systems [Panganiban and Temin 1983; Colicelli and Goff 1985, 1988].

We attribute the surprising efficiency of the Ty1 reaction utilizing artificial substrates to the very small amount of Ty1 DNA found in the VLPs. Probably, there are many Ty VLPs that lack Ty DNA but contain the appropriate protein(s) required for transposition. Moreover, the 20% efficiency of TYA/TYB frameshifting [Clare et al. 1988] and efficient incorporation of p190-TYA/TYB, the precursor to Ty1 IN protein, into VLPs [Adams et al. 1987, Muller et al. 1987, Youngren et al. 1988] suggest that there are dozens to hundreds of IN proteins per VLP, whereas there should be only one reverse transcript per VLP, if retroviral precedents hold here [Panganiban and Fiore 1988]. Finally, many of the Ty1 DNA molecules inside VLPs are heterogeneous in end structure and presumably nonfunctional, suggesting that the inefficiency of reverse transcriptase and/or RNase H activities encoded by Ty1 IN limit transposition frequency. Indeed, the number of Ty1 VLPs observed by electron microscopy in cells in which high-level transposition has been induced [hundreds to thousands] far exceeds the number of transposition events observed over a period of several days in those cells [typically one to five per genome [Boeke et al. 1985, 1988]]. In vitro, the activity of Ty1 reverse transcriptase is sluggish compared to that of retroviral enzymes [Garfinkel et al. 1985, Mellor et al. 1985]. Ty1-encoded RNase H activity has not yet been directly demonstrated or measured, although it too may be relatively inactive; many of the double-stranded linear molecules isolated from Ty1 VLPs are resistant to cleavage by restriction endonucleases [Eichinger and Boeke 1988], consistent with the existence of RNA tracts within many in vivo Ty1 re-

Table 3. Transposition of deletion mutant substrates

| Substrate | Number of transpositions (50 ng DNA)* |
|-----------|-------------------------------------|
| δr1-0     | 3.7 × 10^6                          |
| δr1-0 U3Δ12 | 3.4 × 10^6                          |
| δr1-0 U5Δ12 | 1.7 × 10^6                          |
| δr1-0 U3Δ12/U5Δ12 | 1.5 × 10^6                        |

*These are the average obtained with two independent constructs, except the double mutant, which was obtained using pooled DNA from two independent constructs.
verse transcripts (M. Ciriacy, U. Pott, W. Laufer, C. Drewke, and F. Muller, pers. comm.).

Materials and methods

Plasmid constructions

$\delta$1 was constructed as follows. Two oligonucleotides, JB39 (GGTCCGCAATGTTATTCATGTTGGAATAGAATC) and JB40 (GGTCCGCAATGTTATTCATGTTGGAATAGAATC), were used as primers in a PCR reaction that used a restriction fragment containing the 5' LTR of Ty1-H3, which contains a Xhol site, as the template. The PCR product of the appropriate size was phenol-extracted and digested with SalI. The SalI PCR fragment was then cloned into the M13mp18 SalI site, to give M138-9. Plasmid 8mr7, digested with SalI, was cloned into the unique Xhol site of M138-9, to give M138mr1.

M138mr2, which has HindIII sites flanking the 8, was constructed as follows. M138mr1 was digested with FokI and filled in. The 8-containing FokI fragment was isolated and ligated to M13mp18 SalI site, to give M138-9. Plasmid 8mr7, digested with SalI, was cloned into the unique Xhol site of M138-9, to give M1388mr1.

Deletion derivatives of M138mr1, 8mr1U3A12 and 8mr1U5A12 were constructed as described [Kunkel 1985]. The oligonucleotides used as primers were JB51 (GGATCTCACTGCCTTCCTGTGGAGT) for U3A12 and JB49 (GGTCCGCAATGTTATTCATGTTGGAATAGAATC) for U5A12. The double mutant was constructed by remutagenizing 8mr1U3A12 with JB49. The termini of all constructs were sequenced and shown to be exactly as indicated.

Preparation of 8 substrates

8mr1-0 was produced by cleaving M138mr1 with FokI and filling in the ends. 8mr1-1 was produced by FokI cleavage of M138mr1. 8mr1-2 was produced by cleavage with FokI, filling in the presence of [a-32P]dCTP, electrophoretic purification of the product, filling in in the presence of dATP and dTTP, electrophoretic purification of the product, and filling in with dCTP and 2',3'-dATP (a control aliquot was filled in in the presence of dCTP and dTTP and was fully active). 8mr1-3 is prepared by treating FokI-digested M138mr1 with calf intestine alkaline phosphatase, followed by filling in. An aliquot of the reaction product was tested for phosphate removal by self-ligation, no ligation of 8mr1-3 was observed, whereas control 8mr1-0 made at the same time was >90% ligated. 8mr1-4 was produced by NsiI cleavage of M138mr1. Note that the terminal dincucleotide of this substrate, CA, is identical to the functional ends of 8mr1-0. 8mr2-1 is produced by cleaving M138mr2 with HindIII. (A control aliquot of 8mr2-1 subsequently treated with S1 nuclease was fully active.)

All substrates were agarose gel-purified prior to use in in vitro integration reactions.

VLP isolation and in vitro transposition reactions

VLPs were first purified over sucrose step gradients, as described [Eichinger and Boeke 1988] using buffer B/EDTA throughout the procedure, followed by centrifugation of the three peak 1.2-ml fractions on a second, linear 15–50% sucrose gradient. The RT-positive fractions were pooled and concentrated by pelleting at 80,000g for 1 hr at 4°C. Pellets from each 500-ml culture were resuspended in 100 μl of H2O and stored (–80°C). In vitro transposition reactions were performed by combining in the following order: H2O (to obtain a final volume of 20 μl), 2 μl of 8 substrate (0–25 ng/μl), 2 μl of VLP suspension, 2.5 μl of XgtWES · λB concatenated DNA (400 ng/μl), 2 μl (1 μg) of 10× transposition buffer, and 5 μl of 20% PEG8000. Contents were mixed by gentle tapping and incubated (30°C) for 30 min (or as noted otherwise). Subsequent steps [solvent extraction, packaging reactions, etc., were as described [Eichinger and Boeke 1988]]. Total transposition events per reaction were calculated using the formula A/B · C/D · E, described by Brown et al. [1987]. [A = number of plaques on CES200; B = number of plaques on LE392; C = number of XgtWES genomes in 1.0 μg of XgtWES; D = fraction of the XgtWES genome tolerant of insertion (0.27); E = a correction factor for the differential packaging/plating efficiency between XgtWES and recombinant phage.]

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References

Adams, S.E., J. Mellor, K. Gull, R.B. Sim, M.F. Tuite, S.M. Kingsman, and A.J. Kingsman. 1987. The functions and relationships of Ty-VLP proteins in yeast reflect those of mammalian proteins. Cell 49: 111–119.

Benjamin, H.W. and N. Kleckner. 1989. Intramolecular transposition by Tn10. Cell 59: 373–383.

Boeke, J.D., D.J. Garfinkel, C.A. Styles, and G.R. Fink. 1985. Ty elements transpose through an RNA intermediate. Cell 40: 491–500.

Figure 6. Relationship of primer binding site and LTR sequences in Ty1 and retroviruses. The U5 sequences are separated from the primer binding site sequences by a vertical line. Regions of complementarity between Ty RNA and tRNA^{Met} [Eibel et al. 1981] and Moloney RNA and tRNA^{Pro} [Chen and Barker 1984] are shaded. The sequences of Ty1 and Moloney murine leukemia virus RNAs and the corresponding (–) strong stop DNAAs are shown as the upper double-stranded sequences of each pair. The lower double-stranded sequences are the predicted terminal structures of the right ends of the completed reverse transcripts, in the case of Moloney, the 3' terminal TT is removed by IN prior to its joining to target DNA. No such reaction is expected to be required for Ty1.
Boeke, J.D., H. Xu, and G.R. Fink. 1988. A general method for the chromosomal amplification of genes in yeast. *Science* 239: 280–282.

Bowerman, B., P.O. Brown, J.M. Bishop, and H.E. Varmus. 1989. A nucleoprotein complex mediates the integration of retroviral DNA. *Genes Dev.* 3: 469–478.

Brown, P.O., B. Bowerman, H.E. Varmus, and J.M. Bishop. 1987. Correct integration of retroviral DNA in vitro. *Cell* 49: 347–356.

——. 1989. Retroviral integration: structure of the initial covalent product and its precursor, and a role for the viral IN protein. *Proc. Natl. Acad. Sci.* 86: 2525–2529.

Chen, H.R. and W.C. Barker. 1984. Nucleotide sequences of the retroviral long terminal repeats and their adjacent regions. *Nucleic Acids Res.* 12: 1767–1778.

Clare, J.J., M. Belcourt, and P.J. Farabaugh. 1988. Efficient translational frameshifting occurs within a conserved sequence of the overlap between the two genes of a yeast Ty1 transposon. *Proc. Natl. Acad. Sci.* 85: 6816–6820.

Colicelli, J. and S.P. Goff. 1985. Mutants and pseudorevertants of Moloney murine leukemia virus with alterations at the integration site. *Cell* 42: 507–517.

——. 1988. Sequence and spacing requirements of a retrovirus integration site. *Cell* 54: 497–504.

Craigie, R. and K. Mizuuchi. 1985. Mechanism of transposition of bacteriophage Mu: structure of a transposition intermediate. *Cell* 41: 867–876.

Eichel, H., J. Gafner, A. Stotz, and P. Philippsen. 1981. Characterization of the yeast mobile genetic element Ty1. *Cold Spring Harbor Symp. Quant. Biol.* 45: 609–617.

Eichinger, D.J. and J.D. Boeke. 1988. The DNA intermediate in yeast Ty1 element transposition copurifies with virus-like particles: Cell-free Ty1 transposition. *Cell* 54: 955–966.

Fujisawa, T. and K. Mizuuchi. 1988. Retroviral DNA integration: structure of an integration intermediate. *Cell* 54: 497–504.

Garfinkel, D.J., J.D. Boeke, and G.R. Fink. 1985. Ty element transposition: Reverse transcriptase and virus-like particles. *Cell* 42: 507–517.

Kunkel, T.A. 1985. Rapid and efficient site-specific mutagenesis without phenotypic selection. *Proc. Natl. Acad. Sci.* 82: 488–492.

Mellor, J., M.H. Malim, K. Gull, M.F. Tuite, S. McCready, T. Dibbayawan, S.M. Kingsman, and A.J. Kingsman. 1985. Reverse transcriptase activity and Ty RNA are associated with virus-like particles in yeast. *Nature* 318: 583–586.

Müller, F., K.-H. Bruhl, K. Freidel, K.V. Kowallik, and M. Giriac. 1987. Processing of Ty1 proteins and formation of Ty1 virus-like particles in *Saccharomyces cerevisiae*. *Mol. Gen. Genet.* 207: 421–429.

Panganiban, A.T. and D. Fiore. 1988. Ordered interstrand and intrastrand DNA transfer during reverse transcription. *Science* 241: 1064–1069.

Panganiban, A.T. and H.M. Temin. 1983. The terminal nucleotides of retrovirus DNA are required for integration but not virus production. *Nature* 306: 155–160.

Roth, M.J., P.L. Schwartzberg, and S.P. Goff. 1989. Structure of the termini of DNA intermediates in the integration of retroviral DNA: Dependence on IN function and terminal DNA sequence. *Cell* 58: 47–54.

Sanger, F., A.R. Coulson, G.F. Hong, D.F. Hill, and G.B. Petersen. 1982. Nucleotide sequence of bacteriophage lambda DNA. *J. Mol. Biol.* 162: 729–773.
A specific terminal structure is required for Ty1 transposition.

D J Eichinger and J D Boeke

Genes Dev. 1990, 4:
Access the most recent version at doi:10.1101/gad.4.3.324

References
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