An *Entamoeba histolytica* LINE/SINE Pair Inserts at Common Target Sites Cleaved by the Restriction Enzyme-Like LINE-Encoded Endonuclease

Prabhat K. Mandal,1 Anindya Bagchi,2† Alok Bhattacharya,1 and Sudha Bhattacharya2*

School of Environmental Sciences2 and School of Life Sciences,1 Jawaharlal Nehru University, New Delhi 110067, India

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The advent of large-scale genome sequencing will make it possible to understand the origin and evolution of transposable elements and their dynamic relationship with the resident genome. In this respect the study of ancient lineages of eukaryotes, for example, the amitochondrial human pathogens *Giardia lamblia* (1, 7) and *Entamoeba histolytica* (26), have recently been shown to harbor non-long-terminal-repeat (non-LTR) retrotransposons. These retrotransposons (also called long interspersed repetitive elements [LINEs]) are considered more primitive than the LTR retrotransposons (13, 35). In keeping with this contention, it is interesting that non-LTR elements are found abundantly in parasitic protozoa while there is only one report so far of an LTR element in these organisms (3).

Data from whole-genome sequencing efforts have revealed the presence of multiple LINE families in *E. histolytica* (30). Of these, the most abundant seems to be the 4.8-kb *E. histolytica* retrotransposon-like element (EhRLE), a LINE family from *E. histolytica* that had been described previously (26). In addition, short repetitive sequences of about 0.5 kb that are abundantly transcribed have also been identified (10, 32, 33). These, the most abundant seems to be the 4.8-kb *E. histolytica* genome, known to be occupied by EhSINE, were analyzed. Both elements resulted in short target site duplications (TSD) upon insertion. A common feature was the presence of a short T-rich stretch just upstream of the TSD in most insertion sites. By sequence analysis an empty target site in the *E. histolytica* genome, known to be occupied by EhSINE, was identified. When a 176-bp fragment containing the empty site was used as a substrate for EN, it was prominently nicked on the bottom strand at the precise point of insertion of EhSINE1, showing that this SINE could use the LINE-encoded endonuclease for its insertion. The nick on the bottom strand was toward the right of the TSD, which is uncommon. The lack of strict target site-specificity of the restriction enzyme-like EN encoded by EhLINE1 is also exceptional. A model for retrotransposition of EhLINE1/SINE1 is presented.

The non-long-terminal-repeat (non-LTR) retrotransposons (also called long interspersed repetitive elements [LINEs]) are among the oldest retroelements. Here we describe the properties of such an element from a primitive protozoan parasite, *Entamoeba histolytica*, that infects the human gut. This 4.8-kb element, called EhLINE1, is present in about 140 copies dispersed throughout the genome. The element belongs to the R4 clade of non-LTR elements. It has a centrally located reverse transcriptase domain and a restriction enzyme-like endonuclease (EN) domain at the carboxy terminus. We have cloned and expressed a 794-bp fragment containing the EN domain in *Escherichia coli*. The purified protein could nick supercoiled pBluescript DNA to yield open circular and linear DNAs. The conserved PDX12–14D motif was required for activity. Genomic sequences flanking the sites of insertion of EhLINE1 and the putative partner short interspersed repetitive element (SINE), EhSINE1, were analyzed. Both elements resulted in short target site duplications (TSD) upon insertion. A common feature was the presence of a short T-rich stretch just upstream of the TSD in most insertion sites. By sequence analysis an empty target site in the *E. histolytica* genome, known to be occupied by EhSINE1, was identified. When a 176-bp fragment containing the empty site was used as a substrate for EN, it was prominently nicked on the bottom strand at the precise point of insertion of EhSINE1, showing that this SINE could use the LINE-encoded endonuclease for its insertion. The nick on the bottom strand was toward the right of the TSD, which is uncommon. The lack of strict target site-specificity of the restriction enzyme-like EN encoded by EhLINE1 is also exceptional. A model for retrotransposition of EhLINE1/SINE1 is presented.
recently been demonstrated for the eel LINE/SINE pair (19). Although sequence conservation of the 3' tail of the E. histolytica LINE/SINE pair is suggestive of SINE mobilization by LINE-encoded enzymes, there is no experimental evidence yet in favor of this hypothesis. Nothing is known about the mechanism of retrotransposition in this primitive parasite. The general model that has served to explain LINE retrotransposition is target-primed reverse transcription, proposed by Eickbush and coworkers from their work on the silk moth element R2Bm (20). According to this model, the LINE-encoded endonuclease nicks the bottom strand of the target site, generating a 3'-OH group that primes reverse transcription of the element RNA.

In order to understand the mode of transmission of EhRLE in the E. histolytica genome, we have cloned and expressed the EN domain in Escherichia coli and studied its properties with respect to target site specificity of nicking in vitro. Here we show that the enzyme is not site specific and that it nicks a variety of sequences that fall into a loose consensus. This is the first clear example of an element belonging to the R2 group that does not insert in a site-specific manner. Some R6x6 elements of vertebrates also probably lack strict target site specificity, but this has yet to be clearly defined (31). We also show that the enzyme can nick an empty target site, known to be occupied by IE/Ehapt2 in vivo, at the precise point of insertion. This provides strong functional evidence that IE/Ehapt2 is a SINE element that utilizes the enzymatic machinery of EhRLE for its insertion. For the sake of uniformity in nomenclature, we henceforth refer to EhRLE as EhLINE1 and to IE/Ehapt2 as EhSINE1.

MATERIALS AND METHODS

Cloning of EhLINE1 EN. The clones from the shotgun library used for E. histolytica genome sequencing (accession numbers AZ669903 and AZ541056) were selected for cloning the EN domain. These were a kind gift from B. Loftus, The Institute for Genomic Research (TIGR). A 276-bp EcorI-Nov fragment from AZ669903 and a 600-bp Ndel-BamHI fragment from AZ541056 were ligated and cloned into pBS [pBlueScript II KS (+); Stratagene]. For expression of EhLINE1 EN in E. coli, the EcorI-Nov fragment from pBS was cloned into the EcoRI-Nov site of pET30b (Novagen) and transformed into E. coli BL21(DE3) to yield pET-EN. The PDIXD-D→PAN-X mutation was generated by a QuickChange site-directed mutagenesis kit (Stratagene) using primers 5'-CAAATATCACAATAACCTAACATAGTATATATAGA3' and 5'-TTT ATTATAATAGATTTTCTGTATTTTTATATGGC-3'. The sequence of the wild-type (pET-EN) and mutant (pET-ENM) constructs were confirmed by sequencing.

Expression and purification of EhLINE1 EN. E. coli cells (BL21; 200 ml) containing pET-EN were grown at 30°C in Luria-Bertani medium containing 30 µg of kanamycin/ml to an OD600 of 0.7. For induction of His6-tagged EN and mutagenized EN (ENM) proteins, isopropyl-β-D-thiogalactopyranoside (IPTG) was added to a final concentration of 1 mM and the cells were further incubated for 3 h. Recombinant proteins were purified by Ni²⁺-nitritotriacetic acid-agarose affinity chromatography as described by the supplier (Qiagen). The cells were harvested by centrifugation and then resuspended in 10 ml of buffer A (50 mM NaH₂PO₄ [pH 8.0], 300 mM NaCl, 20 mM imidazole [pH 8.0], 10 mM β-mercaptoethanol, 0.5% Triton X-100, and 2 mM phenylmethylsulfonyl fluoride). These were then lysed by three cycles of freeze-thawing, followed by sonication on ice. The lysate was centrifuged at 12,000 × g for 30 min at 4°C. The supernatant was passed through a 0.45-µm pore-size filter (Millipore) and incubated with 1 ml of preequilibrated Ni²⁺-nitritotriacetic acid-agarose (Qiagen) for 2 h at 4°C with gentle mixing. It was then packed in a C10/10 column (Amersham Pharmacia Biosciences) and washed several times with buffer A containing 50 mM imidazole. The recombinant protein was eluted with an imidazole gradient (150 to 500 mM). The EhLINE1 EN protein was found to elute at an imidazole concentration of 150 to 200 mM. Fractions containing EhLINE1 EN were identified by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis and then pooled and concentrated by dialyzing against buffer B (50 mM Tris-HCl [pH 7.5], 100 mM NaCl, 20 mM MgCl₂, 2 mM dithiothreitol [DTT], 20% sucrose). The sucrose was removed by dialyzing against buffer C (50 mM Tris-HCl [pH 7.5], 100 mM NaCl, 20 mM MgCl₂, 2 mM DTT, 10% glycerol). The purified protein was stored in aliquots of 2 to 5 µg in 30% glycerol at −20°C.

Immunoblotting. Proteins were separated on an SDS–10% polyacrylamide gel and transferred to Hybond ECL nitrocellulose paper (Amersham Pharmacia Biosciences). After transfer, the membrane was blocked with 5% bovine serum albumin in TBS-T (20 mM Tris-HCl, 150 mM NaCl, 0.1% Tween 20 [pH 7.6]) overnight and subsequently incubated with the anti-EN tag antibody (Amersham Pharmacia Biosciences) for 1 h at room temperature. The membrane was washed with TBS-T three times and incubated for 1 h at room temperature with a horseradish peroxidase-conjugated secondary antibody. Subsequently, the membrane was washed three times with TBS-T. The proteins were visualized with an enhanced chemiluminescence kit (Amersham Pharmacia Biosciences).

Nicking assay. Supercoiled plasmid DNA was purified by using a Qiagen tip100 plasmid purification kit. The EhLINE1 EN reaction mixture contained 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 20 mM MgCl₂, 1 mM DTT, 0.2 µg of supercoiled pBS DNA, and 80 ng of purified protein in a total reaction volume of 30 µl. The reaction was carried out at 37°C for 1 h and was stopped by addition of 25 mM EDTA. Products were separated on a 0.8% agarose gel containing 0.5 µg of ethidium bromide/ml.

Growth of E. histolytica and genomic DNA purification. E. histolytica strain HM1:IMSS was maintained in TYI-S-33 medium (11) at 36°C with appropriate antibiotics. Total DNA was purified from cells grown to late-log phase, as described previously (2).

Mapping of nicking sites. The primer (50 pmol) was end labeled in a 20-µl reaction mixture by using 50 µCi of [γ-³²P]ATP (Amersham Pharmacia Biotech) and T4 polynucleotide kinase (New England Biolabs). The reaction was stopped by incubation at 65°C for 20 min, and the labeled primer was purified by being passed through a Sephadex G-25 column (Amersham Pharmacia Biosciences). (25). The DNA substrates were generated by PCR with E. histolytica genomic DNA as the template and a combination of one end-labeled primer and one unlabeled primer. The sequences of the primers used are indicated in the figure legends. After PCR the products were separated on a 6% native polyacrylamide gel, and the DNA band corresponding to the full-length product was excised from the gel. The DNA was recovered by the “crush-and-soak” method (25).

The DNA substrate (100 ng) was incubated with 40 ng of protein in a 10-µl reaction mixture for 1 h at 37°C. The protein was inactivated as mentioned above. For denaturing electrophoresis on 6% polyacrylamide gels, a 2-µl aliquot of the reaction product was mixed with 8 µl of formamide gel loading dye (95% formamide, 20 mM EDTA, 0.05% bromophenol blue, and 0.05% xylene cyanol FF), boiled for 5 min, and chilled on ice before loading. The parallel sequencing reaction was carried out by using a Thermo Sequenase cycle sequencing kit (Amersham Pharmacia Biosciences). Template DNA (100 to 150 ng) and 1 to 2 pmol of primer were used for each sequencing reaction. Electrophoresis was carried out at 65 W for 1 to 3 h with the gel temperature maintained at 45 to 50°C. The gels were fixed, dried, and exposed to X-ray film.

Phylogenetic analysis. Sequences used for phylogenetic analysis were taken from the work of Burke et al. (7). The sequences of EhLINE1, EhLINE2, and EhSINE1 were assembled from the following entries in the Genome Sequence Survey (GSS) database: EhLINE1, accession no. AZ539252, AZ684839, AZ542852, AZ669999, AZ687050, AZ533295, AZ546488, AZ678960, AZ699093, and AZ341065; EhLINE2, accession no. AZ547248, BH152719, AZ687791, AZ534852, AZ546599, BH135183, AZ679659, AZ688833, AZ545802, AZ534795, AZ542853, AZ545537, and BH146735; EhSINE1, accession no. AZ532904, BH135862, AZ687777 (removed stop codon), AZ544710, AZ542964, AZ542119, AZ679616, AZ530593, AZ530263, AZ550894 (corrected frameshift), and AZ549664. The RT and EN domains were aligned by using the Clustal X alignment program (29). The rooted phylogenetic tree was derived by the neighbor-joining method using the PAM250 matrix of PHYLIP (14) and maximum-parsimony heuristic options in PAUP (27). Bootstrapping was carried out by using PAUP, with 100 data sets.

RESULTS

Cloning and purification of the EN polypeptide. From nucleotide sequence analysis of EhLINE1, the conserved motifs of the EN domain were localized to nucleotide positions 4122 to 4479 (Fig. 1A). Two GSS entries encompassing this region
and lacking any stop codons were used to clone the EN domain. The clones corresponding to these entries (accession no. AZ669903 and AZ541065) were a kind gift from TIGR. The 794-bp EcoRI-BamHI fragment (nucleotide positions 4010 to 4804) containing the EN domain was cloned into the E. coli expression vector pET30b. The expressed protein contained a His tag, and together with other vector sequences at the amino terminus, it was 309 amino acids long, with an expected molecular mass of 35.5 kDa. It was purified by nickel-agarose chromatography, and its identity was confirmed by using an anti-His tag antibody. As a control, the PDX 14D motif in the EN domain, thought to be essential for enzyme activity, was mutated (37). The ENM motif was changed to PAX 14D and expressed in E. coli. The recombinant EN protein purified from E. coli was stable for at least 6 months in 30% glycerol at −20°C.

Lack of a strict target-site consensus sequence for insertion of EhLINE1 and EhSINE1. To design a suitable substrate for testing the endonucleolytic activity of the EN protein, we looked at the flanking genomic sequences at the sites of insertion of EhLINE1 and its putative partner SINE, EhSINE1. Sequences were obtained from the GSS and contig databases and were aligned using ClustalW. The 5′ and 3′ ends of the elements were defined as follows. Both elements started with 5′-AGATC, after which the sequence of the EhLINE1 family diverged from that of the EhSINE1 family, but within each family the sequence was highly conserved. The 3′ ends of the two families shared a 74-bp sequence, which ended with CTT TTTATTT-3′, with minor variations. The sequences upstream and downstream of EhLINE1 and EhSINE1 showed that these elements do not insert in a strictly site-specific manner. However, a T-rich stretch of 15 to 20 nucleotides was almost uni-
versally present about 13 to 19 nucleotides upstream of the insertion site of both elements.

The EN protein can cleave a nonspecific substrate, pBS. The lack of a strict target site consensus for insertion of EhLINE1 prompted us to check if the EN protein could utilize a non-specific substrate, pBS, as found for the human L1 endonuclease (15). Supercoiled pBS DNA was efficiently nicked by the purified EN protein to yield open circular and linear DNAs.
by Southern hybridization. The appearance of discrete bands end-labeled linear DNAs were used as substrates. The nicks also borne out by the experiments described below in which absence of exonuclease activity in our enzyme preparation is a low level of exonucleolytic activity cannot be ruled out. The due to closely spaced nicks on the two DNA strands, although DNA even after 240 min, indicates that smears are probably presence of detectable levels of high-molecular-weight linear DNAs as substrates. The slow appearance of smears, and the below) shows that the enzyme can use both circular and linear molecular-weight molecules. This (and experiments reported On further incubation, the linear DNA gave a smear of low-molecular-weight molecules. This (and experiments reported first that of open circular DNA, the enzyme appears to make pre- produced at early time points was very small compared with 5 min, following which the enzyme was denatured and DNA was linearized with EcoRI. The fragments generated were detected by Southern hybridization. The appearance of discrete bands along with a background smear showed that the enzyme did have a preference for some sites (data not shown).

Nicking activity of EN at the target sites occupied by EhLINE1 and EhSINE1. To assay the activity of EN on its natural substrate, we looked for sites of insertion of EhLINE1/SINE1 in the E. histolytica genome, which may be unoccupied. Since E. histolytica is thought to be polyploid (n = 4) (9), it may be possible to find homologous regions of chromosomes, some of which harbor these elements while others are unoccupied. To look for unoccupied sites, it is necessary to have contigs containing the full-length element, from which flanking sequences can be obtained. Since the assembly of the E. histolytica genome sequences is still in progress, we undertook this exercise only for the 0.55-kb EhSINE1 at this point. The GSS database was adequate for this purpose, since it contains, on average, 0.8 kb of sequence available from either end of a clone. Thirty-eight entries containing the full-length EhSINE1 could be retrieved from the database. Sequences flanking the ends of the element in each entry were stitched together and used to search the database for unoccupied sites. Only one unoccupied site was found (AZ669709). The corresponding EhSINE1-containing sequence was AZ550231. The infrequent occurrence of unoccupied sites may be a limitation of the database, or it may be a consequence of the transposition mechanism, which does not leave unoccupied sites even in a polyploid genome. Both EhSINE1 and EhLINE1 insertions are accompanied by short target site duplications (TSDs), as seen from sequence analysis. In the case of AZ550231, EhSINE1 insertion resulted in a 22-bp TSD, which is present only once in AZ669709.

| Band  | Unoccupied site | EhSINE1 insertion | EhLINE1 insertion |
|-------|----------------|-------------------|-------------------|
|       | Bottom strand | Top strand | 3' end | 5' end | 3' end | 5' end |
| 1     | 5'-GTATG     | 5'-TTACTT | 5'-GTATG | 5'-GGAGA | 5'-GATAAT | 5'-GTATTT |
| 2     | 5'-GCATT     | 5'-TTAATT | 5'-GCATT | 5'-GAGTC | 5'-GATAA | 5'-GCAATT |
| 3     | 5'-GTATTT    | 5'-TTATT  | 5'-GTATT | 5'-GTGTTT | 5'-GCAATT |
| 4     | 5'-TAATA     | 5'-TTAAT  | 5'-GTATT | 5'-GCCACC | 5'-GTATT |
| 5     | 5'-GAGTA     | 5'-CCTCT  |         |         |         |
| 6     | 5'-TTATT     |         |         |         |         |
| 7     | 5'-GTATT     |         |         |         |         |
| 8     | 5'-CAATG     |         |         |         |         |
| 9     | 5'-CAATA     |         |         |         |         |
| 10    | 5'-GTCTTT    |         |         |         |         |
| 11    | 5'-TACCTT    |         |         |         |         |
| 12    | 5'-ACATA     |         |         |         |         |
| 13    | 5'-CACAA     |         |         |         |         |

* Data summarized from Fig. 2 and 3.
A 176-bp fragment containing the unoccupied site was obtained by PCR amplification of *E. histolytica* genomic DNA using primers designed from the sequence in AZ669709 (Fig. 2). The fragment was sequenced to confirm its identity with the sequence of the GSS entry. The fragment was then labeled either in the bottom strand or in the top strand by PCR amplification using the corresponding end-labeled primer. It was incubated with the endonuclease, and the resulting products were denatured and separated on a sequencing gel along with a sequencing reaction of the same template with each primer. The sites of nicking by the endonuclease on each strand were identified from the parallel sequencing run. The substrate in which the bottom strand was labeled showed three major nicking sites in which the products were doublets (Fig. 2, double arrows 1 to 3) and two less-preferred sites in which the products were primarily single bands (Fig. 2, arrows 4 and 5). Of these, bands 1 and 3 were the brightest. Band 3 corresponded with the exact site of insertion of EhSINE1. The pattern of nicks on the top strand was different from that on the bottom strand. The enzyme acted on many sites, but none was as strongly preferred as the sites in bands 1 and 3 on the bottom strand. In addition, most of the bands generated from the top strand appeared to be singlets (Fig. 2). The sequences surrounding the nicks on the two strands are listed in Table 1. From this it appears that the enzyme nicks at some preferred sequences and not randomly, since clear patterns could be discerned. The differences between the nicking patterns of the top and bottom strands could, in part, be due to the fact that the top strand was T rich while the bottom strand was A rich. The property of this endonuclease to nick at one of two consecutive phosphodiester bonds has also been observed with human L1 endonuclease (15) and R2Bm endonuclease (34). We were not able to
repeat this experiment with more unoccupied sites of EhSINE1, since only one such site could be identified in the database used for this study.

To further understand the nicking preferences of the endonuclease, we used as substrates fragments containing the boundary regions of sites where EhSINE1 and EhLINE1 had inserted. For EhSINE1 the GSS entry AZ550231 contained the entire element. Primers were designed to PCR amplify two fragments (189 and 163 bp, respectively), one from the 5' and one from the 3' end, containing a part of the element and adjoining genomic sequence (Fig. 3A). Eh. histolytica genomic DNA was used for PCR amplification. To obtain radioactively labeled fragments, the bottom strand primer was end labeled in each case. After incubation with the endonuclease, the resulting products were separated on a sequencing gel as described for Fig. 2. Several bands of various intensities were observed with both fragments. Of these, the four most prominent bands are marked (Fig. 3A). In the fragment from the 3' end, there was a prominent nick at the 3' boundary with the TSD (band 3), which corroborates the data for the unoccupied site in Fig. 2. The sequences at the nicking sites are listed in Table 1.

The same experiment was repeated for EhLINE1 by using contig 8160 from the E. histolytica genome database at the Sanger Centre. This contig contains the entire EhLINE1. Again, the boundary fragments were obtained by PCR amplification, and the bottom strand was labeled. The nicking pattern is shown in Fig. 3B, and the nicked sequences are listed in Table 1. Here, too, the enzyme nicked the bottom strand very close to the 3' end of the TSD in the fragment derived from the 3' end (band 2). However, unlike the corresponding band in the EhSINE1 insertion site, this band was not the most prominent (Fig. 2 and 3A). It is likely that mutations occurred after the original transposition event took place, making the sequence in contig 8160 a less favorable substrate. The fragment from the 5' end did not give very prominent nicks.

Once better assembly of the E. histolytica genome sequence is available, it may be possible to find some unoccupied sites of

FIG. 4. Conserved sequence features of EhLINE1 and EhSINE1 insertion. EhLINE1 sequences were obtained from the E. histolytica genome database at TIGR (sequences 1 to 6) and the Sanger Centre (sequences 7 to 13), while EhSINE1 sequences were from the GSS database. Boxes show the 3'-most sequence of the element. The TSDs at the two ends are uppercased. Regions of identity between them are underlined, while nonidentical nucleotides are marked by dots. The sequences immediately upstream of the TSD and at the 3' end of the element, which share close identity, are lowercased. Their regions of identity are underlined, and nonidentical nucleotides are marked by dots. The nearest C flanking the 3' TSD at its 3' end is boxed.
EhLINE1 and use them as substrates. Although the EhLINE1-encoded endonuclease lacks site specificity, it is clear from Table 1 that it nicks at some preferred sites. For example, T was the most common nucleotide found both 3ʾ and 5ʾ of the nick, followed by A. Another common feature was the presence of a G 3 to 4 nucleotides upstream of the nick in a large number of sites.

To gain further insight into the possible mode of retrotransposition of EhLINE1/SINE1, we analyzed the sequences flanking the inserted element, as available in the database. We selected entries containing full-length EhLINE1 and EhSINE1 (Fig. 4). The TSD could be readily identified in most cases and ranged from 3 to 22 bp. Two important features that emerged from this analysis are as follows. First, of 13 entries containing full-length EhLINE1 and 17 entries containing full-length EhSINE1 shown in Fig. 4, 21 of 30 contained a C (in the bottom strand) within 0 to 5 nucleotides from the TSD 3ʾ end, while the average occurrence of C in a 60-nucleotide stretch surrounding the TSDs in these entries was calculated to be 8%. Since a large number of nicks made by the endonuclease lay close to a 5ʾ G residue (Table 1), this finding strengthens the possibility that the G in the bottom strand may be involved in target site recognition and nicking by the endonuclease, leading to insertion of the element. Second, the TSD upstream of the element was almost always immediately preceded by a T-rich stretch. This stretch showed a very good sequence match with the 3ʾ ends of EhLINE1 and EhSINE1 (underlined in Fig. 4). Taking into account the features observed, we propose a model for retrotransposition of these elements (Fig. 5).

**DISCUSSION**

The non-LTR retrotransposons described so far can be classified into two broad categories based on the natures of the endonucleases encoded by the elements. One class encodes the apurinic endonuclease (APE), while the other encodes a restriction enzyme-like endonuclease (EN). Based on RT phylogeny, all the elements of the latter class belong to the R2 group, which is considered to be of ancient origin. The EN domain has been included in phylogenetic analysis, along with the RT domain to improve the resolution (7). Using this approach we show that EhLINE1 (as also EhLINE2 and -3) can be grouped in the R4 clade along with Rex6, Dong, R4, and NeSL (Fig. 6). Excluding some members such as Genie2 of *Giardia*, which is highly degenerate (7), and some vertebrate Rex6 elements where target sites have not been well defined (31), all well-characterized members of the R2 group show some level of site specificity and insert into defined sites (28S rRNA gene, spliced leader gene, TAA repeats, telomere). EhLINE1 is a clear exception. It is widely dispersed in the genome of its host. In addition, the properties of the EN domain of EhLINE1 reported here show that the enzyme is not strictly site specific and nicks a variety of related target sequences in vitro.

It is generally believed that target site selection is determined by the element-encoded endonuclease (28) and that consequently, elements with the restriction enzyme-like (EN) type of endonuclease would be site specific. However, the dispersed insertion pattern of EhLINE1 shows that other factors might be involved. This was also suggested by studies on elements belonging to the APE group, where most members...
FIG. 6. Phylogenetic positions of EhLINEs based on RT and EN domains. The analysis was performed essentially as described elsewhere (7). The entire sequence from the RT domain to the C terminus of the element (~600 amino acid residues) was included in each case. The tree was rooted on the Genie sequence. Numbers indicate bootstrap values at each node. The organisms to which the elements belong are as follows: GENIE1 and -1A, G. lamblia; CRE1 and CRE2, Crithidia fasciculata; SLACS, Trypanosoma brucei; CZAR, Trypanosoma cruzi; Rex6, Orzyias latipas; Dong, B. mori; R4, Ascaris lumbricoides; NeSL, Caenorhabditis elegans; R2-D. melanogaster, Drosophila melanogaster; R2-D. mercatorum, Drosophila mercatorum; R2-Isopod, Porcellio scaber.

are not site specific but many are known to insert at defined target sites (DRE, Tdd-3, and related elements in Dictyostelium discoideum, Zepp elements of Chlorella spp., R1 of Bombyx mori) (13). Although these elements inserted at defined sites, the endonucleases encoded by them were not strictly site specific. It was therefore proposed that apart from the endonuclease, other important factors, such as local chromatin organization and interaction with the transcriptional apparatus of particular genes, might be involved in target site selection (16). It would be interesting to know which factors may have led to the evolution of a loose target specificity of the EhLINE1 endonuclease.

Several studies have shown that SINEs, which do not encode their own transposition functions, exist in partnership with specific LINEs and utilize the enzymatic machinery of the latter for their own transposition (5, 22, 23). The consensus sequences flanking the site of insertion of human and rodent SINEs agreed very well with the experimentally observed hot spots of nicking by the human L1 endonuclease, suggesting that the latter may be involved in SINE mobilization (15, 18). The most compelling evidence in this regard has been the direct in vivo demonstration that eel LINE-encoded functions provided in trans could mobilize a partner SINE which shared the same 3' tail (19). Here we show the LINE/SINE functional linkage by using a different approach, namely, the ability of the LINE-encoded endonuclease to nick an empty target site, known to be occupied by the partner SINE in vivo. Although the EhLINE1 endonuclease is not strictly site specific and nicked the DNA substrate at several sites, it made a very prominent nick exactly at the point where EhSINE1 had inserted into the genome (Fig. 2), making it highly likely that this SINE element is mobilized by EhLINE1.

The nicking specificity of EhLINE1 EN provided important clues about the possible mode of transposition of this element. The endonuclease nicked the bottom strand at the right end of the TSD (Fig. 2 and 3), implying that the top strand nick would be toward the left of the TSD, resulting in 5' overhangs. This nicking orientation is also found for the endonuclease encoded by R2Bm (12, 37), whereas the APE enzymes encoded by L1 (15, 18), Tx1L (8), and R1Bm (16) nick the two DNA strands in the opposite orientation to give 3' overhangs. Due to this difference in nicking orientation, the model proposed for insertion of L1 (18) and R1Bm (16) is not directly applicable to EhLINE1. We propose a speculative model (Fig. 5) to account for TSDs with an enzyme that nicks to produce 5' overhangs. This model takes into account the conserved feature found in most EhLINE1/SINE1 insertions, namely, the presence of a T-rich stretch just upstream of the TSD, as shown in Fig. 4. According to this model, the EhLINE1/SINE1 transcript finds the target site by virtue of its T-rich 3' tail. This could happen either concurrently with or after the nicking of the bottom strand by the endonuclease. The 3'-OH of the RNA is then used by the RT (or a host enzyme) to copy the bottom strand, up to the nick. This disruption is followed by a template shift, such that the 3'-OH of the nick is used to prime reverse transcription of the RNA template, which now has a short DNA extension (the TSD). Alternatively, the RT uses the 3'-OH of the nick in the bottom strand to copy the DNA template for a short stretch (the TSD) until it can begin to reverse transcribe the element RNA by switching its template. The importance of the T-rich stretch upstream of the TSD, and that of other aspects of our model, needs to be experimentally verified. In the context of the RT switching its template, it is interesting that the RT encoded by R2Bm has been shown to jump from the 5' end of one RNA template to the 3' end of another (4). We have also noted the presence of a G upstream of the nicking site in the bottom strand (Table 1 and Fig. 4). Its involvement, if any, in the endonucleolytic cleavage reaction has to be tested by using appropriate oligonucleotides as substrates. This G, if essential, may have evolved to limit the number of insertion sites of EhLINE1/SINE1 in the AT-rich E. histolytica genome (17). In conclusion, the EhLINE1 element is a novel member of the R4 clade, which underscores the mechanistic diversity that may be encountered as more members of these lineages are discovered.

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