Polymorphisms on the right arm of yeast chromosome III associated with Ty transposition and recombination events

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ABSTRACT

The region of Saccharomyces cerevisiae chromosome III centromere-distal to the PGK gene is the site of frequent chromosome polymorphisms. We have sequenced this region from fragments of chromosome III isolated from three different yeast strains, GRF88, CN31C and CF4-16B. The sequence analysis demonstrates that these polymorphisms are associated with the presence of Ty and delta elements and defines a region of the chromosome which is a hot-spot for transposition events (the RAHS). The three strains can be arranged into a logical evolutionary series in which successive transposition and recombination events insert Ty elements and fuse them with consequent deletions of chromosome and of transposon sequences. The influence of such events on yeast genome evolution is discussed.

INTRODUCTION

Most laboratory strains of the yeast Saccharomyces cerevisiae contain ca. 30 copies (1) of the yeast retrotransposon, Ty (for review, see ref. 2). The general structure of these elements consists of a 5.2 kb unique 'epsilon' region flanked by two directly repeated 'delta' sequences of ca. 330 bp each which have identical, or nearly identical, nucleotide sequences. The sequence homology between delta sequences in the same element permits cross-over events to occur between them which results in the excision of the epsilon region together with one copy of delta, a 'solo delta' sequence being left behind in the chromosome. Such delta-delta recombination events appear to be relatively common since laboratory strains of yeast usually contain some 100 solo delta sequences (3). The distribution of Ty and solo delta elements within the yeast genome appears to be random (1). However, the preferential insertion of Ty elements into promoter sequences (4) and adjacent to tRNA genes (5,6) has been recorded.

Most Ty elements whose chromosomal context has been studied in detail are associated with some particular class of mutation for which the investigators selected. In order to examine the dynamics of the interaction...
of Ty with the yeast genome and to understand the factors involved in maintaining an equilibrium number of these transposons within it, an analysis of unselected events is required. Such an analysis is provided by the determination of the DNA sequence of yeast chromosome III which is being performed in our laboratories.

In the course of these sequencing studies, we have discovered two regions of chromosome III which appear to act as transposition 'hot-spots' (Fig. 1a). One of these, the left-arm hot-spot (LAHS) lies immediately centromere distal to the LEU2 gene. DNA sequencing (7) showed the LAHS to be flanked by the 5' ends of two tRNA genes, tRNA\textsubscript{Leu}\textsuperscript{3} and tRNA\textsubscript{Glu}. It contains the complete class II Ty element, Ty 1-17 (ref. 8), a number of delta fragments and a structure in which two solo deltas are inserted into one another. We interpreted this structure to be the result of two successive rounds of transposition and delta-delta recombination and suggested that the delta sequence itself may be a preferred target for transposition events. Our own data (8) and that of others (3,9) have shown that Ty and other repeated elements, such as tau (6) have a propensity to insert into the same region of delta DNA between base pairs 212 and 227. If delta is a preferred target for transposition, then the steady-state number of complete Ty elements would be critically dependent on events occurring within delta sequences. An equilibrium would be established between the deletion of Ty elements by delta-delta recombinations and promotion of new transposition events by the solo delta sequences left behind. Such a scheme has been described as the 'Ty cycle' (10) and in this paper we present a further examination of this cycle based on our sequence analysis of the right arm transposition hot-spot (RAHS).

The RAHS lies centromere-distal to PGK and we have sequenced this region from three different isolates of chromosome III. The data demonstrate that successive transposition and recombination events involving Ty and delta sequences are responsible for the polymorphisms found in this region of the chromosome.

**METHODS**

(1) Strains: the region of chromosome III analysed in this communication was recovered from three yeast strains. GRF 88 was derived from S288c and carries the his4-38 mutation (M. Rose, pers. comm.). The genealogy of S288c is described in ref. 11. CF4-16B was obtained from S.C. Falco. The strain is disomic for chromosome III and carries complementing his4C alleles. It was also derived from S288-related strains. CN31C is a segregant of a
cross between strain 248 (ref. 12) and YNN27 (ref. 13). It carries the four chromosome III markers from strain 248 and the ura3-52 allele from YNN27. The genealogy of strain 248 can be traced back through at least five crosses and its background includes strains from the Cold Spring Harbor Yeast Course, T. Manney and C. McLaughlin (J. Strathern, pers. comm.). The cryl-3 (and perhaps much of the right arm of the strain 248 chromosome III) originated in a strain obtained from C. McLaughlin. The full genotypes of these strains are given in Table 1.

(2) Plasmids: CB15A-2D was recovered from a library constructed in yeast shuttle vector YCp50 from GRF88 DNA partially digested with Sau3A by M. Rose and P. Novick (pers. comm.), using a probe from the centromere-distal end of plasmid 62B5-2D (Fig. 1b). Extensive Southern blot analysis (data not shown) demonstrated that this plasmid is homologous to the ring III chromosome from strain XGl-24b which has been cloned and mapped (12,14). Plasmids 31C-GF and 16B-GF were constructed by gap repair, in yeast, of a linearized plasmid containing sequences that flank the region of interest in the chromosome (15).

The plasmid used for gap filling, B52D-H9G-GF, was constructed in the shuttle vector YCp50 and contains the centromere proximal 1.8kb BamHI-XhoI fragment of plasmid H9G adjacent to the centromere distal 2.8 kb EcoRI-BamHI fragment of plasmid 62B5-2D in their chromosomal orientation (see Fig. 1b). The plasmid was linearized at the unique BamHI site which separates the chromosome III fragments by digestion with BamHI and used to transform strain CN31C or strain CF4-16B. Ura+ transformants containing the gap-repaired plasmid of interest were distinguished from those containing only the recircularized starting plasmid by isolating covalently closed circular DNA using the procedure of
Devenish and Newlon (ref. 16) and looking for large plasmids carrying sequences within the region of interest by Southern blotting and hybridizing with $^{32}$P labelled single copy from plasmid CB15A-2D. All of the Ura$^+$ transformants screened contained extrachromosomal plasmid and approximately half of the transformants carried the gap-filled plasmid. The chromosome III sequences in the plasmids were shown to be the same as those in the intact chromosome by comparing restriction maps of the plasmid with those deduced for the chromosome by Southern analysis.

**Southern Blotting**

Plasmid DNAs were purified, digested with various combinations of restriction enzymes and fractionated on 0.7% TBE-agarose gels (17). The fractionated DNAs were transferred to Hybond N by the procedure of Southern (18). The filters were probed with a Tyl-15 specific probe which comprised a 9.6 kb HindIII restriction fragment from the plasmid pMA91-301 (19) and labelled with $^{32}$P-dCTP (3000 Ci/mol) by nick translation (20). Filter hybridisation conditions used were as recommended by Amersham for Hybond N. After autoradiography, the Ty probe was removed from the filters and they were re-hybridised with a 0.6 kb BglIII-EcoRI fragment from 31C-GF.

**Sequencing of the $d_1$ delta element and its flanking DNA in CB15A-2D and H9GSTy**

The 1.4 kb XhoI-PvuII restriction fragments containing the $d_1$ element of Tyl-161 in CB15A-2D and the single Ty in 16B-GF, were isolated, end repaired with DNA polymerase I (Klenow fragment), and blunt end ligated into the SmaI restriction site of M13mpl8. The fragments were sequenced in both orientations using the 'Bandaid' technique of Dale et al. (21).

**Sequencing of the $d_2$ element and its flanking DNA from CB15A-2D.**

The 2.6 kb SalI restriction fragment spanning the two tandem Ty elements was isolated, and cleaved with BglII and PvuII. The 500 bp BglII-PvuII DNA fragment containing the $d_2$ sequence was sub-cloned into SmaI, BamHI cleaved M13mpl8 and mpl9 and sequenced in both orientations. The 2.6 kb SalI fragment was also cleaved with XhoI and the SalI-XhoI fragments were subcloned into SalI cut M13mpl8 and sequenced.

**Sequencing of the $d_3$ element and its flanking sequences in CB15A-2D and 16B-GF.**

The 2.7 kb EcoRI restriction fragment containing $d_3$ delta sequences was isolated, and cleaved with either BglII, ClaI or XhoI. The resulting fragments were subcloned into M13mpl8 and mpl9 cleaved with the appropriate combinations of EcoRI, BamHI, AccI and SalI and sequenced.

**DNA sequence analysis of the unique 31C-GF region.**

This region along with some flanking chromosome III DNA was isolated on
Figure 1 Transposition hot spots on chromosome III.

a) A map of the 200 kb circular derivative of chromosome III from strain GRF88 which has been cloned and mapped (ref. 14) LAHS and RAHS indicate the positions of the left arm and right arm Ty element insertion hot spots.

b) An expanded map of the region from strain CN31C containing the RAHS. Designations of the cloned BamHI fragments are given above the map. Fragments K3B and H9G are the same in the three strains examined in this report. The polymorphic RAHS reported in this paper extends from the EcoRI site shown in fragment 62B5-2D through the XhoI site shown in fragment HBGF. The pet18 mutation is a deletion of the region shown (ref. 38).

a 4kb XhoI fragment, and an overlapping 2 kb EcoRI fragment (see Fig. 2c).

The XhoI fragment was subcloned into the SalI site of M13mp19 and the EcoRI fragment was subcloned into the EcoRI site of M13 mp18. The two DNA fragments were sequenced in both orientations using the unidirectional deletion strategy of Henikoff et al. (22) except that Mung Bean Nuclease was used in place of S1 nuclease. The M13 mp18 subclones with BamHI and PstI, prior to ExoIII digestion.
Northern blotting and hybridisation

Total RNA was prepared from yeast cultures grown on 1 x 10^7 cells/ml in YEPD medium as described by Elder et al. (23). Total yeast RNA (10 µg) was subjected to electrophoresis on a 1.2% formaldehyde agarose gel and transferred onto Hybond N (Amersham). The filter was hybridised with the 1.4 kb XhoI-BamHI restriction fragment from 31C-GF (fig. 2c) labelled by nick translation. The electrophoresis, blotting and hybridisation procedures were as described by the Amersham 'Hybond N' booklet.

MATERIALS

Restriction endonucleases and DNA modifying enzymes were purchased from BCL, Bethesda Research Labs and Pharmacia. Reaction conditions were as recommended by the manufacturer. Hybond N and radionucleotides were from Amersham International. Other chemicals were purchased from BDH.

RESULTS

Isolation and characterisation of the RAHS from three different chromosomes III

The region centromere distal to the PGK locus on the right arm of chromosome III (Fig. 1) shows considerable restriction enzyme site polymorphism (12). Most of this can be attributed to the presence or absence of one or more Ty elements. It appears that this region of the chromosome is a 'hot spot' for transposition events and, therefore, it was designated the RAHS (for Right Arm Hot Spot). We have cloned the RAHS region from the chromosomes III of three different strains of Saccharomyces cerevisiae: GRF88, which is homologous in this region to the donor of the ring chromosome in the heterozygous diploid XG1-24B (ref 12), CF4-16B (a relative of S288C) and CN31C (which carries chromosome III from strain 248, ref. 12). Our analysis demonstrated that these forms of chromosome III contained, respectively, two, one and no Ty elements in the RAHS region. The evidence for this comes from restriction mapping and Southern hybridisation experiments and is described briefly below.

Plasmid CB15A-2D contains the RAHS region of the GRF88 chromosome III, cloned as an 18 kb Sau3A fragment which was derived from a partial digest of genomic DNA and inserted into the BamHI site of YCp50. The RAHS from the other two strains were also isolated in YCp50 using gap repair (see Methods). Plasmid 31C-GF contains a 21 kb RAHS insert from the chromosome III of strain CN31C, while 16B-GF contains a 26 kb RAHS fragment from strain CF4-16B. Each...
plasmid was analysed using a range of restriction enzymes, both singly and in combination, and the fragments containing Ty sequences determined by Southern hybridisation (18), using a nick-translated (20) HindIII fragment that contains the class I Ty element, Ty 1-15 (ref. 19).

Plasmid CB15A-2D was found to contain two class I Ty elements. The centromere-proximal (left-hand) member of the pair, we designated Ty 1-161 since it occupies the same position as the element of that name originally described by Kingsman et al. (24). Like the original Ty 1-161, the XhoI fragment carrying this element is longer than XhoI fragments carrying other class I transposons and by restriction analysis the element appears to have 1.2 kb insertion at its 5' (left) end. The centromere-distal Ty in the pair we named Ty 1-3R (see fig. 2). No sequences homologous to the Ty 1-15 probe were found in plasmid 31C-GF although prolonged exposure of the Southern blots did reveal some faint bands which subsequent sequence analysis (see below)

Figure 2 Detailed restriction maps of the RAHS from 3 yeast strains, showing Ty element related polymorphisms.
a) The RAHS of strain GRF88 from the plasmid CB15A-2D containing two tandemly linked Ty elements.
b) The RAHS of strain CF4-16B from the plasmid 16B-GF, which contains a single hybrid Ty element.
c) The RAHS of strain CN31C from the plasmid 31C-GF. This strain's RAHS contains 3 complete solo delta elements, a tRNA<sup>lys</sup> gene and an ORF of unknown function.

B = BglII, Ba = BamHI, C = Clai, E = EcoRI, H = HindIII, P = PvuII, S = SalI, X = XhoI.
Figure 3. Nucleotide sequence of the RAHS from strain CN31C. The region of plasmid 31C-GF sequenced is that shown in Fig. 2c. The solo delta sequences are underscored and the tRNA<sup>ys</sup> sequence is overscored.

showed to be due to the presence of solo delta sequences. Prolonged exposure of Southern blots from CB15A-2D and 16B-GF did not reveal any equivalent bands. BglII-EcoRI double digests of 31C-GF produced a 0.6 kb fragment which was not seen in similar digests of the other two plasmids. Further analysis, using this fragment as a probe, demonstrated that 31C-GF contains a 3 kb region of chromosomal DNA which is absent in both CB15A-2D and 16B-GF. Detailed maps for the RAHS region of all three inserts are presented in Fig. 2.

The RAHS from strain CN31C contains three solo delta elements and a tRNA gene
Figure 4. Autoradiograph of a Northern blot of total yeast RNA prepared from strain CN31C, probed with the 1.4 kb XhoI-BamHI fragment from plasmid 31C-GF (Fig. 2c and 4).

5'ATATC 3'; and the third, 31C-GF3 (bp 3838-4167) by 5'AGAAT 3' and 5'TGTGA 3'. This last solo delta is flanked on its centromere-proximal (left) side by the last 64 bp of another delta element arranged in the opposite orientation. This supports the idea that there has been considerable delta-delta interaction in this region via either transposition or recombination. Such delta-delta rearrangements probably explain the absence of chromosomal target site duplications at the ends of these elements. This absence might be the result of sequence degeneration, but this seems unlikely in the case of 31C-GF1 and 31C-GF2 both of whose sequences are highly conserved.

The LAHS region is flanked by the 5'ends of two tRNA genes (7) and others (5,6) have pointed out an association between Ty elements and such genes. A search for tRNA genes in the 31C-GF sequence revealed a tRNA_{Lys} at bp 3601 to 3674 (overacored in Fig. 3). This gene is embedded in a very A + T rich domain which is abutted by the 64 bp delta remnant 5' to the tRNA gene (see Fig. 3). This arrangement is consistent with the previously described association between yeast transposons and transfer RNA genes.

The RAHS from 248 contains a transcribed gene of unknown function

A search of the 31C-GF sequence for potential protein-encoding genes revealed a single open reading-frame (ORF) of 222 amino acids between bp 551 and 1228 and oriented towards the centromere (see Fig. 3). The sequence context of this ORF encourages the belief that it is expressed. Putative TATA
boxes occur at positions 1300 and 1330. Several CT tracts, which are associated with a number of yeast promoters (26), are found downstream of these TATA boxes. The 3' flanking region of the ORF contains two sequences which have been suggested to be yeast transcriptional terminators: first, TAGT at position 377 followed by a T-rich stretch and a TTT at position 309 (ref. 27); second, the sequence TTTTATA is found at position 410 (ref. 28). A Northern blot of total yeast RNA using a XhoI-BamHI fragment (bp 1 to 1410) from 31C-GF probe confirmed the transcription of this putative gene, revealing a transcript of ca. 1.0 kb in length (Fig. 4). The codon bias of the ORF is typical of a poorly expressed yeast gene (29). A search of the Doolittle (30) and PIR (31) protein sequence data banks failed to identify any known proteins with significant homology to the 31C-GF ORF.

Sequence analysis of the RAHS from GRF88 which contains two Ty elements

The restriction enzyme and Southern blot analysis described earlier demonstrated that GRF88 contains a deletion of chromosomal sequences in the RAHS region of chromosome III. In order to determine the nature of the deletion event it was necessary to determine the sequences flanking the tandem Ty insert in CB15A-2D and to compare them with the 31C-GF sequence.

The 1.4 kb XhoI-PvuII fragment containing the left (centromere-proximal) delta elements of Tyl-161 was isolated, end-repaired using Klenow polymerase and the resulting blunt ended fragment cloned into the SmaI site of M13mp18. The 'Bandaid' technique (21) was then used to sequence this DNA in both orientations. The sequence obtained is presented in Fig. 5a. It demonstrates that Ty 1-161 is not a variant class I element but is of normal length. The left delta of Ty 1-161 (d in Fig. 2) has lost its characteristic XhoI site and the XhoI site that was previously inferred to lie within the left delta is in fact contained within unique chromosome III DNA some 940 bp upstream of the 5' end of the left delta of Tyl-161. This delta is in the middle of the 31C-GF reading-frame at position 939 as recorded in Fig. 3.

The junction between the tandem Ty elements was investigated by isolating it on a 2.6 kb SalI fragment (see Fig. 2), and then sub-cloning it as a 0.5 kb BglII-PvuII fragment into M13mp18 and mpl9 cleaved with SmaI and BamHI. This sequence was confirmed using two further sub-clones of the SalI fragment which were obtained by XhoI digest. Fig. 5b gives 790 bp of sequence from the junction region and shows that Tyl-161 and Tyl-3R are joined by a single delta sequence (d in Fig. 2a) of 322 bp with no intervening chromosome III DNA.

The centromere-distal junction between the tandem pair of Ty elements and
chromosomal DNA was isolated on a 2.7 kb EcoRI fragment whose sequence is shown in Fig. 5c. It can be seen that the epsilon sequence of Ty 1-3R terminates in a complete delta sequence of 327 bp (d3 in Fig. 2a). This delta sequence is itself flanked by the last 180 bp of another delta sequence (Rd in Fig. 2a), suggesting that Tyl-3R may originally have been incorporated into chromosome III by transposition into, or recombination with, a pre-existing delta sequence. The 180 bp delta fragment is joined to unique chromosome III DNA at its right end and the sequence is identical to that beginning at position 3986 of 31C-GF. Thus strain GRF88 contains a deletion that extends from nucleotide 939 through 3986 of the 31C-GF sequence shown in Fig. 3 (see Discussion).

The RAHS from the CF4-16B chromosome III which contains a single Ty element

The 0.5 kb SphI - PvuII fragment from 16B-GF that contains the left junction of Tyl-16B was cloned into SphI, Smal digested M13 mpl8 and mpl9 and sequenced. It was found that this fragment had exactly the same nucleotide sequence as the equivalent fragment from CB15A-2D and contained 3C bp of chromosome III DNA from the SphI site to the 5' end of the left delta sequence of Tyl-161, the Tyl-161 left delta itself and the first 140 bp of the Tyl-161 epsilon up to the PvuII site. The 0.8 kb BglII - EcoRI fragment that contains the right junction of Tyl-16B was cloned into BamHI, EcoRI digested M13mpl8 and mpl9 and sequenced. Again, the nucleotide sequence of this fragment was found to be identical to that of the equivalent fragment from CB15A-2D. From the BglII site it contains 20 bp of the 3' epsilon region of Tyl-3R, the Tyl-3R right delta sequence (d3 in Fig. 2a and b), the delta fragment and then unique chromosome III DNA exactly as shown in the CB15A-2D sequence given in Fig. 5c.

The restriction mapping and sequence data indicated that the left delta and epsilon region of Tyl-16B, up until at least the PvuII site, are derived from Tyl-161 (the left Ty element of CB15A-2D). The right end of Tyl-16B, from at least the HindIII site, appears to be derived from Tyl-3R (the right Ty element of the tandem pair in CB15A-2D).

Figure 5. Nucleotide sequence from the RAHS of GRF88 contained in plasmid CB15A-2D.

a) The left delta (d1) of the Tyl-161 element and its flanking sequence. Left most XhoI-PvuII fragment in Fig. 2a.

b) The middle delta (d2) and the flanking Ty epsilon sequences (Fig. 2a).

c) The right delta element of Ty 1-3R (d3) and its flanking sequences, up to the right most EcoRI site in Fig. 2a.
Figure 6. A mechanism to explain the generation of the three Ty element associated polymorphisms characterised.

a) The earliest structure is assumed to be that of the RAHS contained in plasmid 16B-GF from strain CN31C. Two independent transposition events inserted Ty 1-161 and Ty 1-3R into the CN31C RAHS. The two Ty elements are separated by ca. 3 kb of DNA unique to plasmid 16B-GF.

b) A recombination event between the two proximal delta elements brings Ty 1-161 and Ty 1-3R together. This forms the tandem Ty element structure characterised in plasmid CB15A-2D, containing DNA from the RAHS of strain GRF88. The ca. 3 kb of intervening DNA is lost from the RAHS of GRF88.

c) Recombination between the epsilon regions of Ty 1-161 and Ty 1-3R generates the hybrid Ty1 element (Ty 1-10B) found in the RAHS of strain CF4-163. This event could occur from either stage a) when the Ty1 elements are separated or at b) where they are in tandem array.

DISCUSSION

The exact relationship between the three strains of *S. cerevisiae* whose chromosome III sequences we have analysed in the RAHS region is unknown;
Figure 7. Alignment of the delta element sequences from the three Ty element deltas from the RAHS of GRF88 (Fig. 2a).

1 = d1 (31C-GF1); 2 = d2 (31C-GF2) and 3 = d3 (31C-GF3) Nucleotides differing between one or more of the delta elements are shown. The consensus delta sequence is marked below. The probable position of the recombination event between the proximal delta sequences of Ty 1-161 and Ty 1-3R is underscored.

However, the sequence data obtained suggests the following evolutionary relationship.

Strain CN31C contains no complete Ty elements in the RAHS region of chromosome III. Nevertheless the presence of three solo delta sequences in the CN31C RAHS, none of which is bounded by a target site duplication and one of which is inserted into a delta fragment, suggests that even in this strain the RAHS has been the site of a number of successive transposition events followed by delta-delta recombinations. The CN31C RAHS contains two functional genes, the 31C-GF ORF and tRNA\(^{\text{lys}}\). We propose the gene name \textit{CTR1} for the ORF, standing for chromosome three reading-frame.

The RAHS from strain GRF88 can be considered as being derived from strain 248 via two independent transposition events. One transposition inserted Tyl-161 into chromosome III at the position numbered 939 in the 31C-GF sequence (Fig. 3), meaning that the element was actually inserted into the coding sequence of CTR1. This may be considered an unusual event given the reported preference for Ty elements to insert into promoter sequences (4). However, we would reiterate that we have examined unselected events whereas previous workers have investigated Ty insertions associated with particular mutations and so may have produced a biased picture. The second transposition into the CN31C RAHS inserted Tyl-3R into a pre-existing solo delta sequence at position 3986 in the H9G sequence (Fig. 3). Our sequence analysis of the LAHS region of chromosome III (ref. 7) had suggested that delta sequences may themselves be preferred targets for transposition events and Tyl-3R provides

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further support for that hypothesis. Our data (this paper and ref. 7) and that of others (3,3) delimits the delta transposition target as lying between base pairs 150 and 230. This region contains the delta element promoter and provides a further example of Ty elements inserting into yeast promoter sequences.

After the two transposition events, the RAHS would have contained two class I Ty elements (Tyl-161 and Tyl-3R) separated by ca. 3 kb of unique chromosome III DNA containing part of CTR1, two solo delta sequences and the \text{tRNA}^{lys} gene. A recombination event between the right delta sequence of Tyl-161 and the left delta sequence of Tyl-3R could then have occurred fusing the two Ty elements via a single, central, delta sequence and producing the tandem Ty pair found in the RAHS of strain GRF88. Fig. 6 diagrams this sequence of events. A comparison of the sequences of the three deltas contained within that tandem Ty pair (Fig. 7) suggests that the recombination event between the proximal deltas of Tyl-161 and Tyl-3R occurred between base pairs 238 and 244, close to or at the transcription start site. In addition to fusing the two Ty elements, this recombination event also resulted in the deletion of ca. 3 kb of chromosome III DNA. The consequences of this deletion will be considered after the derivation of the RAHS from strain CF4-16B has been explained.

CF4-16B contains a single Ty element, Tyl-16B in its RAHS region. This Ty appears to be the product of a recombination event between the epsilon regions of the tandem Ty pair contained within the chromosome III of GRF88. Strain CF4-16B represents, therefore, the end-point of our proposed evolutionary series. The restriction mapping and sequence analysis of Ty 1-16B presented in the RESULTS section strongly suggests that a recombination event occurred between Tyl-161 and Tyl-3R in the region between the first PvuIIsite in the epsilon sequence of Tyl-161 and the following HindIII site. We believe this is the first example of an epsilon-epsilon recombination event to be reported.

Such recombinations are difficult to detect especially if, as in this case, they occur between two members of the same class of Ty elements. However, a number of factors suggest that epsilon-epsilon recombinations may indeed be rare events. For instance, no element with a restriction pattern which indicates that it is a hybrid between a class I and class II Ty has been reported. Epsilon-epsilon recombinations most frequently result in deletion of chromosomal DNA or produce a translocation. Such events are likely to be selected against and yeast may even have some positive means of repressing them. The epsilon-epsilon recombinations we reported here occurred between
two Ty elements fused in tandem and so produced no deleterious genetic effects. Delta-delta recombinations may occur at a higher frequency than those between epsilon sequences since they usually involve the two delta sequences of a single Ty element and, therefore, are genetically innocuous. However, as we have demonstrated in this paper, recombination between delta sequences on different Ty elements may occur and result in a deletion of chromosomal sequences. Rothstein (33) has presented evidence that the frequency of even delta-delta recombinations is repressed by the action of a chromosomal gene, EDR.

The assignment of epsilon-epsilon recombination as the event which converted the adjacent Ty pair (Fig. 6a) or the tandem Ty pair in the RAHS of GRF88 (Fig. 6b) to the single Ty element of CF4-16B (Fig. 6c) relies on evidence involving sequence differences between the two class I Ty elements, Tyl-161 and Tyl-3R. It is instructive to compare the sequence of these two elements from chromosome III with those of the two class I Ty's which have been sequenced previously, Ty 912 (ref. 34) and Ty109 (ref. 35). The centromere-proximal member of the tandem Ty pair, Tyl-161, differs from Ty 912 at only 7 nucleotides within the sequenced region (341 bp) shown in Figs. 5a and b, whereas it differs from Ty109 at 81 nucleotides. In contrast, the nucleotide sequence of Tyl-3R is more similar to that of Ty109 than that of Ty912. In the 538 bp region shown in Figs. 5b and c, it differs from Ty 109 at 14 nucleotides but from Ty 912 at 93 sites. Tyl-161 and Ty912 are easily distinguished from Tyl-3R and Ty109 since the former pair contain a HindIII and a SalI site which the latter two lack. These differences permit the division of class I yeast transposons into two sub-classes, Tyla (comprising Tyl 1-161 and Ty 912) and Tylb (comprising Tyl-3R and Ty 109).

The delta-delta recombination event which fused Tyl-161 and Tyl-3R resulted in the deletion of 3 kb of unique chromosome III DNA which contained a transcribed open reading-frame (the 31C-GF ORF) and a tRNA^lys^ gene. Since this deletion did not lead to the death of the cell, neither of these genes is essential. This is consistent with the report of Goebl and Petes (36) that 60% of insertions into the yeast genome they studied had no phenotypic effect.

In other regions of chromosome III we have found a number of open reading frames whose deletion or interruption is without lethal effect (ref. 37 and manuscript in preparation). Toh-e el al (38) have reported that the pet18 mutation is associated with another deletion (see Fig. 1b) just centromere distal to Tyl-3R. The pet18 mutation is pleiotrophic in its effect and confers the following phenotypes on yeast cells which harbour it: (i) respiratory
deficiency, (ii) inability to maintain killer dsRNA, and (iii) temperature sensitive growth. Sequence comparisons between pet10 and PET10 strains showed that the deletion removed three open reading frames. All of these data indicate a remarkable level of redundancy in a genome which is less than four times the size of that of E. coli. Nevertheless, most or all of these non-essential genes may confer some selective advantage on yeast and investigators must use more sensitive tests (e.g. 39) in order to determine the magnitude of that advantage.

Fig. 6 shows that the delta-delta recombination must have excised the 3 kb fragment of chromosome III DNA on a circular molecule which contains either a solo delta sequence or an intact Ty element. Such a molecule represents a new class of translocatable element since it may be re-inserted into the yeast genome, at any one of at least 160 homologous sites, by a subsequent delta-delta recombination event. If the excised fragment contained an ARS element the probability of its translocation would be increased since it could replicate as an episome before reinsertion into a chromosome. That would not arise in this particular case since there is no functional ARS in the RAHS region (14,40).

The possibility of such translocations offers a new view of the association of tRNA genes with Ty sequences. Translocation of the portion of chromosome III which has been deleted in strain GRF68 would move the tRNA<sup>lys</sup> gene to a new chromosomal location. A subsequent mating with a partner without the deletion would result in the duplication of the tRNA<sup>lys</sup> gene. It is possible, therefore, that tRNA genes are often located near Ty elements since this association has served to amplify and spread those genes through the yeast genome and that such amplifications have been selectively advantageous. This is but one example of the likely importance of Ty elements in yeast genome evolution (41). This analysis of a transposition hot-spot from the right arm of chromosome III has demonstrated that Ty insertions and recombinations are a major cause of chromosome polymorphism in yeast.

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REFERENCES

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