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Origin of the Human L1 Elements: Proposed Progenitor Genes Deduced from a Consensus DNA Sequence

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A consensus sequence for the human long interspersed repeated DNA element, L1Hs (LINE or KpnI sequence), is presented. The sequence contains two open reading frames (ORFs) which are homologous to ORFs in corresponding regions of L1 elements in other species. The L1Hs ORFs are separated by a small evolutionarily nonconserved region. The 5' end of the consensus contains frequent terminators in all three reading frames and has a relatively high GC content with numerous stretches of weak homology with AluI repeats. The 5' ORF extends for a minimum of 723 bp (241 codons). The 3' ORF is 3843 bp (1281 codons) and predicts a protein of 149 kD which has regions of weak homology to the polymerase domain of various reverse transcriptases. The 3' end of the consensus contains an adenine-rich end. The organization of the L1Hs consensus sequence resembles the structure of eukaryotic mRNAs except for the noncoding region between ORFs. However, due to base substitutions or truncation most elements appear incapable of producing mRNA that can be translated. Our observation that individual elements cluster into subfamilies on the basis of the presence or absence of blocks of sequence, or by the linkage of alternative bases at multiple positions, suggests that most L1 sequences were derived from a small number of structural genes. An estimate of the mammalian L1 substitution rate was derived and used to predict the age of individual human elements. From this it follows that the majority of human L1 sequences have been generated within the last 30 million years. The human L1 elements studied here differ from each other, yet overall the L1Hs sequences demonstrate a pattern of species-specificity when compared to the L1 families of other mammals. Possible mechanisms that may account for the origin and evolution of the L1 family are discussed. These include pseudogene formation (retroposition), transposition, gene conversion, and RNA recombination.

INTRODUCTION

The genomes of man and other mammals contain a large fraction of repeated DNAs. The functions, if any, of these sequences have remained elusive. Repeated DNAs can generally be cataloged as either tandem or interspersed on the basis of their distribution. The interspersed repetitive elements can be further characterized according to length as either short (SINES) or long (LINES) (Singer and Skowronski, 1985). The most commonly described LINES are the L1sp elements (where sp is the species designation). The human L1 elements have also been frequently referred to as KpnI sequences. There are at least 10^8 copies of the L1 element in the human genome, although the number of complete 6-kb elements is much smaller due to truncation (Singer and Skowronski, 1985). L1 elements are present in all mammals that have been studied (including both placental and marsupial mammals; Burton et al., 1986) and, based on their shared sequence homology in distantly related species, they most likely are derived from a common progenitor. Mammalian L1 repeats appear to evolve in concert such that the majority of elements within a species will be more like each other than they will be to the elements of another species. Consequently, Southern blots of restricted genomic DNA will often give species-specific patterns when hybridized with a LINE probe. However, when cloned elements from a particular species are compared, there are often significant differences between individual sequences.

Various investigators have identified open reading frames (ORFs) followed by an adenine-rich end in some L1 repeats from man and other species (Singer and Skowronski, 1985). L1-homologous transcripts have also been described in various cell types.
(Schmeckpeper et al., 1984; Kole et al., 1983; Shaftit-Zagardo et al., 1983) and a 6.5-kb polyadenylated RNA has been observed in teratocarcinoma cells (Skowronska and Singer, 1985). These observations have led to the description of LINEs as "retroposons" or "retrotranscripts" (Temin, 1985), suggesting that they may be derived by the integration of sequences involving either RNA or cDNA intermediates, as occur with retroviruses (Temin, 1985), and may occur in the formation of SINES (Weiner et al., 1986) and processed pseudogenes (Wagner, 1986). Likewise, similarities in size and organization and limited sequence homology with transposable elements from Trypanosoma (Kimmel et al., 1987), Drosophila (Fawcett et al., 1986), and yeast (Fink et al., 1986) suggest that L1 elements may also be related to sequences such as these. If either model is correct, it would follow that a small number of functional genes have given rise to the majority of L1 sequences, which have subsequently diverged from the progenitors by the accumulation of mutations. If such changes have occurred randomly in these nonfunctional copies, then it should be possible to deduce the sequence of the original structural gene or genes from which they were derived. This can be done by determining shared bases in a sufficiently large number of L1 sequences. To that end we have assembled an L1Hs consensus sequence based on L1Hs sequence data published in the literature and generated in our laboratories. As described below, this analysis has yielded information about the origin of the human L1 elements and the organization of proposed progenitor genes.

**MATERIALS AND METHODS**

The human 3' β-globin LINE, an apparently complete 6.2-kb sequence, has been described previously (Adams et al., 1980). The b2 LINE, an autosomal element, was derived from a genomic clone, λHb2, which had been obtained from a λ-phage library of human DNA (Schmeckpeper et al., 1981). The clones X63 and X65 were from a genomic library of AHA-11a DNA, a mouse–human hybrid cell line, containing an X as the only detectable human chromosome (Schmeckpeper et al., 1979). This library was made by partial EcoRI digestion and ligation into Charon 4A λ-phage. λ-Phage X63 and X65 were isolated on the basis of hybridization of L1 subclones derived from λHb2. A gorilla L1 element was obtained from a similarly constructed library (Scott et al., 1984) screened with a human β-globin cDNA probe. Restriction maps of the recombinant λ phases were prepared, and appropriate regions from these were subcloned into plasmids for further analysis. Because a majority of published sequences are homologous to the 3' end of the element, we have accumulated new data from underrepresented portions of the 5' end of the element.

Several sequences used to generate the consensus were derived from genomic fragments with particular restriction enzyme sites. Selection of sequences defined by particular restriction enzyme sites might bias the final consensus by selecting elements of a given type. In order to lessen this potential problem, the new elements analyzed for this paper were selected from genomic clones by hybridization criteria and because they were at least 6 kb long. The presence of particular restriction enzyme sites was not a criterion for selection. Further, many of the published elements were derived either by similar criteria or by the fact that they are adjacent to other genes or sequences. We observed no obvious clustering of elements into subgroups on the basis of how they were selected.

Fragments to be sequenced were ligated into either M13 phage (Messing and Vieira, 1982) or pEMBL plasmids (Dente et al., 1983). Random length clones were prepared by a variety of methods (Dale et al., 1985; Hong, 1982; Poncz et al., 1983; Barnes and Bevan, 1983). Portions of sequence were also obtained using specific synthetic oligomers as primers. DNA sequencing was done by the enzymatic chain-termination method (Sanger et al., 1977) using [35S]dATP. Most samples were run on 6 or 8% acrylamide–urea gels (40 cm long, 0.2- to 0.4-mm wedge gels). Selected samples were also run on long (80 cm long, 0.2 mm thick) water-jacketed gels heated to 55°C (Garoff and Ansorge, 1981). Regions of sequence overlap between clones were located with the aid of homology identification programs (Queen and Korn, 1984), while database searches were conducted using the programs of either Queen and Korn (1984) or Lipman and Pearson (1985).

The sequence of the 3' β-globin LINE used in this analysis was determined in our laboratory. The identical element has been sequenced independently by Hattori et al. (1985). Five differences in 6239 nucleotides were noted between the two sequences which could not be resolved by reexamination of the sequence generated in our laboratory. The GenBank database was searched for L1 homologous sequences that were aligned with respect to the β-globin element. Interspecies comparisons (Fig. 3) were made by aligning each pair of sequences, counting the number of matches per 60-nucleotide window, and then computing a moving average of three windows.

**RESULTS**

Figure 1 shows an overview of the organization of the human L1 element derived from the consensus sequence (Fig. 2). Figure 1 also indicates the relationship of the component sequences used in construction
of the consensus sequence. Each horizontal bar or group of bars per line represents DNA sequence from a separate element (as listed in Table 1). Some elements (e.g., sequences 5 and 6, Fig. 1 and Table 1) used in the consensus have regions that are scrambled relative to the 3' β-globin element. We believe that the β-globin LINE lacks rearranged or atypical regions because its organization best accounts for the placement of blocks of sequence from the other elements. The consensus human L1 DNA sequence and predicted amino acid sequences of the two ORFs are shown in Fig. 2. Because of space constraints, the consensus is shown without the individual sequences used in its assembly (the complete data set is available upon request). At three positions in the consensus sequence there have been insertions or deletions of blocks of sequence in more than one element. The deleted regions are indicated in the consensus by dashed lines. At present, each element characterized at these positions has at least one block of sequence deleted relative to the consensus. Whether there are full-length elements corresponding to the consensus sequence remains to be shown. At a number of positions we observed alternative bases occurring with sufficient frequency that no particular base could be unambiguously chosen (Fig. 2).

**LINE Organization**

As illustrated in Fig. 1 the human L1 consensus has a 5' region of about 1 kb that is presumably noncoding. This portion of the consensus has a noticeably higher GC content (Fig. 1), a number of areas that show weak similarity to human AluI sequences, and frequent termination codons in all three reading frames. The patches of homology to AluI sequence, while small (14 to 32 bases with 60 to 82% homology averaging 72%), were confined largely to the noncoding region 5' of the first ORF. This homology may reflect similar GC composition of the two classes of sequence rather than functional or evolutionary relatedness. Position 1327 has been designated as the beginning of the 5' ORF because a terminator occurs immediately before this position in two of the four individual sequences used in the consensus. Other terminators which are present in at least 50% of the component elements occur at 1179–1181 and 933–935. We presume that these truly reflect the consensus, but sequence from additional elements will be necessary to confirm this. If these positions are not true terminators, then the 5' ORF could be lengthened (although a longer 5' ORF would extend into sequence that is relatively poorly conserved, as discussed below). Figure 2 shows the proposed amino acid sequence of the extended 5' ORF in small letters. An in-frame terminator at position 924–926 occurs in all four elements used to deduce the consensus and presumably marks the most 5' that the first ORF could be extended. This possible extension of the 5' ORF is shown as a box with horizontal lines in Fig. 1. The reading frame beginning at 1327 is 241 codons long and would code for a protein of 218 amino acids (about 26 kDa) if the methionine encoded at position 1395–1397 is the amino terminus. A protein beginning at the methionine at nucleotides 1035–1037 in the extended 5' ORF would be 338 amino acids, approximately 40 kDa. A data base search failed to identify significant homology between the protein predicted by the 5' ORF and other protein sequences.

A small block of noncoding sequence (from 2049 to 2098; indicated by an asterisk in Fig. 1) was observed
FIG. 2. The sequence of the human L1 consensus is shown. Upper- or lowercase lettering is used to indicate the relative frequency at which a particular base occurs. A capital letter signifies that a base is present in 75% or more of the aligned sequences where fewer than six component sequences are available, or at two-thirds of positions when six or more component sequences are present. A lowercase letter indicates that the base occurred 65-74% of the time for fewer than six components, or 50% or better when multiple alternatives (i.e., \( t = t, t, a \) or \( t = t, t, a, c \)) or more than six component sequences were present. If no clear choice was possible, the base is indicated by a N. Where two choices appeared with equal or nearly equal frequency (e.g., 2:2 or 2:3), both alternatives are indicated. Where a particular base is predominant but a second alternative is seen repeatedly, then a combination of upper- and lowercase is used (e.g., c/T = 8T, 3C, 1A). Where two bases are each displaced from the line, they occur with equal or nearly equal frequency. The amino acid sequence (using the single-letter code) for both open reading frames is indicated below the nucleic acid sequence. Small letters are used to indicate the possible extension of the 5' ORF. The residue letter is shown directly below the first letter of the codon. Alternative residues are indicated only in cases where they are equally probable (i.e., residues predicted by codons with the less frequent nucleotides are not indicated). At some positions (especially in the 5' ORF) several alternatives are possible, but for clarity only one is indicated. A terminator occurs as an alternative at position 3765-3767 in the 3' ORF. A small deletion occurs as an alternative base in the 3' ORF at position 5311 and, if true, would shift the reading frame and create a smaller protein. Two insertions also occur in the 3' ORF at 3686 and 3695-3696. The alternative longer ORF would increase by one codon but would remain in frame.
### TABLE 1
Sequences Used to Derive the Consensus

| No. | Sequence name | Percentage homology with consensus | Description | Ref. |
|-----|---------------|-----------------------------------|-------------|------|
| 1   | 3' β-globin LINE | 97.1 (6026) | Genomic sequence | This paper and (19) |
| 2   | b2 LINE | 94.5 (3730) | Genomic sequence | This paper |
| 3   | X63 LINE | 97.1 (2935) | Genomic sequence from X chromosome | This paper |
| 4   | X65 LINE | 95.8 (1633) | Genomic sequence from X chromosome | This paper |
| 5   | HUMRSKPNP1 | 95.6 (3359) | Genomic sequence imbedded in alphoid repeat | (45) |
| 6   | ε-Globin LINE-1 | 72.7 (2223) | Genomic sequence adjacent to ε-globin gene | (6) |
| 7   | ε-Globin LINE-2 | 93.2 (2084) | Genomic sequence adjacent to ε-globin gene | (5) |
| 8   | HUMRSH3 | 96.8 (1905) | Genomic 1.9-kb HindIII restriction fragment | (38) |
| 9   | SHR4 | 98.2 (448) | Genomic sequence | (3) |
| 10  | HUMUG3PA | 89.1 (46) | Genomic flanking sequence near U3.2 small nuclear RNA pseudogene | |
| 11  | HUMRSKPA2 | 98.8 (2109) | Genomic sequence | (41) |
| 12  | HUMRSKPA6 | 97.3 (1756) | Fibroblast cDNA | (11) |
| 13  | HUMFIXG | 96.8 (1914) | Genomic sequence flanking Factor IX gene | (65) |
| 14  | HUMMHDDB1 | 95.5 (1363) | Genomic sequence flanking MHC DR β locus | (33) |
| 15  | HUMRSKPA9 | 96.7 (1229) | Fibroblast cDNA | (11) |
| 16  | HUMBLUR1 | 75.8 (264) | Genomic sequence adjacent to an Alu repeat | (8) |
| 17  | HUMIFNAGS | 89.3 (906) | Genomic sequence near α-interferon gene | (62) |
| 18  | HUMRSKPI | 96.4 (197) | Genomic 1.8-kb KpnI fragment | (42) |
| 19  | HUMRSKPE | 95.7 (1184) | Genomic sequence | (58) |
| 20  | HUMMIGHAD | 94.6 (617) | Genomic sequence near an Ig heavy chain ε3 pseudogene | (61) |
| 21  | 21-RING | 97.1 (751) | Genomic sequence from chromosome 21 | C. Wong et al.* |
| 22  | HUMRSKPA9 | 94.6 (720) | Fibroblast cDNA | (11) |
| 23  | HUMRSKPT | 92.4 (672) | Genomic sequence | (98) |
| 24  | HUMRSKPG | 92.9 (562) | Genomic sequence | (11) |
| 25  | HUMRSKPG8 | 83.4 (465) | Genomic sequence | (11) |
| 26  | HUMRSKPA8 | 95.3 (254) | Genomic sequence | (11) |
| 27  | HUMPPD16 | 95.1 (307) | Genomic sequence | (8) |
| 28  | Kpn6'1 | 76.0 (229) | Genomic sequence identified 5' of ε-globin gene | (35) |
| 29  | Kpn6'D | 96.1 (154) | Genomic sequence identified 5' of ε-globin gene | (35) |
| 30  | KTI | 93.4 (137) | Genomic sequence near an HIV integration site | (43) |
| 31  | HUMRSKPA2 | 98.5 (81) | Genomic sequence interrupting mitochondrial homologous DNA | (42) |
| 32  | PAC-32 | 92.9 (209) | Genomic sequence | J. W. Adams et al.* |
| 33  | HUMRSA16 | 92.2 (115) | Genomic sequence near Alu repeat | (58) |
| 34  | HUMRSKPA1 | 100.0 (94) | Genomic sequence | (41) |
| 35  | HUMRSKPA6 | 97.1 (103) | Genomic sequence | (41) |

**Note.** Numbers correspond to sequences diagrammed in Fig. 1 (except for nonhuman elements). GenBank filenames are given when available. The percentage homologies of individual sequences relative to the consensus used in Fig. 4 are given along with the length of homologous sequence in parentheses.

* Unpublished data.

* Nonhomologous 5' and 3' regions were excluded from this calculation.

* Consensus based on multiple sequences.

between the 5' and 3' ORFs. No obvious splice junctions were found bordering the noncoding region between ORFs. The consensus in the noncoding region is based on six sequences and it is unlikely that the separation of the two reading frames is a consequence of incomplete data. The 3' ORF extends from position 2099 to 5942 for 1281 codons and has homology with various reverse transcriptases (Hattori et al., 1986; Sakaki et al., 1986). If the putative protein begins with the methionine encoded at position 2115-2117, then it...
would be 1276 residues long. The 3' ORF is followed by another nonconserved block of sequence that is 208 bp long with multiple terminators in all three reading frames. The L1 consensus terminates in a short stretch of adenines. A search of the L1Hs sequence in Fig. 2 failed to identify likely ribosome-binding sites near possible initiator methionines (Kozak, 1986). Further, no obvious promoter sequences were identified 5' of either ORF.

In order to validate our observations about the human consensus, we compared published sequences from the mouse and rat to each other and to the human consensus (Fig. 3). Figure 3A shows a comparison of percentage homology of aligned mouse and rat L1 sequences calculated as a running average of three 60-nucleotide windows. A large region of conserved sequence is seen flanked by 5' and 3' ends which have markedly less homology. Furthermore, a short region of reduced homology (approximately 1979 to 2129) can be seen as indicated by the arrow. When the human consensus sequence is then aligned relative to the mouse–rat comparison in Fig. 3A, it can be seen that a good correspondence occurs between the predicted 5' and 3' noncoding regions from the L1Hs consensus and the regions of least homology between the two rodents. The sequence in L1Hs between the 5' and 3' ORFs aligns well with the area of reduced homology in the rodent comparison (asterisk in Fig. 3A). However, the region of reduced homology is longer than the gap between L1Hs ORFs because approximately the last 23 codons of the 5' ORF and the first 10 codons of the 3' ORF extend into nonconserved region. When either the mouse or the rat sequence is compared directly to the human sequence (Figs. 3B and 3C), the region between the human ORFs shows reduced homology in each case. The region corresponding to the 3' ORF is relatively well conserved for both comparisons, since the sequence homology is as high as 65%. However, it appears that the length of the conserved sequence in the 5' ORF is shorter in the human–rodent comparison than in the rat–mouse comparison. We also observed that this portion of the 5' ORF gene product was less well conserved than the 3' product, suggesting a possible species-specific function for this sequence. On the basis of these comparisons we are confident that the functional human L1 element (and by extension the rodent elements as well) must code for two separate proteins. Two ORFs have also been reported for the mouse L1 element, although in that species the 5' reading frame extends (in a different register) into the 3' ORF (Loeb et al., 1986). The fact that the 5' end of the element is poorly conserved, not only between the human and rodent elements but also between mouse and rat sequences, implies that this region does not contain large blocks of functionally important sequence (at least as measured by the criterion of evolutionary conservation).

Subfamily Analysis

Comparison of individual L1Hs elements to the consensus sequence (Fig. 2) indicates that subfamilies can be defined both on the basis of the association of alternative bases and by the presence or absence of blocks of sequence that occur at the same positions in different elements. Three such blocks (at L1Hs consensus positions 143–157, 710–716, and 782–909) can be used to define at least three subfamilies at the 5' end as shown in Table 2. Subfamily 1 (as typified by the 3' β-globin LINE) is characterized by a deletion from 710 to 716. Subfamily 2 (the X63 and Kpn-A elements) has a deletion from 782 to 909. In subfamily 3 (the b2 LINE) deletions occur from 143 to 157 and 710 to 716. PAC-32 LINE also has the subfamily 3 deletion from 782 to 909. In subfamily 3 (the b2 LINE) deletions occur from 143 to 157 and 710 to 716. PAC-32 LINE also has the subfamily 3 deletion from 143 to 157; however, since PAC-32 sequence does not extend into the region of the consensus where the other deletions are found, it is not yet possible to assign it to a subfamily. It could belong to subfamily 3 or represent a fourth subfamily. Other elements whose sequences include only a portion of this region may represent additional subfamilies.

FIG. 3. Interspecies comparisons of L1 sequences are shown. DNA sequences were aligned and matched bases were counted in windows of 60 nucleotides. A moving average of three windows was calculated and plotted. Each of the plots is shown aligned with a schematic of the human element. Note that the scale for the rat–mouse comparison is changed to reflect the greater similarity between these species. Arrows above the asterisks point to the local homology minima corresponding to the evolutionarily nonconserved region between ORFs. (A) Mouse vs rat (elements 41 and 40, respectively; Table 1). (B) Human consensus vs rat. (C) Human consensus vs mouse.
TABLE 2
Variation at the 5' End of L1Hs Defines Subfamilies

| Subfamily type | L1Hs consensus sequence position | Example |
|----------------|---------------------------------|---------|
| 1              | +                               | 3' fl-Globin LINE [1] |
| 2              | +                               | X63 LINE [3] |
| 3              | -                               | Kpn-A LINE [5] |
| ?              | -                               | b2 LINE [2] |

Note. Three major subfamilies at 5' end of the consensus are defined by the presence (+) or absence (−) of sequence blocks (see Fig. 2). Numbers in brackets refer to sequences listed in Table 1.

However, it should be noted that all of the subfamilies can be accounted for by mixing blocks of sequence from as few as two different progenitor sequences. For example, 5' subfamily 1 could result from a small conversion of a type 2 element by sequence from a type 3 element that includes the two blocks between 710 and 909. Additional evidence for subfamily groups can also be seen by the concordance of particular alternative bases at several positions at the 3' end (Table 3). Several elements can be clearly assigned to either of two subfamilies, although it is important to note that evidence for mixing and the possibility of further subdivisions are suggested by the data. The existence of multiple subfamilies has also been inferred from restriction mapping studies of genomic DNA (Jubier-Maurin et al., 1985; B. J. Schmeckpeper et al., unpublished data). Nevertheless, the analysis presented here clearly suggests that the majority of L1 elements are derived from only a few classes of similar L1 progenitors, perhaps as few as two functional genes.

Age of L1Hs Elements Estimated by Evolutionary Analysis

Figure 4A illustrates the data presented in Table 1 on the homology between individual L1Hs sequences (Fig. 1) and the consensus sequence (Fig. 2). Most elements are less than 5% different from the consensus, although a few sequences are as much as 25% different. For this analysis a match to an alternative base in the consensus was accepted as perfect homology. Thus, differences among the subfamilies should not cause an overestimation of the percentage divergence of individual elements from either of the two proposed L1 progenitor sequences. During this analysis we also observed that the percentage divergence by which any particular element differs from the consensus tends to be the same throughout the entire length of the element.

Figure 4B illustrates L1 sequence homology between the human consensus (Fig. 2) and sequence data from other mammals (Table 1). The first comparison is from sequence at the 5' end of the orthologous L1 element adjacent to the gorilla β-globin gene. In 164 bp of homologous sequence there are only three differences. This value of about 2% change between man and gorilla, which diverged 8 to 10 million years (MY) ago, was also seen in other regions of the β-globin cluster (Scott et al., 1984). A second comparison is available from the sequence of elements from the African green monkey (Cercopithecus aethiops), a cat-

TABLE 3
Variation at the 3' End of L1Hs Elements Defines Two Major Subfamilies

| Subfamily type | L1Hs consensus sequence position | Example |
|----------------|---------------------------------|---------|
| 1              | 3' fl-Globin [1]                | T C A T |
| 1 HUMRSK04     | [15] g                         | T C A T |
| 1 HUMRSK08     | [12] g                         | T C A T |
| 1 HUMFIXG      | [13] T C A T                    | T C A T |
| 2              | c-Globin 2 [7]                 | A T G C |
| 2 HUMIGHAD     | [20] g                         | A T G C |
| 2 HUMIFNAGS    | [17] A T G C                    | A T G C |
| 2 HUMRSKPT     | [23] g                         | A T G C |
| 2 HUMRSKP84    | [25] g                         | A T G C |

Note. Two major subfamilies can be identified by associated bases from several elements at the 3' end of the consensus from position 5103 to 6000. Most individual elements (Table 1) clearly fall into one of these subfamilies. Bases that do not correspond to either subfamily are shown in small letters. Discordant bases, which occur in a sequence from one subfamily but are most often seen in the other, are underlined.
tarbine primate which separated from the hominoids about 25 MY ago. Three partial monkey L1 sequences were compared to the human consensus. Overall homology with the human consensus is 95% for two and 92% for the third. If we assume that there are no, as yet uncharacterized, monkey elements more like the human consensus, then the 5% difference between monkey and human sequences can be used as an estimate of the percentage divergence that has occurred in 25 MY. The percentage similarity of rabbit, rat, and mouse sequences to the human consensus ranges from 64 to 69%. Since the mammalian radiation occurred about 80 MY ago, these species establish a third measure of the amount of sequence divergence with time. Altogether, the interspecies comparisons can be used to generate an approximation of the mammalian L1 substitution rate.

If the substitution rate calculated from the interspecies comparisons can be applied to elements within a species, then the age of particular L1Hs sequences can be determined. By such an analysis most are between 10 and 30 MY old. The validity of this analysis is dependent on several factors, the most important of which is how well the consensus reflects the proposed L1 structural genes. If there are more than the two L1 structural genes which we have inferred from the subfamily data, then the percentage homology will be greater and the predicted age of particular elements will be reduced. Nevertheless, a testable hypothesis follows from this model: that the estimated age of insertion of particular L1Hs elements can be independently determined by observing whether orthologous L1 sequences are present in the genomes of other primates. For example, of the five separate LINES in the human β-globin gene cluster (open bars, Fig. 4A) two are apparently quite old. The two oldest elements (Table 1, Nos. 6 and 28) appear to have inserted near the ε-globin gene about 65 MY ago, while the most recent element (the 3' ε-globin L1) may have integrated as recently as 25 MY ago. We know that the element present near the gorilla β-globin gene is the result of the same integration event as the human sequence, because its position relative to the gene is identical in both species and the first variable sequence block (position 143-155) is present in both elements. Although sequence data are not available, restriction maps and hybridization studies indicate that the 3' β-globin L1 is also present in orangutans and chimpanzees but absent in the Old World monkeys (Fujita et al., 1987). This is consistent with an age of insertion occurring approximately 25 MY ago which coincides with the estimated time of the hominoid–cattarine divergence. Therefore, its integration may have occurred shortly after the divergence of the Old World monkey and hominoid lineages. By the same reasoning those elements least similar to the consensus, such as those near the ε-globin gene, might be present in primates with earlier divergence dates. Indeed, Rogan et al. (1987) have shown that at least one of the ε-globin LINES is present in the Galago (a prosimian) which diverged from the higher primates at least 60 MY ago. Thus, the existing data, though limited, support the assumptions used to date the human elements.

**FIG. 4.** (A) The percentage homology of individual human elements relative to the L1Hs consensus sequence (Table 1). Open bars, from left to right, represent the 3' β-globin element, Kpn512, ε-globin LINE-2, Kpn51, and ε-globin LINE-1. (B) The percentage homology of nonhuman elements to the human consensus. Estimated divergence dates are indicated above the figure. The position of the gorilla β-globin element is indicated (G) and represents the percentage similarity between the gorilla and human orthologous sequences rather than the similarity between the gorilla element and the human consensus. The African green monkey (M) sequences (elements 36-38; Table 1) are plotted relative to the consensus and are shown at 25 MY. Other mammals include (from left to right) the rabbit, mouse, and rat (elements 39-41; Table 1) with a divergence date of 80 MY. The percentage homology for the non-primate comparisons is calculated for sequence corresponding to a portion of the 3' ORF from the rabbit and from the combined 5' and 3' ORFs of the mouse and rat. The less homologous flanking regions were excluded for this analysis.

**DISCUSSION**

Given that at least 1% of the human genome consists of L1 elements, how can we account for their origin and proliferation? The data presented here support the view that a relatively small number of similar L1 structural genes are the source of a much larger number of apparently nonfunctional elements which resemble processed pseudogenes. In the discussion below we make the assumption that transcription of an L1 structural gene produces a 6-kb mRNA containing two separate reading frames. However, transcription beginning at sites within the proposed structural genes is also possible. The observation that many elements appear to be truncated at the 5' end could be accounted for by reintegration of
RNAs derived from multiple sites of initiation in the L1 structural genes or from various dispersed elements that remain transcriptionally active but do not produce a translatable mRNA. Indeed, C. T. Comey et al. (unpublished data) have recently shown that a portion of the 3′ β-globin LINE within the 3′ ORF has specific pol II promoter activity in an in vitro transcription system. Truncation of 5′ sequence is also consistent with a retroposition model in which L1 mRNAs are incompletely copied by a polymerase such as reverse transcriptase.

A retroposition model for the origin of the L1 family has often been proposed. In support of this model, several groups have analyzed the relationship between the putative L1 gene products and reverse transcriptases. As noted above, the amino acid sequence of the 3′ ORF predicts a protein very similar to that of Hattori et al. (1986) which contains regions of weak homology to the polymerase domain (Johnson et al., 1986) of various reverse transcriptases (see Hattori et al. (1986) and Sakaki et al. (1986) for amino acid sequence alignments with these enzymes and the gene product of the 3′ ORF). Recently, Fanning and Singer (1987) constructed a mammalian amino acid consensus sequence for portions of the 3′ ORF by comparing the predicted L1 proteins from four different orders of mammals (cat, mouse, rabbit, and a combined human and monkey sequence). They concluded that the most highly conserved regions between species also corresponded to regions shown by Hattori et al. (1986) and Sakaki et al. (1986) to be homologous with various reverse transcriptases. Although these homologies are provocative, experimental evidence substantiating the retroposition model is needed. If the gene product of the 3′ ORF can be shown to have a polymerase activity, it would greatly strengthen the hypothesis that L1 elements originated by a retroposition mechanism and may be related to transposable elements or the general class of RNA viruses.

The origin of the L1Hs family can also be investigated by analysis of the subfamilies that we observed (Tables 2 and 3). As noted above, the study of individual elements suggests that the subfamilies can be accounted for by the mixing of blocks of sequences from as few as two structural genes. This could be the consequence of a process such as gene conversion involving DNA exchange between homologous sequences. Gene conversion in mammals often appears to be the result of homologous recombination between nearly identical, tandemly linked sequences, such as the γ-globin loci. Whenever conversion occurs at the level of the L1 structural genes, transcripts with mixed blocks of sequence would be generated. The reintroduction of such transcripts would result in the accumulation of dispersed elements with the observed “patchwork” appearance. Another mechanism that would account for the apparent mixing of L1 sequences is one similar to that proposed for the recombination seen among certain RNA viruses (Keele et al., 1987; Kirkegaard and Baltimore, 1986). For example, poliovirus, which contains a (+)-strand RNA genome, is copied by RNA polymerase to a (−)-strand intermediate. Evidence has accumulated that the RNA polymerase can switch to a second strand if it encounters a break during replication (Kirkegaard and Baltimore, 1986). If virus-infected cells contain two heterologous RNA (+)-strands, a hybrid (−)-strand can be formed. By analogy, if RNA recombination also occurs between L1 transcripts arising from the two proposed structural genes, then the subfamilies of these dispersed elements could be explained as retroposition events involving recombinant transcripts.

L1 elements occur in relatively high numbers in all mammals that have been studied (Burton et al., 1986). They not only share homology between species, but they also show a pattern of species specificity detectable by both DNA sequence analysis and the presence of discreetly sized bands following Southern blotting with L1 probes. This pattern of within-species similarity is most obvious when distantly related species are compared, but it has also been observed when elements from different mouse species were analyzed (Jubier-Maurin et al., 1985; Martin et al., 1985) or when monkey and human elements were aligned (A. F. Scott, unpublished data). The relative similarity of the L1Hs elements used in constructing the consensus sequence (as manifested by the clustering of elements in Fig. 4A) is another measure of the within-species homogeneity of this repeated DNA family. At least three models can account for the species-specificity. The first would require the generation of the L1 repeated DNA family by the sudden amplification in different taxa of a small number of functional L1 elements, as proposed for the SINEs (Weiner et al., 1986). If true, such amplifications would have had to occur recently in each species in order that individual elements would not have had sufficient time to diverge by the accumulation of random base substitutions. However, a difficulty in accounting for all intraspecies similarity of L1 sequences by this model is the necessary assumption that independent amplification events have occurred in each mammalian species. It is possible that such a predisposition to generate large numbers of dispersed elements is an intrinsic characteristic of the functional L1 genes. However, the observation (Fig. 4A) that some of the L1Hs elements are quite old argues against a single burst of amplification as the source of the L1Hs repeated DNA family.

A second model that would account for the within-species similarity of L1 sequences involves gene con-
version. These conversion events would have to affect a large fraction of the thousands of dispersed L1 elements in order to maintain the observed intraspecies homogeneity of this repeated DNA family. Such widespread conversion could occur by an RNA-mediated mechanism involving transcripts newly generated from the structural genes or by exchange of DNA sequences between various dispersed elements. In both yeast and fungi, conversion between dispersed tRNA genes occurs at relatively high frequency (Heyer et al., 1986) and may involve RNA intermediates (Doolittle, 1985). In the case of the L1Hs family, conversion of dispersed elements is unlikely, because the alternative bases in Fig. 2 and Table 3 cluster into two groups with relatively little discordance. In addition, the percentage by which a given element differs from the consensus appears to be the same throughout the length of that element. This observation is opposite to the expectation that conversion of dispersed elements might produce LINEs with “new” sequence imbedded in otherwise “old” elements.

If neither conversion of dispersed elements nor recent bursts of amplification are adequate to account for within-species homogenization, how else can we account for this process? A third model that might explain the characteristics of the L1 repeats would require their continuous generation in each mammalian lineage. Indeed, evidence for new L1 integrations has been presented from studies of a canine myc gene (Katzir et al., 1985) and from the rat Mbi-2 (Economou-Pachnis et al., 1985), immunoglobin heavy chain (Economou-Pachnis et al., 1985), and insulin (Lakshmikumaran et al., 1985) loci. In each case allelic forms can be distinguished by the presence or absence of an associated element. If L1 elements are continuously produced, then we would expect an even distribution of sequences with a range of homology to the consensus instead of the clustering of “new” elements seen in Fig. 4A. The observed distribution could be accounted for either by a bias of ascertainment, so that older elements are less likely to be detected with probes most like the consensus, or by processes that remove DNA from the genome. As noted above, we detected no obvious bias of ascertainment of the elements in Table 1. We favor, therefore, the latter hypothesis. Support for this process can be found from several observations including differences in the lengths of intervening sequences and in the organization of nonfunctional DNA adjacent to orthologous genes in different species. Likewise, the fact that about 5% of mutant genes characterized in a variety of human genetic disorders are the result of deletions (Kazazian and Antonarakis, 1987) suggests that loss of large blocks of sequence is not uncommon. If removal of DNA from the genome occurs with moderate frequency, then as a consequence L1 sequences will be removed. Therefore, DNA loss coupled with new integration events and the decay of older L1 sequences by base substitution could produce the appearance of relatively recent L1 amplifications. Clearly, the mechanism or combination of mechanisms that accounts for the generation and evolution of these sequences remains to be fully understood.

The L1 consensus sequence suggests that transcripts of the L1 structural genes are distinct from typical processed eukaryotic mRNAs in that they have two ORFs separated by a noncoding region. The two reading frames and the reverse transcriptase homology of the 3' ORF are characteristics shared with retroviruses, the Ty elements of yeast (Fink et al., 1986), and the I factor transposable elements of Drosophila (Fawcett et al., 1986). The possibility arises that all of these sequences are functional analogs and may have had a common evolutionary origin. If the 3' ORF codes for a reverse transcriptase, then, as with retroviruses, that activity might enhance the copying and reintegration of L1 transcripts into the genome (Temin, 1985; Hattori et al., 1986). The human L1 consensus sequence differs from that of typical retroviruses in that it lacks terminal repeats and has 5' and 3' nontranslated and nonconserved flanking regions together with an adenine-rich 3' terminus. Although the origin of the L1 sequences is unclear at present, it is apparent that this class of DNA is very old. Perhaps the original L1 structural gene was derived from the integration of an RNA virus or transposable element prior to the mammalian radiation, and with selection the sequence was modified to become a conventional eukaryotic gene which continues to retain features of its origin. Assuming a retroposon mechanism for the generation of the high copy number for L1 elements, it follows that the structural genes may be transcribed in germ line cells. The demonstration of a large polyadenylated L1 RNA in certain teratocarcinoma cells (Skowronski and Singer, 1985) suggests that the structural genes may be expressed in ova. Because mammalian eggs are present in females at birth and can be maintained for long periods before fertilization, their DNA may represent excellent targets for invasion by sequences transcribed in these cells (Wagner, 1986). Although a functional role for L1 sequences remains unknown, it is not unreasonable to expect that their integration in or near genes might alter the expression of adjacent sequences and that they might serve as insertional mutagens. While single elements may have a small influence on neighboring genes, the cumulative effect of thousands of elements throughout the genome may be profound and, in part, account for the rapid rate of phenotypic change seen in mammalian evolution.

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