Selection on the Genes of *Euplotes crassus* Tec1 and Tec2 Transposons: Evolutionary Appearance of a Programmed Frame-shift in a Tec2 Gene Encoding a Tyrosine Family Site-Specific Recombinase

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Received 24 June 2002/Accepted 5 September 2002

The Tec1 and Tec2 transposons of the ciliate *Euplotes crassus* carry a gene for a tyrosine-type site-specific recombinase. The expression of the Tec2 gene apparently uses a programmed +1 frameshift. To test this hypothesis, we examined whether the frameshift has been under purifying selection in Tec1 and Tec2. Each element carries three genes, and each has evolved under purifying selection for the function of its encoded protein, as evidenced by a dearth of nonsynonymous changes. This distortion of divergence is apparent in codons both 5′ and 3′ of the frameshift site. Thus, Tec2 transposons have diverged from each other while using a programmed +1 frameshift to produce recombinase, the function of which is under purifying selection. What might this function be? Tyrosine-type site-specific recombinases are rarely found in eukaryotes, and Tec elements also encode a widespread transposase. The Tec recombinase might function in transposition, resolve products of transposition (bacterial replicative transposons use recombinase or resolvase to separate joined replicons), or provide a function that benefits the ciliate host. Transposons in ciliated protozoa are removed from the macronucleus, and it has been proposed that the transposons provide this “excisase” activity.

We have studied the evolution of ciliate Tec transposon sequences to compare their evolution in *Euplotes crassus* to that of TBE1 transposons in *Oxytricha* ciliates (49) as well as to examine the appearance of a frameshift site in a Tec2 gene (22). We have confirmed that this frameshift has been a functional mechanism, and we demonstrate that this gene and its homolog in Tec1 elements encode a tyrosine-type site-specific recombinase.

The Tec and TBE elements encode a DDE transposase (conserved Asp, Asp, Gln residues in the active site) (10) and appear to be type II elements (“cut-and-paste” or “DNA-mediated” elements) (5). The evolution of eukaryotic type II transposons has proved difficult to understand: purifying selection for transposase function is not expected to act on the transposase genes of eukaryotic cut-and-paste transposons within a host population and is generally not found (36, 37, 48, 49). However, TBE1 transposons are a notable exception to this expectation. TBE1 transposon genes have evolved under strong selection for the function of their encoded proteins within their host ciliates (49); that is, nonsynonymous—or missense—mutations have been selected against and so removed from the pool of transposons in the host genome, while synonymous mutations have accumulated unabated. The reason for this selection is at present unknown, but it has been proposed that TBE1 transposons are under selection for a specific host function (28, 46, 49). Ciliated protozoa are unique transposon hosts because they carry two types of nuclei, a micronucleus and a macronucleus. The macronucleus is a terminally differentiated version of the micronucleus and provides all of the transcripts necessary for cell growth. When a new macronucleus differentiates, all ~4,000 TBE1 insertions are removed, at least some by precise excision (for a review, see reference 28). This process regenerates many genes that were inactivated in the micronucleus due to TBE1 insertions. Developmental excision is, in essence, “wholesale” somatic reversion of germ line insertion mutations. Our proposal was that the three TBE1 genes are selected for function because they are responsible for the excision of TBE1 insertions during differentiation. This proposal is no longer tenable, because these genes are not detectably expressed during macronuclear development (see below).

The Tec1 and Tec2 elements of *E. crassus* are ciliate transposons comparable to TBE1 in their general features. They are present in high copy numbers and are precisely excised during macronuclear development. Tec genes are not expressed at high levels sufficient to excise the ~12,000 Tec1 elements and ~12,000 Tec2 elements (24). Both Tec1 and Tec2 elements are 5.7-kb elements and have identical organization of their three genes and 690-bp inverted terminal repeats (ITRs). The encoded protein sequences of Tec1 and Tec2 are easily aligned but are only ~40% identical. The noncoding sequences within the ITRs have diverged to the point at which they cannot be aligned; thus, Tec1 and Tec2 diverged from each other a very
FIG. 1. Evidence of the action of purifying selection on the genes of Tec1 and Tec2 elements. (A) Window analysis of divergence in the ORF 2A-ORF 2B region of Tec2. Divergences ($d_s$ and $d_n$) were calculated in the three forward reading frames for all 10 pairwise comparisons among five aligned Tec2 sequences by using a 150-bp window and scanning across the alignment in one-codon increments. Ratios of the averaged $d_s$ and $d_n$ values are plotted (on a logarithmic scale) for each window and each reading frame; frame 1 is the open frame of ORF 2A, and frame 2 is that of ORF 2B. ORF 2A and ORF 2B are drawn to the scale of the plot at its lower boundary. The nucleotide sequence in the immediate vicinity of the putative frameshift is shown, with codons in the two reading frames delimited by tick marks; relevant start and stop codons are underlined (ORF 2B) or overlined (ORF 2A). The remarkably low $d_s/d_n$ ratios in out-of-frame intervals is presumably due to selection on coincident in-frame intervals. In sequence maps for Tec2-1 and Tec1-1, sequences are depicted to scale, and ORFs (bars, labeled) and ITRs (arrows) are shown. Regions sequenced from homologous cloned elements are represented by lines aligned beneath their respective “type” elements. Divergence analyses were performed with pairs of sequences drawn from alignments made with ClustalW (43). (B) Scatter plot of synonymous and nonsynonymous divergences ($d_s$ and $d_n$) calculated for all pairwise comparisons among sequences within alignments of five ORFs: Tec1 ORFs 1, 2, and 3 and Tec2 ORFs 2A and 2B. Divergences were calculated by the method of Nei and Gojobori (33) as implemented by Ina et al. (21), but with the genetic code of Euplotes (17). Divergence in the absence
long time ago and have since evolved into two quite independent families of elements.

The Tec and TBE elements are not demonstrably related except by the inclusion of their transposases in the IS630/Tc1/ 
Mariner subfamily of the D,D35E transposase superfamily (10). That is, Tec and TBE transposases do not form a 
ciliate clade of transposases: they are as like Tc1 or IS630 as they are 
like each other. Nonetheless, both Tec and TBE1 elements 
carry three genes (11, 22), whereas eukaryotic type II trans-
posons typically carry only a transposase gene (5). Other 
than the transposase genes, the other Tec1 or Tec2 and TBE genes 
are unrelated.

Oddly, in all Tec1 elements, open reading frame (ORF) 2 is 
a single ORF, while in all Tec2 elements, ORF 2 is divided by 
a frameshifting single-base-pair insertion into ORF 2A and 
ORF 2B. Jahn et al. (22) proposed that the complete Tec2 
gene product is made by a programmed frameshift. Here we 
demonstrate that the two sections of ORF 2 have been under 
selection for protein function since the appearance of the 
frameshifting insertion, indicating that a programmed +1 
frameshift has been used to produce the protein. We also show 
that ORF2 encodes a tyrosine family site-specific recombinase 
(34). Site-specific recombinases, of either the serine or the 
yeastine families, are extremely rare in eukaryotes, and Tec 
elements are, to our knowledge, the first eukaryotic type II 
transposons found to encode a site-specific recombinase. Ja-
cobs et al. (21a) report that the newly described Tec3 elements 
also encode such an enzyme. In prokaryotes, recombinases are 
associated with transposons that transpose replicatively, form-
ing cointegrates that the recombinase subsequently resolves. 
That Tec elements possess a recombinase suggests that Tec 
elements might transpose replicatively, a complicated process 
in organisms with linear chromosomes.

MATERIALS AND METHODS

Tec element DNA sequences were determined as described by Jahn et al. (22). 
The partial ORF 2 sequence from Tec1-3 was not analyzed. Figure 1A indicates 
the sequences that we have analyzed: a single complete sequence each of a Tec1 
element and of a Tec2 element, multiple sequences of all three Tec1 genes, and 
the sequences determined here are AF159907 to AF159921 for individual ORF 
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a full-length ORF 2 recombinase, the function of which was under selection.

Since this analysis detected only selection that has acted in the evolutionary past, the gene may no longer be under selection now. The divergence analysis of ORF 2B (Fig. 1B) shows two clusters of points representing two sets of divergences (d_3), ancient and intermediate (d_4 values, ~0.3 and ~0.12, respectively). Selection thus has acted during both ancient evolution and more recent evolution of this ORF. Since this data set lacks recently diverged elements, we cannot whether selection has acted even more recently. Several of the ORFs in the full data set are clearly damaged: of 20 Tec gene sequences analyzed, 6 have stop codons in frame, in either the original frame or following a frameshifting insertion or deletion (other than the programmed frameshift in Tec2 ORF 2). These particular damaged genes are no longer under selection: as soon as a gene suffers a disabling mutation, it ceases to be under selection and will diverge with a subsequent d_4/d_3 ratio of 1.

Homology of Tec ORF 2 proteins to tyrosine-type site-specific recombinases. We had been unable to identify the Tec ORF 2 protein function on the basis of homology. However, in a PSI-Blast search (2) of public databases with a profile of the Tec1 and Tec2 ORF 2 proteins, the best match was a Bergeyella integrase (E value, 0.006). The Bergeyella integrase is a member of a diverse set of recombinases, named “tyrosine-type” after the tyrosine residue that forms a transient covalent bond with the DNA substrate, described in detail by Nunes-Duby et al. (34). The family includes phage integrases, yeast FLP recombinases, the transposition function (int) of Tn916, and XisA of the heterocyst-excised nifD gene of Anabaena. A profile representing the Bergeyella protein and the two Tec proteins matched several other recognized family members aligned by Nunes-Duby et al. (34). The Tec sequences were aligned (Fig. 2) with a well-studied member of each functional subfamily defined by Nunes-Duby et al. (34) as well as with best matches from the profile search. We also included the protein sequence from the recently discovered Tec3 element (21a). The alignment clearly shows the homology of the Tec proteins to the recombinase family. For instance, the Tec proteins share the conserved active-site tyrosine and basic residues in boxes I and II. Further, the regions of greatest similarity between the Tec1 and Tec2 proteins correspond to the most conserved regions of the family (the boxes” and “patches” defined by Nunes-Duby et al. [34]) (Fig. 2), a finding supporting the inferred homology of the ORF 2 proteins to the tyrosine recombinases. While the Tec3 protein is not particularly similar to the Tec1 and Tec2 proteins (21a), it is clearly a member of the family overall.

Very few eukaryotic tyrosine recombinases are known, making the Tec genes nearly unique. The other known eukaryotic tyrosine recombinases are the FLP enzymes of yeasts, which form a clade very diverged from the rest of the recombinases (34); baculovirus VFL-1 (50); and the recently characterized recombinases of the DIRS retrotransposon family (14). Do eukaryotic tyrosine recombinases form a clade? The Tec recombinases do not match the FLPs at all and only match VFL-1 strongly in the box II region immediately adjacent to the catalytic tyrosine (Fig. 2). There is also no particular similarity to the DIRS recombinases. These findings fail to support the existence of a unique eukaryotic subfamily of recombinases and suggest that there have been multiple transfers of prokaryotic recombinases into eukaryotic lineages.

DISCUSSION

We find that the three Tec element genes have evolved under selection for the functions of their encoded proteins (Fig. 1B). The Tec2 ORF 2A-ORF 2B protein product is produced by a programmed frameshift (Fig. 1A) and is a tyrosine site-specific recombinase (Fig. 2).

What selection has acted on Tec transposon genes? It has been difficult to explain the selection that we have detected on Tec genes (Fig. 1B) and the selection on TBE1 genes (11, 29, 49). Genes must be expressed in order for selection to act, but very little, if any, expression is seen for these genes. Jarzewska et al. (24) found only very low levels of Tec gene transcripts in exconjugant cells: ~60 polyadenylated messages per cell, enough to support Tec transposition in the germ line. TBE1 genes are expressed at undetectably low levels (K. Williams, T. G. Doak, and G. Herrick, unpublished results). Given the lack of expression, it is difficult to formulate a function for the transposon genes that would lead to selection.

We have considered two explanations (not mutually exclusive) for the selection on Tec and TBE1 genes. Either transposon genes are under selection because elements with functional genes transpose more often than do elements with inactivated genes (selection for transposition), or the genes are selected due to a host benefit that they provide (selection for host fitness).

A host fitness explanation for TBE1 selection was proposed by Witherspoon et al. (49). That is, functional TBE1 genes increase the fitness of host cells that carry them, relative to host cells that carry defective TBE1 genes, and so functional TBE1 genes are more likely to be replicated. In particular, Williams et al. proposed that TBE1 transposase catalyzes the precise excision of TBE1 genes during macronuclear development (46). Precise excision of a TBE1 element (or Tec element or most other internal eliminated sequences) removes the element and one flanking target site duplication (for a review, see reference 28) and restores functional genes in the macronucleus. Thus, the selection acting on TBE1 genes would be based on their essential role in precise excision (49). In such a model, the “price” of selection is “paid for” in reduced host fitness. Selection for Tec functions can be imagined in the same fashion (28). However, the low levels of both Tec and TBE1 gene expression seem to rule out this explanation: for both Tec and TBE1, there are thousands of elements to be excised, and it seems implausible that so few transcripts could provide enough transposase to catalyze massive precise excision (24). Furthermore, Tec excision products do not appear to be formed by a transposase-like activity (23, 27), as is also true for the intermediates and products of TBE1 excision (9; K. Williams and T. G. Doak, unpublished data). It is possible that both Tec and TBE1 genes were under selection for excision functions until recently but that now their hosts have independently assumed alternative excision processes and the transcription of thousands of Tec and TBE1 elements has been suppressed (a refinement of the “invade, bloom, abdicate, and fade” model of Klobutcher and Herrick [28]); however, this possibility also seems unlikely.
FIG. 2. Alignment of the sequences of the proteins encoded by the Tec1 and Tec2 ORF 2 genes and the Tec3 ORF 1 gene with sequences of tyrosine-type site-specific recombinases. Only the C-terminal sections of the Tec proteins are aligned, with truncated recombinase sequences limited to their catalytic domains; the position of the first residue is indicated, as are the numbers of residues removed from the C terminus, if any. Known recombinase family members aligned are of two kinds: (i) several well-studied members (Int, Cre, VLF-1, XerD, Tn916, XisC, and FimB) and (ii) several recognized members with particularly strong matches in a PSI-Blast search with a profile representing the proteins of Tec1, Tec2, and Weeksella (Bergeyella) zoohelcum (W.zooTnpA): PL2, phiCTX, Clostridium butyricum (C. butyricum), Methanococcus jannaschii (M. jannaschii), W.zooTnpA, and Ye24-shuf; two good hits—RP2 and Int-I3—are not shown in the final alignment because they lengthened the alignment unnecessarily. The sequences are named as described by Nunes-Duby et al. (34). Alignment was achieved in consultation with S. Nunes-Duby (personal communication), yielding a set of five aligned sequences (the Tec1 and Tec2 proteins, Int, Cre, and VLF-1); this core alignment was used as a profile to align the remaining sequences by using ClustalX (42). The alignment was manually adjusted to conform in most cases to the alignment of Nunes-Duby et al. (34). The Tec3 sequence was later aligned against this entire alignment by using ClustalX (42). The two boxes and three patches of strong conservation noted by Nunes-Duby et al. (34) are enclosed in boxes; patches I and III were assigned by S. Nunes-Duby (personal communication). The three absolutely conserved positions (including the catalytic Y) are indicated by asterisks. Residue symbols have been colored to indicate similarity between family members. Colors were assigned to residues as follows. First, a consensus was calculated for each position in the alignment (not shown); the consensus was either an amino acid or an amino acid similarity group (31 or 50%, respectively, as described by Nunes-Duby et al. [34]). Second, based on this consensus, residue colors were designated for alignment positions with five or more nongapped residues. The exchange groups of Dayhoff et al. (7) used for the consensus calculation were SPAGT, RKH, FYW, DEQN, and LIMV. Residues were colored according to chemical type when they contributed to the consensus or fell into a similarity group with the consensus residue. Colors of chemical types are as follows (from reference 34): hydrophobic (I, L, or V; F, Y, or W; M; and P), pink; hydrophilic (S or T and Q or N), green; acidic (D or E), magenta; small (G and A), yellow; and basic (K, R, or H), cyan. The residues of the conserved RRY triad are represented by white letters within dark blue (R) and dark red (Y) backgrounds.
Alternatively, selection could be directed at transposition functions by one of two mechanisms. First, preferential cis action of transposase on the particular element that encodes it (a known mechanism in prokaryotes; see, e.g., reference 8) would impose selection (26). However, it is difficult to imagine a molecular mechanism in eukaryotes whereby proteins encoded by one of many nearly indistinguishable DNA elements and then translated in the cytoplasm would act preferentially on those particular elements in the nucleus. In contrast, this mechanism appears to function with retroelements, whereby a reverse transcriptase molecule acts in the cytoplasm on the retroelement RNA molecule that encoded it (44). Thus, retrotransposon transposition genes are generally found to be under strong selection (see, e.g., references 19, 25, 31, and 39; D. J. Witherspoon, unpublished results), due to this selection for transposition. While it remains possible that Tec and TBE elements transpose via an RNA intermediate, there is no evidence against the assumption that they are simple type II DNA elements.

A second situation that leads to selection for transposition functions has been described by Witherspoon (47, 48). The force of the selection is dependent on the variance in the proportions of wild-type elements in individual hosts; simply stated, in hosts with more wild-type elements, transposition is more frequent, so that wild-type elements are preferentially transposed. This population structure model can explain selection on the Tec transposition functions under certain conditions, in particular, if the rate of element turnover (loss, balanced by transposition) is sufficiently high. Conditions exist under which a small number of competent elements would be maintained in a large pool of mutant elements, consistent with the large number of damaged Tec elements that we have found. This selective force is very weak in the presence of lower turnover rates, and in such instances, all elements will become mutant, as has been observed for Mariner elements (see, e.g., reference 37). We have not determined the rate of Tec or TBE1 transposition but plan to do so for TBE1 as a test of this population structure model (Witherspoon, unpublished).

Programmed +1 frameshift in the Tec2 ORF 2 gene. Jahn et al. (22) proposed that the Tec2 gene was expressed as a single fusion protein, colinear with the homologous Tec1 protein. A +1 frameshift in the overlap of ORF 2A and ORF 2B was proposed that skips one of the A’s in a run of three, adjusting the reading frame to that of ORF 2B and avoiding an immediate TAG stop codon (Fig. 1A). The resulting Tec2 fusion protein would have the same amino acids (KN) at the shift site as the Tec1 protein at the homologous position. The Tec2 elements share the same sequence, AAATAG, with an “extra A,” indicating that this sequence and the need for a frameshift were present in their most recent common ancestor. Descendants of Tec2 ORF 2A and ORF 2B have diverged from that ancestor under selection for the function of the encoded recombinase. Thus, the evolutionary pattern of Tec2 recombinase gene evolution is unique proof of a functional programmed frameshift.

Further evidence of functional frameshifting in the genus *Euplotes* was recently documented for several genes, those encoding the p43 telomerase subunit (1) and two protein kinases, *Eopkar* (40) and *Eondr2* (41). Most recently, Z. Karamysheva et al. (submitted for publication) identified two functional +1 frameshift sequences in an *E. crassus* gene for the telomerase protein TERT. In all five examples, amino acid homology between a known protein and the candidate frameshifted protein is evidence for a programmed frameshift. The reasoning is that the similarity would not have persisted in the absence of selection for function of the encoded protein, a notion which in turn implies that the full-length protein was expressed via the programmed frameshift. For p43 and TERT, a protein of the predicted full length is observed, and a predicted p43 peptide has been sequenced.

A common feature of sequences that promote frameshifting is the ability to cause the ribosome to pause while translating a codon paired to a tRNA that also can pair with the overlapping codon in the new frame (for reviews, see references 12 and 13). This codon is often located in a string of identical nucleotides (in the present example, a string of three A’s) (Fig. 1A), allowing pairing in the two different frames (with AAA or AAU). A common pause-causing sequence is an in-frame stop codon terminating the first ORF just 3’ of the shift codon (see, e.g., references 6, 16, 30, and 45). The *Euplotes* +1 frameshift consensus, now AAATAA/G (41; Karamysheva et al., submitted), fits this pattern. Another feature that can cause pausing is a 3’ segment with a strong secondary structure; however, we have not found convincing structures in the ORF 2 gene sequence. A codon that requires a rare tRNA just before the shift can also impede translation and stimulate shifting, but there is no rare codon near the end of ORF 2A (see reference 20 for *Euplotes* codon usage tables).

Regardless of the details, it is clear that a robust mechanism for frameshifting functions in euplotids. Hence, the appearance of a frameshift mutation in Tec2 ORF 2 might have been effectively silent. It could not have been very deleterious, since the same purifying selection that has maintained the surrounding coding sequence would have eliminated it. Most frameshift sequences represent the independent mutation of nonhomologous sequences to analogous frameshifting signals. Since such signals can be quite short, random mutation is likely to produce them in genes at an appreciable frequency. Thus, a frameshifting sequence may serve no function at all. Note that Tec1 elements, which lack the ORF 2 frameshift, have been as successful as Tec2 elements. Alternatively, the frameshift may have a function such as producing stoichiometrically balanced levels of full-length recombinase and the N-terminal fragment, with the fragment modulating the activity of the recombinase. Many programmed frameshifts used by transposons serve such functions (for a review, see reference 4).

We have considered two alternative explanations for the frameshift and conservation that we have seen in the Tec2 ORF 2A-ORF 2B gene. The first explanation is that the Tec2 ORF 2A-ORF 2B gene is now expressed as a two-peptide recombinase by translational reinitiation in the ORF 2B frame; given the mounting number of frameshift examples in euplotids, we do not favor this explanation.

The second explanation is that gene conversion patched an inactivating frameshift mutation into all modern Tec2 elements, after they had diverged under selection, and that modern Tec2 elements are inactivated by this frameshift. Such a pervasive directional gene conversion seems inherently unlikely, but our data further rule it out. The proposed gene
conversion would have erased differences between the Tec2 elements near the frameshift site, but there are base pair differences within 15 nucleotides of the frameshift site on either side, and nucleotide diversity is not reduced near the frameshift site (data not shown). If gene conversion were so frequent, then it should also have acted on the rest of the Tec sequence, but we still saw evidence of selection, which would have had to accumulate after the proposed gene conversion. Therefore, we must still propose an ancestral Tec2 under selection for the function of ORF 2A-ORF 2B with the ancestral frameshift.

**Eukaryotic transposons encoding a site-specific recombinase.** The ORF 2 gene of Tec elements encodes a site-specific recombinase (Fig. 2), while the ORF 1 gene encodes a conventional DDE transposase (10). Given that the detected purifying selection might indicate that both of these genes are required for Tec transposition, we now consider how such transposition might occur. A tyrosine recombinase performs the transposition reaction (in the absence of a DDE enzyme) for members of the bacterial Tn916 family (35) and may as well for the eukaryotic retroelement DIRS (14). We are unaware of any eukaryotic type II transposon, other than the Tec elements (see also Jacobs et al. [21a]), that encodes a recombinase in addition to a DDE transposase. However, transposons that carry a gene for both a DDE transposase and a site-specific recombinase are common in bacteria. Such elements transpose replicatively, creating a temporary cointegrate structure during interreplicon transposition (3, 38; for a review, see reference 18). When both the donor and the target replicons are circles, as in bacteria, the cointegrate is a composite circle.

The generation of the bacterial cointegrate requires two steps. First, the DDE transposase nicks 3′ of each of the donor element strands, and then each 3′ OH end is covalently linked to the target in two strand transfer reactions. Second, the two displaced flanking target 3′ OH ends prime replication across each strand of the element, generating old and new heteroduplexes. The two old and new element copies lie at the junctions of the fused replicons, oriented in the same direction around the circle. The completion of transposition is accomplished by the resolution of the cointegrate circle back into the two original replicons. This resolution reaction is effected by a site-specific recombinase (usually a serine type; not homologous but analogous to the Tec tyrosine recombinase). The recombinase synapses the resolution site from each copy of the element and catalyzes recombination between them. This process unlinks the two replicons and leaves an element copy at the original location on the donor and an element integrated at the target site on the acceptor. The result is a new copy of the element at the target site location.

Do Tec elements transpose replicatively? If they do, they must have evolved solutions to problems arising from transposition within a genome of linear eukaryotic chromosomes and arising from the large number of Tec elements in the genome. During bacterial interreplicon transposition, the two replicons are temporarily joined in a single structure, the cointegrate. This circle serves to keep the two elements physically linked and in the proper orientation, so that they can participate in the subsequent resolution step. Transposition immunity would ensure that only two elements are present in this structure—the two that must participate to unlink the two replicons and restore them to their initial structures.

Neither condition pertains to Tec elements. First, like other eukaryotic genomes, the *E. crassus* germ line genome consists of multiple linear mitotic chromosomes. Second, there are hundreds of Tec elements within each chromosome. Replicative transposition by itself, without resolution, would cause chromosome rearrangements. Intrachromosomal transposition without resolution would cause one of two rearrangements: a deletion would result if the new element was inserted into the target in the same orientation as the donor element, or an inversion would result if the new insertion was in the opposite orientation. Similarly, without resolution, transposition between linear chromosomes would result in two possible structures, depending on the orientation of the two elements with respect to centromeres and proximal telomeres of their chromosomes. The two outcomes are a reciprocal (balanced) translocation, if the elements have the same relative orientations, or an acentric fragment and a dicentric fusion resulting from two elements with opposite orientations.

These cointegrate intermediate analogs would not present a problem if resolved before host DNA replication or mitosis. However, with thousands of transposon copies in the genome, the two cointegrate elements must be held together until resolution is complete: otherwise, they could not be correctly rejoined. These two element copies are generated side by side during the local DNA replication that completes cointegrate formation: if a mechanism to hold them together initiates at that point, then they could be held together for subsequent resolution. A precedent for such a replication-dependent mechanism is sister chromatid cohesion (32).

If the ORF 2 protein acts as a site-specific recombinase in transposition, a recognition or recombination site should reside on the element; resolution sites of serine-type recombinases in general consist of an 11- to 13-bp inverted repeat, split by 6 to 8 bp of a central sequence (for a review, see reference 18). We have not found a credible resolution site in the Tec1 or Tec2 sequences.

**Summary.** By examining patterns of sequence divergence, we have shown that the *E. crassus* Tec families of transposons have evolved under purifying selection for their protein products, and we have identified a functional programmed frameshift site in the Tec2 transposons. This site joins a number of other, similar identified shift sites in *Euplotes*. The *Euplotes* protein produced as a result of this frameshift is a member of the tyrosine site-specific recombinase family. This finding is surprising, given that tyrosine site-specific recombinases are extremely rare in eukaryotes and that no other known eukaryotic cut-and-paste transposon carries a tyrosine recombinase gene. The selection that we have characterized insists that the protein is needed for either element transposition or host fitness. We consider that Tec elements may transpose replicatively, the recombinase reverting the chromosomal rearrangements that replicative transposition creates.

**ACKNOWLEDGMENTS**

This work was supported by Public Health Service grants GM25203 to G.H. and GM37661 to C.L.J. from the National Institutes of Health.

We thank Mark Krikau, John Jaraczewski, and John Frels for determination of Tec gene sequences. We also thank Dorothy Shippen,
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