A Family of Developmentally Excised DNA Elements in Tetrahymena is under Selective Pressure to Maintain an Open Reading Frame Encoding an Integrase-Like Protein

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A family of developmentally excised DNA elements in *Tetrahymena* is under selective pressure to maintain an open reading frame encoding an integrase-like protein

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**ABSTRACT**

Tlr1 is a member of a family of ~20-30 DNA elements that undergo developmentally regulated excision during formation of the macronucleus in the ciliated protozoan *Tetrahymena*. Analysis of sequence internal to the right boundary of Tlr1 revealed the presence of a 2 kb open reading frame (ORF) encoding a deduced protein with similarity to retrotransposon integrases. The ORFs of five unique clones were sequenced. The ORFs have 98% sequence conservation and align without frameshifts, although one has an additional trinucleotide at codon 561. Nucleotide changes among the five clones are highly non-random with respect to the position in the codon and 93% of the nucleotide changes among the five clones encode identical or similar amino acids, suggesting that the ORF has evolved under selective pressure to preserve a functional protein. Nineteen T/C transitions in T/CAG and T/CAG codons suggest selection has occurred in the context of the *Tetrahymena* genome, where TAA and TAG encode Gin. Similarities between the ORF and those encoding retrotransposon integrases suggest that the Tlr family of elements may encode a polynucleotide transferase. Possible roles for the protein in transposition of the elements within the micronuclear genome and/or their developmentally regulated excision from the macronucleus are discussed.

**INTRODUCTION**

DNA is remarkably mobile. Two major categories of DNA rearrangements are those resulting from the invasion and proliferation of independent transposable elements and those which occur as part of a developmental program. The relationship between the two kinds of events is not yet understood.

Transposons are invasive elements that integrate into host genomes. Class I elements, or retrotransposons, transpose via an RNA intermediate. Class II elements transpose via a DNA intermediate. Both types of elements encode polynucleotidyl-transferases, which catalyze transposition, and an integrase or a transposase in the class I and class II elements, respectively (1). The active sites of these proteins are structurally similar in that they contain regularly spaced acidic amino acids, called DDE motifs, which chelate metal ions essential for enzymatic activity.

In a variety of phylogenetically diverse organisms an integral part of cellular development involves genomic reorganization associated with developmentally regulated DNA deletion. One example is the rearrangement of immunoglobulin genes and T cell receptors of vertebrate immune systems. Recent evidence has shown that the catalytic activity of RAG1, a recombinase that mediates the rearrangement of these genes, is dependent on three acidic residues, two of which may be involved in binding metal ions (2,3). This raises the possibility that there may be mechanistic similarities between the developmentally regulated DNA rearrangement and transposition.

Ciliated protozoa are particularly good model systems for the analysis of DNA rearrangement because thousands of programmed DNA deletions occur during development of the somatic genome. Ciliates are single cell eukaryotes that contain two structurally and functionally distinct nuclei that share the same genetic origin. The transcriptionally active somatic macronucleus sustains the cell during vegetative growth while the germline micronucleus remains transcriptionally silent. When ciliates reproduce sexually, the existing macronucleus is degraded and a new macronucleus develops from a mitotic copy of the fertilization micronucleus. Macronuclear development is associated with removal of thousands of internal eliminated sequences (IESs) (4,5).

The developmentally regulated deletion of IESs has been studied extensively in two classes of ciliates. In *Oxytricha* and *Euplotes* (formerly hypotrichs, these ciliates are currently placed in the class Spirotrichia; 6) ~93% of the DNA is eliminated during the transition from the micronucleus to the macronuclear genome (5,7). Specific elements are removed precisely and with 100% efficiency.

A substantial fraction of the eliminated DNA in the spirotrichs belongs to families of thousands of repetitive transposon-like elements. Examples of these include the Tec elements (transposon-like, *Euplotes crassus*) in *Euplotes* and the TBEs
(telomere-bearing elements) of Oxytricha. Tec elements are 5.3 kb in length and have ~700 bp of terminal inverted repeat sequence (8). TBE elements are 4.1 kb in length and have 78 bp terminal inverted repeats which include 17 bp of telomere-like G4T4 repeats (9–11). The Tec and TBE IESs are thought to be class II transposable elements. They contain terminal inverted repeats; there are short direct repeats that resemble target site duplications at the ends of the elements (5,10,12); and the presence of TBEI is allele-specific, which is consistent with a recent transposition event (13). Furthermore, both types of elements have open reading frames encoding putative transposases that are under selective pressure to maintain a functional protein (11,14,15).

In addition to the transposon-like IESs, the micronuclear genomes of the spirotrichs contain thousands of short, 10–539 bp, non-coding IESs that are AT-rich. These IESs are unique or of low copy number. One structural feature the short IESs of *Euplotes* share with Tec elements is the presence of a terminal TA direct repeat (5). Deletion of the two types of elements is similar in that both are excised as circular molecules with heteroduplex junctions (12) and their removal is coordinated with DNA replication during polytenization of the DNA in the micronuclear anlagen (16). Thus it seems likely that the two types of elements are deleted by similar mechanisms.

The process of developmentally regulated DNA deletion differs between spirotrichs and the evolutionarily distant Oligohymenophora class of ciliates. The most extensively studied member of the Oligohymenophora is *Tetrahymena thermophila*, wherein approximately 6000 elements, constituting 15% of the genome, are deleted during macronuclear development. As in the spirotrichs, IES deletion in *Tetrahymena* is 100% efficient. However, in *Tetrahymena* excision is imprecise. Several elements have alternative excision boundaries ranging over a few hundred base pairs (17–20) and most show sequence microheterogeneity at the rearrangement junctions (20–22). Another striking difference is in the structure of the excision products. In *Tetrahymena* the short IESs are more likely to be excised as linear molecules, in contrast to the circular heteroduplex excision products found in *Euplotes crassus* (12,23).

Most of the IESs that have been analyzed to date in *Tetrahymena* are relatively short, ranging from 0.6 to 2.9 kb, and consist of presumably non-coding AT-rich sequence (24). The elements have no common structural features. Some have short terminal direct repeats, but these are not required for excision of the element (25). The largest deletion element characterized to date is Tlr1, for *Tetrahymena* long repeat 1, which deletes >25 kb of sequence. Tlr1 belongs to a family of ~20–30 micro-nuclear-specific elements (19). The most striking structural characteristic of Tlr1 is an 825 bp terminal inverted repeat near the element termini. In this respect, Tlr1 structurally resembles the transposon-like elements found in the spirotrichs.

In order to determine whether the Tlr elements contain an open reading frame encoding a transposase, genomic clones of several family members were isolated and sequenced. This study describes the identification and sequence analysis of a 2 kb open reading frame within the Tlr family of elements. This open reading frame is highly conserved and encodes a putative protein with similarity to the integrases encoded by retroviruses and retrotransposons.

**MATERIALS AND METHODS**

**Accession numbers**

The Tlr Int clone sequences have been deposited in the GenBank database. Accession numbers are: Tlr1 Int, AF232246; Tlr Int A, AF232242; Tlr Int B, AF232243; Tlr Int C, AF232244; Tlr Int D, AF232245; Tlr Int E, AF232247.

**Strains**

*Tetrahymena thermophila* strain CU428 Mpr/Mpr (6-methylpurine-sensitive, VI) was obtained from Peter Bruns (Cornell University, Ithaca, NY). Strain CU399 was used as a source for the pMBR micronuclear plasmid library.

**Micronuclear and macronuclear DNA isolation**

Strain CU428 DNA was isolated according to the method described by Gorovsky et al. (26). Following separation by differential centrifugation, the nuclei were treated with 500 µg proteinase K in 20 mM Tris–HCl pH 7.5, 40 mM EDTA and 1% SDS at 37°C for 1 h. DNA was extracted with phenol/chloroform (1:1) and ethanol precipitated.

**Plasmid DNA isolation**

Plasmid DNA was isolated from bacterial cell transformants using either the Wizard Plus Maxiprep DNA purification protocol (Promega) or the Qiagen plasmid purification protocol (Qiagen).

**Oligonucleotides and primers**

Oligonucleotides for circularization of HindIII–EcoRI fragments of micronuclear DNA were 5’-AGCTTAGCTTCTGACGATCG-3’ and 5’-AATTGATGCTCTAGACTGAGCTA-3’. The circularized DNA was amplified by inverse PCR with primers 1 (5’-TCTATTACATCATTTTCTTA-3’) and 2 (5’-TTAATTTATGTAAGTGAAGCTT-3’). Primer 3, (5’-TCTAGTATTTATATCTTTTCTAG-3’), derived from the sequence of the inverse PCR product, was used against primer 2 to amplify Tlr family sequences from micronuclear DNA. Tlr1 sequences were amplified with the Tlr1 specific primer 4 (5’-AATGGAATTTGCAGTGGT-3’) and internal primer 5 (5’-GATGATCTAATTTATAGTTTCTC-3’).

**Southern hybridization**

Purified micronuclear DNA isolated from strain CU428 was digested with the appropriate restriction endonucleases, fractionated through 0.8–1% agarose gels and transferred via capillary action onto NEF 976 Genescreen Plus transfer membrane (NEN Research Products). DNA hybridization probes were radioactively labeled by the random primer protocol (Boehringer Mannheim). The nylon membranes were washed in 2x SSC (1x SSC is 0.15 M NaCl, 0.015 M sodium citrate) at room temperature, 2x SSC, 1% SDS at 65°C and 0.1x SSC at 58°C.

**Genomic library screening**

The pMBR plasmid library was constructed from strain CU399 micronuclear DNA partially digested with *MboI* and cloned into the plasmid vector pUC19, as described previously (27). *Escherichia coli* SURE (Promega) cells were transformed by electroporation with 100 ng of the library. Cells were spread onto Luria–Bertani broth (LB) plates containing 50 µg/ml.
carbenicillin. Colonies were screened by colony hybridization according to the Southern hybridization protocol.

**DNA sequencing**

DNA was sequenced on an Applied Biosystems sequencer using the ABI Prism™ dye terminator cycle sequencing ready reaction kit with AmpliTaq DNA polymerase at the University of Wisconsin, Milwaukee, automated sequencing center.

**Sequence alignment**

Sequences were aligned with Clustal 1.7 found under the multiple sequence alignment program using the Baylor College of Medicine search launcher (www.hgsc.bcm.tmc.edu/search-launcher/) sequence utilities.

**RESULTS**

Cloning of the Tlr family of deleted elements

The Tlr1 element, with long inverted repeats near the termini, is structurally similar to class II transposable elements, which undergo 'cut and paste' transposition, often mediated by transposases. The family of sequences is micronuclear-limited. The elements are unusual in that the copy number differs for different regions of the inverted repeat. Southern blot analysis showed that the innermost half of the Tlr1 inverted repeat hybridized to a family of 20–30 elements. This defines the Tlr family. The outermost region of the Tlr1 inverted repeat has sequence homology to a subfamily of sequences with a copy number of only 6–7 (19).

In order to determine whether the Tlr elements contained open reading frames that might encode a transposase, sequences internal to the inverted repeat at the right end of Tlr1 were cloned by inverse PCR (Fig. 1). A HindIII site is located 45 bp outside the right boundary of Tlr1. Genomic restriction mapping placed an EcoRI site -2.5 kb inside the element. Micronuclear DNA was digested with HindIII and EcoRI and circularized by ligation of the DNA in the presence of two complementary oligonucleotides which hybridize to each other to form a linker with HindIII and EcoRI sticky ends (Materials and Methods). DNA sequences internal to the Tlr1 inverted repeat were amplified by PCR with primers 1 and 2. The resulting inverse PCR product was partially sequenced and primer 3 was synthesized from this sequence. Micronuclear DNA was PCR amplified using primers 2 and 3. Since both primers are within sequences that are conserved in the Tlr family, this reaction contained a mixture of PCR products corresponding to various members of the family. A 1.2 kb cloned fragment contained the expected sequences with homology to the Tlr1 inverted repeat and 1079 bp of additional sequence internal to the repeat.

The innermost 369 bp of the 1.2 kb PCR product contained an open reading frame with similarity to Caenorhabditis elegans retrotransposon CER1 integrase. This open reading frame contains two Asp residues that are spaced proportionally to the Asp residues in the Dn/SpDnPE signature motif that is present in all integrases. In order to obtain the complete open reading frame, genomic clones were isolated from a plasmid library of micronuclear DNA (Materials and Methods). The library was screened with the 264 bp XbaI–HindIII and 971 bp XbaI–XbaI fragments internal to the Tlr1 inverted repeat (Fig. 1). Four clones with large inserts were selected for sequencing. All of these clones contained a 2 kb open reading frame encoding deduced proteins with homology to retroelement integrases. In Figure 2 the open reading frames of clones Tlr Int A, Tlr Int B, Tlr Int C and Tlr Int D are aligned. The sizes of the micronuclear DNA fragments demonstrate that they were independent clones and slight differences in the nucleotide sequence between clones indicated that they were four different members of the Tlr family of elements.

Tlr Int A–D are members of a family of elements that is repeated in the micronuclear genome. In order to obtain the integrase-like ORF associated with Tlr1, the corresponding fragment of micronuclear DNA was PCR amplified with Tlr1 specific primer 4, located in unique micronucleus-destined DNA to the right of Tlr1, and primer 5, in conserved DNA 3' to the open reading frame of the genomic clones (Fig. 2). The sequence at the right end of the 4.4 kb PCR product matched the known sequence of Tlr1 and the clone was designated Tlr1 Int. The integrase-like open reading frame begins 1583 nt from

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**Figure 1.** Restriction maps of the Tlr1 regions of micronuclear and macronuclear DNA. The cross-hatched line represents micronuclear-destined sequence. The thick solid line represents the micronuclear-limited sequence. The bold arrows represent the 825 bp Tlr1 inverted repeat. The symbol * marks the location of 102 bp tandem repeats. The angled arrow depicts the location of the 2 kb open reading frame identified in this study. Arrowheads represent PCR primers. Restriction sites: B, BglII; C, CiaI; E, EcoRI; H, HindIII; S, Sau3A; X, XbaI. The HindIII restriction site indicated by the dashed line is not present in Tlr1, but is present in other family members. The boxes represent fragments used in hybridization for isolation of Tlr Int clones.

**Figure 2.** Alignment of Tlr clones according to shared sequence. The vertical line in clone Tlr1 Int indicates the location of the right Tlr1 boundary and the arrow the inverted repeat. Mic, micronuclear-limited sequence; mac, macronucleus-destined sequence; angled arrow, the 2 kb open reading frame; triangles, the primers used to PCR amplify clone Tlr1 Int from micronuclear DNA.
Figure 3. Alignment of the first 227 amino acid residues in the Tlr Int consensus sequence with proteins or putative proteins that have similarity according to NCBI BLAST. Amino acids that are identical in at least five sequences are shaded black. Amino acids that are similar in at least five sequences are shaded gray. The active site DDE residues are marked with an asterisk. The integrase HHCC domain residues are indicated by circles. E values and accession numbers of the sequences are: C. elegans cosmid Y57GlI C. elegans, Z99281; X. maculatus HAS I putative protein, U43331; Trichoplusia ni TED retrotransposon, X95908; D. melanogaster mdg3 retrotransposon, X95908; S. cerevisiae TYB protein, Z47746.

Similarity of the conceptual protein to retroelement integrases

The Tlr Int clone and Tlr Int A–C each contained an open reading frame of 1998 nt encoding a conceptual protein of 666 amino acids. The open reading frame of Tlr Int D encoded one additional amino acid, due to insertion of a trinucleotide encoding Glu at residue 561. To find similarity to previously identified proteins, the DNA sequence of the 2 kb open reading frame was submitted to NCBI BLASTX (Basic Local Alignment Search Tool) (28). The closest matches to the Tlr family open reading frame were sequences found in transposons or putative transposons. An alignment of the deduced proteins from a variety of organisms is shown in Figure 3. The best characterized of these was the TYB protein of Saccharomyces cerevisiae, which encodes a retrotransposon integrase. All integrases and many transposases contain three acidic amino acids with characteristic spacing (D_{36-38}D_{15}E) which comprise the catalytic core of the proteins. Each of these residues is independently essential for catalytic activity involved in the reactions of transposition (29–35). The Tetrahymena open reading frame encodes these acidic residues, along with characteristic blocks of conserved amino acid residues surrounding each of the signature amino acids.

Integrase and transposases can be distinguished on the basis of the conserved amino acids surrounding the DDE signature (36; Fig. 4). The first Asp of the signature is generally followed by an acidic residue in transposases and more often by a hydrophobic amino acid in integrases. The second Asp is followed by an Asn in both classes of enzymes. Most notably, the integrase Glu is in a consensus sequence ERMNRIKTILK. In the transposases the consensus sequence in the region of the Glu is SPDLNPILH/L. The alignment in Figure 4 suggests that the deduced protein encoded by the Tlr family of elements in Tetrahymena is more closely related to the integrases than the right Tlr1 boundary and 710 nt internal to the inverted repeat (Fig. 1).
open reading frames, and likelihood of interchanges. A
assigned a number from 0 to 6 based on the structural identity
and likelihood of interchanges. A score of 4 or 5, according to
the SG Matrix scoring system (38), where amino acids are
scored on the basis of their structural identity and likelihood of interchanges. A score of 4 or 5 indicates identity.
For the analysis in Table 1, similarity was defined as an
SG score of 4 or 5. Of the 196 nucleotide changes among the five open reading frames, 140 encoded identical amino acids,
42 encoded similar amino acids and 14 encoded dissimilar
amino acids (Table 1). Therefore, 93% of the nucleotide
changes altered codons to encode either identical or similar
amino acids and the amino acid sequence of the putative proteins encoded by the five elements is 99% conserved.

Table 1. Nucleotide and amino acid substitutions among the Int open reading frames

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Nucleotide position</th>
<th>Total nucleotides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Identical</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>Similar</td>
<td>13 ^a</td>
<td>16</td>
</tr>
<tr>
<td>Dissimilar</td>
<td>4 ^c</td>
<td>9 ^d</td>
</tr>
<tr>
<td>Total</td>
<td>37</td>
<td>25</td>
</tr>
</tbody>
</table>

^cFour codons with two nucleotides different from the consensus sequence.

Conservation of the amino acid sequence despite nucleotide
changes suggested that the open reading frame has evolved
under selective pressure to conserve a functional protein.
Statistical analysis of the synonymous versus non-synonymous
nucleotide changes was done in order to determine the signifi-
cance of the amino acid conservation (39). According to this
method, nucleotide substitutions that encode similar or identical
amino acids are synonymous (p S) whereas nucleotide
substitutions that encode dissimilar amino acids are non-
synonymous (p N). If there is selective pressure to conserve a
protein coding region, p S is expected to be greater than p N. In
a one-tailed 1-test the p S value was significantly greater than p N
for each pair of open reading frames at P < 0.001.

One unusual feature of the Tetrahymena genetic code is that
the canonical stop codons TAA and TAG encode Gln (40,41).
Thus T→C transitions in the first nucleotide of T/CAA or T/CAG
Gln codons are silent. Such transitions accounted for 19 of the
20 nucleotide changes in the first nucleotide of a codon that
resulted in identical amino acids (Table 1), suggesting that the
open reading frame has evolved in the context of the Tetra-
hymena genome.

The N-terminus of the Tetrahymena integrase-like protein
had the highest degree of similarity to integrases of other
organisms. In order to determine whether selection among
the Tlr family members is uniform over the length of the putative
protein, the open reading frame was arbitrarily divided
into thirds, with the DDE motifs of the active site located in
the N-terminal region of the protein. Interestingly, 1 /2 analysis
indicated that although the nucleotide substitutions occur
randomly across the gene, dissimilar amino acid changes were
preferentially located in the C-terminal region of the open
reading frame (P < 0.01). Thus, although the C-terminal region
of the protein is under strong selection, it is not as stringent as
the selection in the N-terminal two-thirds of the protein.

Copy number and genome specificity of the Tlr family
of sequences

Whereas the innermost portion of the Tlr1 inverted repeat is
repeated 20- to 30-fold in the micronuclear genome, the outer-
most sequences, as indicated by hybridization of the tandemly
This was consistent with the sequence data suggesting that Xbal sequences within the element are highly conserved among the TlrI inverted repeat. Between the TlrI inverted repeat and the integrase open reading frame (19). Thus it was of interest to determine the copy number of sequences fragment have a copy number similar to that of the inner part of family members. There was no hybridization to macronuclear and macronuclear (MA) DNA digested with EcoRV, thus the family of elements is efficiently eliminated during macronuclear development. The presence of macronuclear DNA was confirmed by ethidium bromide staining of the gel prior to transfer (data not shown).

A similar copy number was found for sequences in the open reading frame. Sequence analysis of the five clones at hand revealed a conserved EcoRV site within the open reading frame and no additional EcoRV sites in the conserved sequence 5' to the open reading frame. Therefore, Southern analysis of DNA digested with EcoRV and probed with the Clal-EcoRV fragment located entirely within the open reading frame provided an estimate of the number of elements that contain the open reading frame (Fig. 5C). The observed pattern of 20–30 bands was similar in number to that found for sequences 5' to the open reading frame and for the innermost part of the TlrI inverted repeat. As in the previous Southern blot, the probe hybridized only to micronuclear DNA, suggesting that the 2 kb open reading frame is located within a family of micronuclear-limited elements.

The similarity in copy number and the micronucleus-limited character suggested that the integrase-like open reading frame may be generally associated with sequences in the TlrI inverted repeat. The limited data available for individual clones supports this model. Of the clones that cover the integrase-like open reading frame, only Tlr Int A extends far enough 5' of the gene to contain DNA with homology to the TlrI inverted repeat. Another clone, Tlr Int E, which contained only the 5'-end of the integrase-like open reading frame, extended far enough to include sequences with homology to the TlrI inverted repeat. Sequence data from these three clones suggests that the integrase-like open reading frame is generally associated with the Tlr family, among which the innermost ~275 bp of the TlrI inverted repeat are conserved.

The Southern blots also support the sequence data indicating a high degree of conservation among the integrase-like genes. Three major bands were seen in DNA digested with Clal and EcoRI (Fig. 5C). One of these had the mobility expected for the 937 bp Clal-EcoRV fragment that is present in all of the sequenced clones. The two other bands at 1.4 and 3.5 kb indicate that there are three major variations in the Clal + EcoRV restriction pattern among the various family members.

**DISCUSSION**

The Tlr family of micronucleus-limited elements in *Tetrahymena* encodes a putative protein with similarity to retrotransposon integrases. The open reading frames from five of the 20–30 family members have been cloned and sequenced. Nucleotide changes among the five family members are highly non-random, suggesting that the elements are under selective pressure to maintain a functional protein in most or all of the elements.

The role of the integrase-like protein in *Tetrahymena* biology is unknown. One possibility is that the Tlr family of IESs encodes the machinery for its own developmentally regulated excision and the integrase-like enzyme is a poly-nucleotidytransferase that is part of that machinery. A functional relationship between transposable elements and developmentally regulated DNA excision was first proposed to account for conservative selection of the open reading frames in the Tec and TBE elements of the spirotrichs (5). According to this model, IESs are degenerate transposons. Developmentally repeated 19mers, belongs to a smaller family of only six to seven elements (19). Thus it was of interest to determine the copy number of sequences internal to the inverted repeat and of the open reading frame encoding the putative integrase. Micronuclear and macronuclear DNA were digested with various restriction enzymes and hybridized in a Southern analysis with the 714 bp HindIII-XbaI fragment containing sequences between the TlrI inverted repeat and the integrase open reading frame (Fig. 5B). In the lanes containing micronuclear DNA digested with EcoRI, EcoRV and HindIII –20–30 fragments hybridized, suggesting that sequences homologous to this fragment have a copy number similar to that of the inner part of the TlrI inverted repeat. In micronuclear DNA digested with XbaI the majority of the hybridization was to a 1 kb fragment. This was consistent with the sequence data suggesting that sequences within the element are highly conserved among family members. There was no hybridization to macronuclear DNA, thus the family of elements is efficiently eliminated during macronuclear development.
regulated DNA deletion evolved as a mechanism to remove them from the transcriptionally active macronucleus, with the transposase serving as the excisase. In *Euplotes* and *Oxytricha*, where many of the IESs occur within coding sequences, efficient IES excision might be expected to exert a powerful selective pressure to conserve the active transposase (42). This model is somewhat less compelling in *Tetrahymena*, where no protein coding sequences are known to be interrupted by IESs. However, it is possible that there are other selective advantages to removing IESs from the macronuclear genome. For example, they might play a role in micronuclear functions such as chromatin condensation and/or transcriptional silencing that are dispensable in the macronucleus (43).

Although conservative selection for functional protein might be expected if the integrase-like open reading frame of the Tlr family encodes excisase, it is not clear what selective mechanism would maintain functional genes in multiple copies of the element, since such an enzyme would presumably act in trans. Constructs containing the inverted repeats of Tlr1, but lacking the integrase-like gene, undergo efficient and accurate rearrangement in vivo (44). Thus developmentally regulated DNA excision of the Tlr family does not require an active integrase-like gene in cis.

Multiple genes encoding the integrase-like protein might be required to synthesize sufficient amounts of the protein to catalyze excision of the family of elements within a specific and brief developmental time period. However, transcripts of the Tec transposase gene of *Euplotes* were detected only by extremely sensitive methods involving Southern hybridization of RT–PCR products and the transcript levels are thought to be insufficient to account for the en masse excision of ~10^6 Tec elements/polytene nucleus in a 2–4 h time period (45,46). Similarly, transcripts of the integrase-like genes in *Tetrahymena* have not been detected by standard northern blot analysis (J.A.Gershan, unpublished data).

A second possible function of the integrase-like gene is that the Tlr family of sequences are mobile elements and the integrase-like enzyme is responsible for their transposition in the micronuclear genome. Although the enzyme encoded by the Tlr family of elements has a DDE motif similar to that of retroelements, the Tlr elements differ from retrotransposons in several respects. At >25 kb (J.D.Wuitschick and J.A.Gershan, unpublished data), they are larger than most active retroelements and the terminal repeats are inverted rather than direct repeats. Analysis of sequences surrounding the Int open reading frame has not revealed a discernible gag–pol gene structure characteristic of retroelements and the putative integrase gene lacks the N-terminal HHCC zinc finger domain that is required for retroelement cDNA processing (30,37). (A complete analysis of the structure of these large elements is in progress and will be published elsewhere.) The long inverted repeats of Tlr1 are a structural characteristic common to many class II elements. Perhaps the Tlr elements are class II elements in which the motifs surrounding the active site DDE residues of the transposases are more similar to those in the integrases than to those of the majority of transposases. This would be unusual, but not unprecedented (36). The bacterial IS30 element is an example of a class II element with integrase-like active site motifs. The putative enzyme encoded by the Tlr family is like the transposase of the bacterial IS30 elements in that both lack the N-terminal HHCC integrase domain (47).

If the Tlr family of sequences are transposons, then it is necessary to account for the fact that these elements are found exclusively in the 15% of the micronuclear genome that undergoes developmentally regulated elimination. Excision of the Tlr elements during macronuclear development could be a secondary consequence of transposon targeting into micronuclear IESs. Two observations support this hypothesis. First, Tlr1 is removed from the macronuclear genome at the same stage of macronuclear development as the smaller, AT-rich IESs (17; Capowski and K.M.Karrer, unpublished data). Second, flanking sequences have been shown to play a role in the delineation of the rearrangement boundaries of Tlr1 (44). This is not an expected feature of transposon excision, but is consistent with the hypothesis that Tlr1 resides within an IES because, in *Tetrahymena*, cis-acting sequences in the flanking DNA regulate the deletion of IESs (21,22,48).

Unique features of chromatin structure might serve to target transposition of Tlr elements to IESs. There is a precedent for transposon targeting to regions of distinct chromatin structure in yeast, where Ty5 elements are selectively inserted into regions of silent chromatin by association of the integration complexes with localized host factors (49–52). Differences between the chromatin structure of eliminated sequences and macronucleus-destined sequences have been detected in *Euplotes* by analysis of chromatin digested with micrococcal nuclease (53). In *Tetrahymena* the chromatin of IES-containing regions of the genome is distinct from that of the bulk of the genome during macronuclear development. The abundant stage-specific proteins Pdd1p and Pdd2p, both of which are required for developmentally programmed DNA deletion in *Tetrahymena* (54,55), are preferentially associated with euchromatin and heterochromatin regions containing IESs (56,57).

If the integrase-like protein encoded by the Tlr family of elements is a transposase and is not functioning as the excisase, then the need to maintain sufficient enzyme for IES excision cannot be invoked to explain the apparent selection on these genes. The transposases of class II elements can be subject to selective pressure for functions other than the transposase activity, such as repression of transposition (58).

Five of the estimated 20–30 integrase-like open reading frames, a significant fraction of the total Tlr family, were analyzed in this study. Despite numerous nucleotide changes amongst the genes, all of the family members examined maintained the open reading frame and a high degree of protein similarity. Whatever the biological role of the integrase-like protein, the data indicate that there is strong selective pressure to maintain an active gene in most or all of the Tlr family members.

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