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# Evidence for retrotransposition of the *I* factor, a *LINE* element of *Drosophila melanogaster*

(transposition/reverse transcription/intron/hybrid dysgenesis)

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ABSTRACT LINEs are transposable elements found in various eukaryotes such as plants, protists, insects, and mammals. Their transposition is usually difficult to study, particularly in humans, where some diseases have been shown to result from LINE insertion mutations. This is due to the fact that most copies of any particular family of elements are defective and that their transposition frequency is low. By contrast, the I factor of Drosophila melanogaster transposes at high frequency during I-R hybrid dysgenesis and is a good model for studying the LINE element superfamily. LINEs encode putative polypeptides showing similarities with viral reverse transcriptases but, unlike viral retrotransposons, they do not have terminal repeats and their ability to transpose by reverse transcription has previously only been inferred from structural analysis. Here we present direct evidence for LINE retrotransposition. Transposition of an I factor marked by an intron resulted in accurate removal of the intron.

The I factor is a transposable element of Drosophila melanogaster related to LINEs. These elements are found in a variety of phyla including plants, protists, insects, and mammals (1). Particularly significant is their quantitative and qualitative contribution to mammalian genome organization and expression. For instance, in humans up to 5% of the genome consists of the same LINE family (LI). LI elements have been shown to transpose, and de novo insertions of these elements have been found associated with hemophilia A and breast carcinoma (2, 3).

LINEs are devoid of terminal repeats but have an A-rich sequence, usually preceded by a polyadenylylation signal, at their 3' ends. They contain two long open reading frames, one of which encodes a putative polypeptide showing similarities with viral reverse transcriptases (1). They are thought to transpose by reverse transcription of an RNA intermediate, but this is based only on structural studies (4-6). Analysis of the mechanism of LINE element transposition is particularly difficult in mammals because a high proportion of these elements are defective and the frequency of transposition is very low.

I factors, which are responsible for the I-R system of hybrid dysgenesis in D. melanogaster (7, 8), have properties that overcome these difficulties. I-R hybrid dysgenesis is a syndrome that includes female infertility and increased frequencies of mutations and is associated with frequent replicative transposition of I factors. There are two categories of strains in D. melanogaster with respect to the I-R system: inducer strains that contain active I factors and reactive strains that do not. Crosses between inducer males and reactive females produce dysgenic  $F_1$  females that are poorly fertile. Mutations occur at high frequency in their germ line. I factors are stable in inducer strains but are activated and

transpose at high frequency in the germ line of dysgenic females (9). They never transpose in males.

Active I factors have been cloned. They are 5.4 kilobases (kb) long and are devoid of terminal repeats. They terminate at the 3' end by several repeats of the sequence TAA. They contain two long open reading frames. One of them, ORF2, encodes a putative polypeptide showing similarities with reverse transcriptases and RNase H (10).

P-M dysgenesis is an independent system of hybrid dysgenesis that has some characteristics in common with I-R dysgenesis but involves transposable elements, P elements, that use a strictly DNA-dependent transposition mechanism (11). Strains completely devoid of P elements are called M strains, whereas strains containing active P elements are called P strains.

Transposition by reverse transcription of an RNA intermediate has been clearly demonstrated for TyI elements in  $Saccharomyces\ cerevisiae$  by showing that an intron inserted within the element is removed during transposition (12). We have used a similar method to investigate the mechanism of transposition of the I factor of D. melanogaster, a member of a class of elements that are structurally quite distinct from retrovirus-like elements such as TyI. For this, we have introduced an intron into an I factor. Its transposition resulted in precise removal of the intron, providing direct evidence for retrotransposition of LINEs. Our results also indicate that trans-complementation of elements of this type is possible.

### **MATERIALS AND METHODS**

Construction of the  $\pi I$  Element. A marked I factor  $(\pi I)$  was constructed by introducing an intron-containing fragment of the P element of D. melanogaster into the functional I factor of clone pI407 (13, 14) (see Fig. 1). For this, the Pvu II/Pvu II restriction fragment of clone  $p\pi 25.7$  (15), which contains the second intron of the P factor, was cloned in the same transcriptional orientation as I factor sequences in Kpn I-cleaved pI407 (8) after treatment with T4 DNA polymerase. The 7.1-kb Sal I/Sal I fragment containing this marked I element,  $\pi I$ , was cloned into the Sal I site of the pUChsneo transformation vector (16), giving clone pPI.

Construction of Transgenic Lines. Clone pPI was microinjected into embryos  $(G_0)$  of the reactive strain Cha together with the  $p\pi 25.7$  wing-clipped helper plasmid as described (17). Transgenic flies were selected among the progeny of these embryos by growing  $G_1$  larvae on food containing G418 (1 mg/ml). We recovered one transformed line in which the  $\pi I$  element containing P-element transposon was located on the X chromosome. It was called  $\pi IX$ . This insertion was subsequently moved three times to chromosome 3 of the inducer strain DcxF by P-mediated transposition using the

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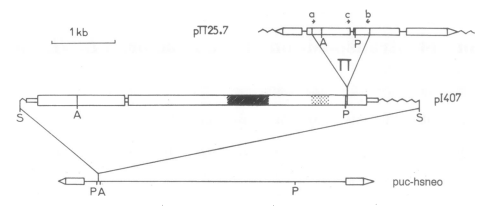


Fig. 1. Map of the  $\pi I$  element. The P and I elements in clones  $p\pi 25.7$  (15) and p1407 (13) are drawn in the conventional 5' to 3' orientation. They are both flanked by genomic DNA represented by wavy lines. The terminal noncoding sequences and the exons and introns of P are represented by open arrowheads, open boxes, and thin lines, respectively. The orientations and positions of the primers used to amplify (a and b) and sequence (c) the  $\pi$  fragment are shown by arrows. The boxes in the I factor correspond to ORF1 and ORF2 and the hatched and stippled regions represent amino acid sequences similar to reverse transcriptase and RNase H, respectively. The pUChsneo transformation vector (16) contains pUC8 sequences and the hance selectable marker flanked by P factor terminal repeats. A, Ava I; P, Pst I; S, Sal I.

P(ry<sup>+</sup>  $\Delta 2$ -3)(99B) chromosome as a donor of transposase (11). The corresponding transgenic strains were called  $\pi$ ID1,  $\pi$ ID2, and  $\pi$ ID3.

**DNA** Amplification Using PCR. Genomic DNAs were extracted according to a proteinase K/phenol/RNase/phenol procedure. The *Pst* I fragment of  $\pi I$  was amplified by two successive steps of PCR (18). One microgram of genomic DNA was mixed with 0.25  $\mu g$  each of oligonucleotides a and b (see

Fig. 1) and  $0.5 \mu l$  (2.5 units) of Taq polymerase (Perkins–Elmer/Cetus) in 100  $\mu l$  of PCR buffer [6.7 mM MgCl<sub>2</sub>/16.6 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>/67 mM Tris·HCl, pH 8.8/0.25 mM each dNTP]. The reaction mixture was overlaid with mineral oil and subjected to 30 temperature cycles (94°C denaturing, 1 min; 60°C annealing, 1 min; 72°C extension, 1.5 min). About 2 pg ( $10^{-5}$  of the PCR product) was amplified again in the same conditions as described above for sequencing.

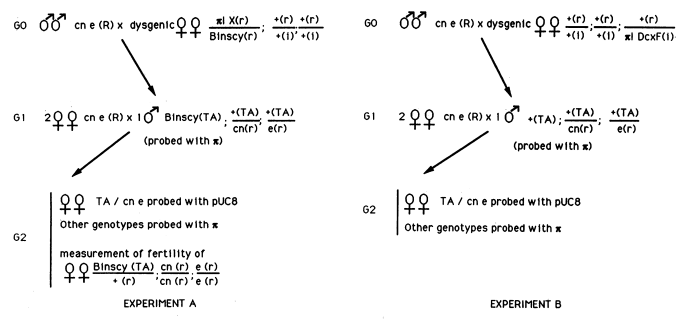


Fig. 2. Mating scheme to detect I-R dysgenesis-dependent transposition of the  $\pi I$  element from various insertion sites. Only M strains were used so that no transposition events could result from P-element activity. Chromosomes originating from an inducer or a reactive origin are symbolized by (i) and (r), respectively. (TA) is a target chromosome that might have acquired I and  $\pi I$  elements in the germ line of the  $G_0$  dysgenic females. Experiment A was designed to detect transposed copies of the  $\pi I$  element located on the X chromosome of the  $\pi IX$  stock and experiment B was designed to detect transposed copies of the  $\pi I$  element located on the DcxF chromosome of stocks  $\pi$ ID1,  $\pi$ ID2, and  $\pi$ ID3. In experiment A, the  $G_0$  dysgenic females were obtained by mating  $\pi IX$  reactive females with males heterozygous for inducer and reactive chromosomes providing the Binscy reactive X chromosome. In experiment B, the dysgenic females were obtained by crossing females of the wild-type reactive strain Cha and inducer males providing one or another of the three DcxF chromosomes containing the  $\pi I$  element. The Binscy and DcxF chromosomes contain crossing-over inhibitors to prevent recombination between ml carrying chromosomes and the homologous chromosomes of the target genome, TA. The TA genome was transmitted to  $G_1$  males that were mated individually with cn; e reactive females (cn and e are recessive markers of the second and third chromosomes, respectively). They were then hybridized with the  $\pi$  probe according to the squash blot procedure (20) to select individuals carrying sequences from the  $\pi I$  element. To eliminate translocations of the whole pUChsneo  $\pi I$  vector as a result of exceptional recombination, the wild-type female progeny of positive G<sub>1</sub> males were probed with pUC8. Any line having homology to this probe was discarded. Lines positive with the  $\pi$  probe and negative with pUC8 contained transposed copies of the marked  $\pi I$  elements. The chromosomal locations of these transposed  $\pi I$  elements were determined by probing flies of various  $G_2$  genotypes with  $\pi$ . Two transposed copies were recovered in one  $G_1$  male from  $\pi ID1$  in experiment B. One was located on the X chromosome ( $\pi ITX$ ) and the other was on chromosome 3 ( $\pi$ IT3).

Sequencing Reactions. The Sequenase kit (United States Biochemical) was used to determine the nucleotide sequence as follows. One-third of the microconcentrator-purified PCR product obtained as described above was denatured in 0.2 M NaOH and annealed with 50 pM primer (see Fig. 1, oligonucleotide c). Synthesis was performed with 3 pM  $[\alpha^{-32}P]$ dATP (3000 Ci/mmol; 1 Ci = 37 GBq) and the labeling mix was diluted 10 times. These experiments, including genomic DNA extraction and PCR amplification, were repeated twice for each DNA. Identical results were obtained in each case.

Other Procedures. All other procedures were carried out as described (19).

#### RESULTS AND DISCUSSION

The  $\pi I$  Element Cannot Transpose Autonomously. We have constructed an element,  $\pi I$ , to test whether I factors transpose via an RNA intermediate. This was made by inserting an intron-containing fragment into the second open reading frame of an active I factor (13, 14). An 896-base-pair fragment ( $\pi$ ) containing the second intron of the P factor was used (Fig. 1). The marked  $\pi I$  element was introduced into the germ line of a reactive strain by a standard P-mediated transformation technique (16, 17). Four transgenic lines were obtained from this experiment. In one of them ( $\pi IX$ ), the marked  $\pi I$  element was inserted on the X chromosome. In the other three ( $\pi ID1$ ,  $\pi ID2$ , and  $\pi ID3$ ) it was inserted on a third chromosome carrying the DcxF rearrangement. The  $\pi I$  elements in the transgenic lines did not show visible rearrangements (see Fig. 3, lanes X, D1, D2, and D3).

The marked I elements are unable to transpose in these strains (data not shown) because of the mutation caused by the insertion of P sequences into a coding region. They were therefore transferred to dysgenic females by appropriate crosses so that they could be complemented by transposing I factors (Fig. 2).

The Intron Is Accurately Spliced During Transposition of the  $\pi I$  Element. For each transgenic line,  $\approx 150~\rm G_1$  males were individually screened for homology to the  $\pi$  probe in experiments A and B (Fig. 2). From the  $\pi I$  element of line  $\pi ID1$ , two bona fide transposition events were isolated in the progeny of one  $\rm G_1$  male according to the procedure described in Fig. 2. The chromosomes carrying the transposed  $\pi I$  elements in this  $\rm G_1$  male were identified by screening its  $\rm G_2$  progeny for  $\pi$  sequences in squash blot experiments (20). Females of the genotype  $\rm TA/+;cn/cn;e/e$  and males of the genotype  $\rm +/Y;cn/cn;TA/e$  were positive while all  $\rm +/Y;TA/cn;e/e$  males were negative. This indicates that one transposed copy of the marked  $\pi I$  element is inserted on the X chromosome while the other is inserted on chromosome 3.

Restriction analysis of these chromosomes, called  $\pi ITX$  and  $\pi IT3$ , is shown in Fig. 3 (lanes TX and T3). Sequences hybridizing to the  $\pi$  probe are on Sal I fragments that are larger than the 7.1-kb fragment of the donor (Fig. 3a), whereas the 4.3-kb internal Ava I fragment of the  $\pi I$  factor has been conserved (Fig. 3b). These results confirm that the  $\pi I$  element transposed independently of flanking sequences from the P transformation vector. This appears to have happened twice in the same germ cell of a dysgenic female (see Fig. 2). The Pst I fragments that contain the intron in the  $\pi I$  donor element are  $\approx 60$  bp shorter in  $\pi ITX$  and  $\pi IT3$  than in the  $\pi I$  element (Fig. 3c). The sequence of the appropriate region of these fragments indicates that this difference results from precise removal of the intron in each case (Fig. 4).

Therefore, the  $\pi I$  element must have transposed via an RNA intermediate that was reverse transcribed after being processed. Reverse transcription is presumably the mechanism by which other LINE elements transpose. A full-length I factor RNA has been recently identified in the ovaries of

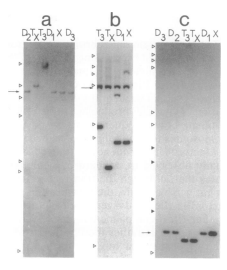


Fig. 3. Restriction analysis of various  $\pi I$  insertions. The following restriction enzymes were used: Sal I (a), Ava I (b), and Pst I (c). X is the  $\pi IX$  stock containing the original  $\pi I$  transgene on the X chromosome. D1, D2, and D3 indicate stocks with the three  $\pi l$ carrying DexF chromosomes. TX and T3 are the  $\pi$ ITX and  $\pi$ IT3 stocks isolated from experiment B in Fig. 2. The positions of  $\lambda$  23.6-, 9.6-, 6.7-, 4.3-, 2.3-, 2.0-, and 0.56-kb HindIII (open arrowheads) and λ 1.57-, 1.32-, 0.93-, and 0.84-kb *HindIII/Eco*RI fragments (solid arrowheads) are indicated. Arrows indicate genomic fragments corresponding in size to the control fragments of the pPI construct. Genomic DNA (3-4  $\mu$ g) was digested with various restriction enzymes, electrophoresed on 0.7% (a), 1% (b), and 1.3% (c) agarose gels, transferred to Amersham Hybond N filters, and hybridized with the Pst I/Pst I fragment covering most of the  $\pi$  marker (see Fig. 1). Hybridization, washing, and autoradiography followed the suppliers' procedures. In b, a common band corresponding to the Ava I fragment internal to the  $\pi I$  element (see Fig. 1) is observed in all lines. The other major band corresponds to the fragment included between the Ava I site of the P marker inserted within  $\pi I$  and the Ava I site to the right of  $\pi I$  in Fig. 1, which is either in the pUChsneo vector (X and D1) or in the target sequences flanking  $\pi I$  (T3 and TX), confirming that in  $\pi IT3$  and  $\pi ITX$ , the  $\pi I$  element transposed independently of pUChsneo sequences.

dysgenic females (21). It could be the intermediate required for transposition.

**Trans-Complementation of I Elements Can Occur.** From the experiments reported above, it appears that frequency of transposition of the  $\pi I$  element in the  $G_0$  dysgenic females (Fig. 2) was  $<10^{-2}$ . This is very low compared to the transposition frequency of active I factors usually observed in dysgenic females. In experiment A (Fig. 2), this frequency

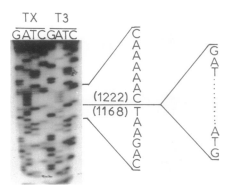


FIG. 4. Nucleotide sequence of the  $\pi$  fragments after transposition of the  $\pi I$  element. TX and T3 correspond to  $\pi$ ITX and  $\pi$ IT3 stocks isolated from experiment B (Fig. 2). The nucleotide sequence at the breakpoint of the deletion is given with coordinates of the P factor sequence (15); 1168 and 1222 are the end of the second exon and the beginning of the third exon, respectively.

was estimated by measuring the fertility of 11 samples of G<sub>2</sub> females of the genotype Binscy (TA)/+(r); cn(r)/cn(r); e(r)/cn(r)e(r). Such females contain only chromosomes from reactive origin and were obtained by crossing females of the reactive strain cn;e with 11 individual G<sub>1</sub> males. Acquisition of an active I factor by the Binscy (TA) chromosome in the G<sub>0</sub> dysgenic females should result in sterility of these G<sub>2</sub> females. The proportion of  $G_1$  males producing such sterile  $G_2$  females gives an estimate of the frequency of transposition. Four of the 11 chromosomes tested in this way had acquired at least one active I factor. This frequency of transposition is similar to that usually observed in standard dysgenic females (22). This indicates that, although the transposition frequency of the  $\pi I$  element was very low, the transposition frequency of active I factors in the G<sub>0</sub> dysgenic females was very high, suggesting that trans-complementation of transposition of elements of this type is rather inefficient. However, it can occur, and this provides a tool for further more detailed studies of the mechanism of transposition.

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