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Genome Res. 2002 12: 391-399

Access the most recent version at doi:[10.1101/gr.216902](https://doi.org/10.1101/gr.216902)

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Cold Spring Harbor Laboratory Press

Processed Pseudogenes of Human Endogenous Retroviruses Generated by LINEs: Their Integration, Stability, and Distribution

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We report here the presence of numerous processed pseudogenes derived from the W family of endogenous retroviruses in the human genome. These pseudogenes are structurally colinear with the retroviral mRNA followed by a poly(A) tail. Our analysis of insertion sites of HERV-W processed pseudogenes shows a strong preference for the insertion motif of long interspersed nuclear element (LINE) retrotransposons. The genomic distribution, stability during evolution, and frequent truncations at the 5' end resemble those of the pseudogenes generated by LINEs. We therefore suggest that HERV-W processed pseudogenes arose by multiple and independent LINE-mediated retrotransposition of retroviral mRNA. These data document that the majority of HERV-W copies are actually nontranscribed promoterless pseudogenes. The current search for HERV-Ws associated with several human diseases should concentrate on a small subset of transcriptionally competent elements.

[Online supplementary material available at <http://www.genome.org>]

The human genome contains two major classes of autonomous, retrotransposition-competent elements encoding the reverse transcriptase and mobilized via an RNA intermediate, long interspersed nuclear elements (LINEs or L1) and human endogenous retroviruses (HERVs). Long terminal repeat (LTR)-containing HERVs comprise ~4%–5% of the genome (International Human Genome Sequencing Consortium 2001). LINEs, retrotransposons lacking LTRs and containing poly(A) tails, are the most active autonomous transposable elements in the human genome (Kazazian and Moran 1998; Prak and Kazazian 2000). They are estimated to be present in >500,000 copies, comprising 17% of the genome (Smit 1996a, 1999; International Human Genome Sequencing Consortium 2001). However, due to 5' truncations and deleterious mutations, only 30–60 LINEs per haploid genome are active and transpose along the genome (Sassaman et al 1997; Kazazian 1999; The database of Retrotransposon Insertion into the Human Genome http://www.med.upenn.edu/genetics/labs/retrotrans_table.html). LINE elements have a broad impact on the genome structure; beside occupying a large portion of the genome by themselves, the LINE copies are most probably involved in the expansion of Alu elements, which comprise 10% of the genome (Smit 1996a, 1999; International Human Genome Sequencing Consortium 2001). LINE transduction of the 3'-flanking sequences is estimated to account for a further 0.5%–1% of the genome (Goodier et al. 2000; Pickeral et al. 2000; International Human Genome Sequencing Consortium 2001).

Another type of genomic element generated by the LINE machinery is called processed pseudogenes. They were first described as pseudogenes structurally colinear with gene mRNA, lacking promoters, introns, and, in general, without protein-coding capacity due to mutations and frequent stop codons. Their mRNA-derived structure, poly(A) tails at the 3' end and the presence of direct repeats of variable (5–15 bp) length led to the hypothesis that their formation requires reverse transcriptase, and these pseudogenes were termed processed pseudogenes (Vanin 1985; Weiner et al. 1986). In search of this reverse transcriptase activity, Tchènio et al. (1993) showed the generation of processed pseudogenes from intron-containing proviral structures in murine cells. Elimination of sequences essential in *cis* for the retroviral life cycle indicated that endogenous retroviruses are not involved in this process. Later, this endogenous reverse transcriptase activity was shown to generate processed pseudogenes also from nonviral (non-LTR) constructs in somatic HeLa cells (Maestre et al. 1995). Along the same line, evidence available from other experiments disclosed that retroviral infection, as well as forced expression of retrovirus-like structures, resulted in all cases in cDNA genes without the typical structure of processed pseudogenes (Dornburg and Temin 1990; Levine et al. 1990; Derr et al. 1991). In direct *in vitro* tests, LINEs, and particularly LINE ORF2, were necessary for production of the typical cDNA structures similar to processed pseudogenes, whereas Moloney murine leukemia virus as well as human immunodeficiency virus type 1 (HIV-1) were unable to form the expected cDNAs, probably due to the lack of essential structures such as the primer-binding site for tRNA in the vector (Dhelliin et al. 1997). An independent support in favor of LINEs in the generation of processed pseudogenes comes from the computational comparison of insertion sites of LINEs, Alus, and processed pseudogenes. They all share simi-

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Article and publication are at <http://www.genome.org/cgi/doi/10.1101/gr.216902>. Article published online before print in February 2002.

lar features including a common TT|AAAA insertion motif and a variable-length (5–15 bp) target site duplication (TSD) (Jurka 1997), whereas retroviruses are characterized by short 4–6-bp direct repeats and no preference for a motif similar to TT|AAAA. Moreover, poly(A) tails and frequent truncation at the 5' end of processed pseudogenes are typical for LINES (Voliva et al. 1983; Smit 1999), indicating again that LINES are the master mobile elements in the human genome. Finally, an *in vitro* assay clearly showed creation of reporter gene copies by the LINE machinery with all hallmarks of the processed pseudogene (Esnault et al. 2000).

The estimated portion of processed pseudogenes in the human genome is ~0.5% (Dunham et al. 1999), with copy number 23,000–33,000 (Goncalves et al. 2000). The LINE machinery influences not only the pseudogene structure, but also the pseudogene distribution. We reported recently that not only young LINES and Alus, but also processed pseudogenes, preferentially reside in GC-poor parts of the genome (Pavlíček et al. 2001).

During our work on the Human Endogenous Retrovirus database (<http://herv.img.cas.cz>; Pačes et al. 2002), we found unusual retroviral structures lacking the common proviral organization. They are structurally similar to retroviral mRNA, followed by a poly(A) tail. These elements are frequently observed in the HERV-W family (Blond et al. 1999), described previously as LM7 or multiple sclerosis-associated virus (Perron et al. 1989, 1997), and as endogenous HERV17 family in the Repbase Update database (<http://www.girinst.org/>; Jurka 1998, 2000; Smit 1999). The HERV-W family is of special interest because of suggestions of a role in several human diseases including multiple sclerosis (Perron et al. 1997; Kormurian-Pradel et al. 1999), rheumatoid arthritis (Gaudin et al. 1997, 2000), and schizophrenia (Karlsson et al. 2001). Moreover, the *env* gene of a prototype HERV-W element on chromosome 7 was suggested as the human gene coding for the syncytin protein responsible for cell fusion during the differentiation of the syncytiotrophoblast in human placenta (Smit 1999; Blond et al. 2000; Mi et al. 2000). This is one of the best-documented examples of recruitment of retroviral genes by the host genome.

This work describes, for the first time, the presence of processed pseudogenes from mRNA of HERV-Ws in the human genome. These pseudogenes are remarkably numerous in the HERV-W family and are not found at comparable frequencies in related retroviral families of the class 1 HERVs. This finding represents not only an interesting connection

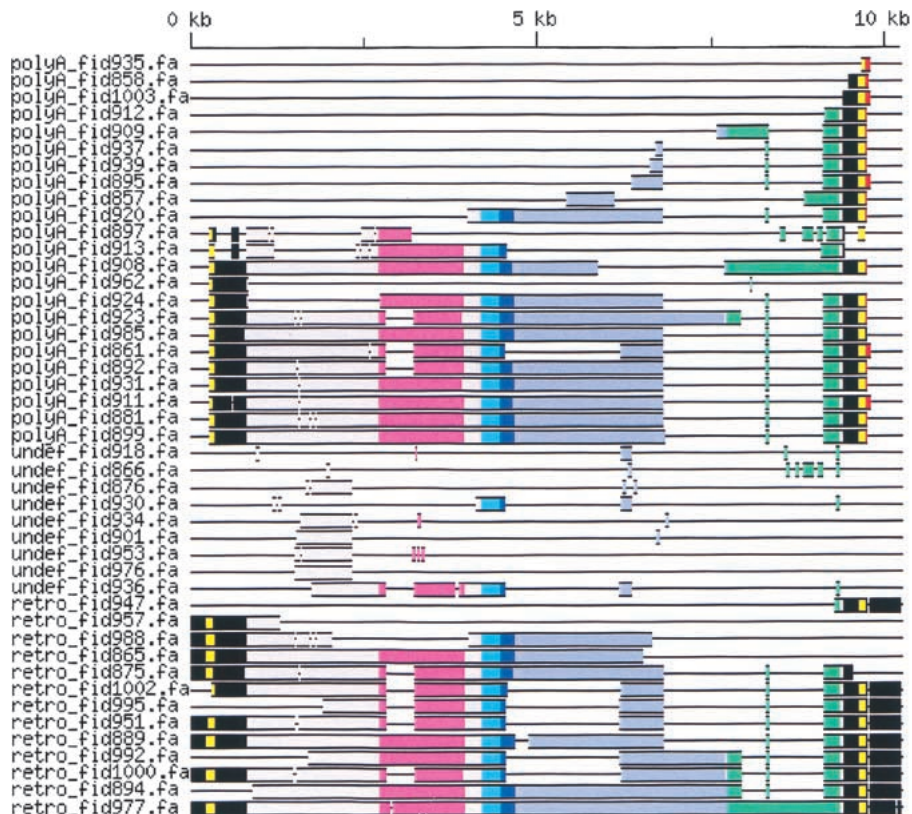


Figure 1 Genomic structure of HERV-W copies on chromosomes 6 and 7 shown schematically as a multiple alignment of elements. Only internal sequences containing elements are included, 35 soloLTRs are not shown. Sequences were aligned to the Repbase Update consensus (see Methods). In the consensus of 3' LTR, we introduced 50 A nucleotides to show the poly(A) tail (positions 9733–9782 in red). The LTR U5 and U3 regions are shown in black (positions 1–780 and 9407–10236), LTR R regions are in yellow. They were defined from TATA-box (TATAAA sequence, pos. 228–233 and 9634–9639 for the 5' and 3' LTR, respectively) to poly(A) signal (ATTAAA, pos. 308–313 and 9714–9719). ORFs are colored as follows: *gag* (pos. 2718–3923) in purple, *pro* (pos. 4192–4641) in cyan blue, *pol* (pos. 4450–7692) in light blue, and *env* (pos. 7720–9348) in green. The overlapping region of *pro* and *pol* is in dark blue. Other regions are contrasted in gray. In addition to several deletions, several splice variants are visible, see also Supplementary Figure 1S (online at <http://www.genome.org>). The last element, *retro_fid977*, is the syncytin-coding virus on chromosome 7.

between two main reverse transcriptase-encoding classes of retroelements, LINES and HERVs, but, as we show, also a unique model for comparing LINE and retroviral transposition, mRNA preference, integration specificity, recombination, and genomic stability.

RESULTS

Determination and Genomic Structure of HERV-W Retroviral and Pseudogene Copies

Using the genome-wide RepeatMasker scan, 654 HERV-W family members were found. From their multiple alignment schematically depicted in Figure 1 (see also Supplementary Figure 1S available online at <http://www.genome.org>), it is clear that HERV-W sequences are divided into two major groups. The first group is defined by the complete or partial U3 region in 5' LTR and/or U5 in 3' LTR and will be referred to as the retro group (or retro copies) in the following text, because the mentioned parts of LTRs are completed during retroviral reverse transcription. This retroviral group contains 77 proviral copies with complete or partial internal (non-LTR)

Table 1. Number of Different HERV-W Elements in the Human Genome

Group	Number	Complete	5' truncation ^a	3' truncation ^a	5'3' truncation ^a
poly(A)	176	46	107	23	0
retro (internal sequence-containing)	77	29	15	24	9
retro (soloLTR)	343	306	20	13	4
undef	58	0	0	0	58

^aTruncations were defined according to alignments with a particular consensus sequence for each group. The terminal gap was counted if the start or the end of the element was more distant than 20 bp from the consensus termini.

sequences, and 343 soloLTRs without any internal sequences (Table 1).

The members of the second group are defined by incomplete LTRs that start close to the beginning of the R region of 5' LTR (from nucleotide 256 of the consensus sequence, see Methods) and end at the 3' part of the 3' LTR R region (up to nucleotide 9732 of the consensus). The second characteristic feature is the frequent presence of the poly(A) tail of variable length and, therefore, we term these HERV-W sequences as the poly(A) group [or poly(A) copies] in the following text. The lengths of the tails vary from a few nucleotides to >50 bp. The poly(A) tail starts at position 9732, just 12 nucleotides downstream of the poly(A) signal ATTTAAA, in the 3' LTR (positions 9714–9719). In the poly(A) tails, we often detected A-rich microsatellites of up to 10 repeat units. The poly(A) tails end, in general, with the 3' direct repeat (Fig. 2). We have found 176 members of the poly(A) group (Table 1).

Aside from these two groups, there are 58 truncated elements, which, due to the absence of diagnostic parts, cannot be unambiguously defined and are referred to here as the undef group (Table 1).

Analysis of terminal deletions shows further intergroup differences. Whereas the poly(A) group is, in the great majority of cases, truncated at the 5' end, retro copies and soloLTRs are nearly equally truncated at both ends (Table 1). In addition, soloLTRs are mostly complete, but the poly(A) and the retro copies of HERV-Ws are generally shortened (Table 1).

Interestingly, in addition to the poly(A) tract, HERV-W genomic copies display further oligonucleotide expansion lo-

cated in the leader region (positions 1484–1603 at the consensus sequence). Sometimes, arrays of several hundreds of basepairs are found, composed from mainly AG-rich di- or oligonucleotides.

Characterization of Insertion Sites of HERV-W Poly(A) and Retro Copies

To provide better insight into the transposition processes of HERV-W copies, we have analyzed their insertion sites. From the retro copies, we first extracted a subset of 172 complete elements and complete soloLTRs with untruncated 5' and 3' ends. For 82 of them, we found 4-bp direct repeats in 5' and 3'-flanking sequences, just adjacent to the border of proviruses. On the other hand, in 116 of 176 elements in the poly(A) group, we detected long direct repeats of variable length up to 21 bp, with a mean of 9.51 bp.

Nucleotide frequencies in insertion sites of both groups are shown in Table 2. Three nucleotides upstream and five downstream of the start of each direct repeat were analyzed. For retro copies, there is a weak bias in the nucleotide frequencies toward a TT|ATA(A/T) sequence, in which | denotes the start of the direct repeat (Table 2a). The fifth downstream nucleotide is frequently T, already corresponding to the first nucleotide of HERV-W LTR, which is consensually T, because the general length of HERV-W insertion TSD is 4 bp.

In comparison with the retro group, poly(A) copies have a strong bias in the insertion site motif, namely toward the (A/T)(A/T)T|AAAAA octanucleotide (Table 2b).



Figure 2 The flanking sequences of poly(A) HERV-W copies. For 23 poly(A) elements shown in Fig. 1, the short flanking sequences are given in detail. Retroviral nucleotides are in upper case, flanking regions in lower case characters. Direct repeats are in italics and underlined. Poly(A) signals are highlighted by bold characters. Poly(A) tails are shown as gray boxes, satellite expansions are inside white or light gray-shaded boxes.

Table 2. Characterization of the Insertion Motif of HERV-W Copies

	(a) HERV-W retro copies								(b) HERV-W pseudogene poly(A) copies							
	1	2	3	4	5	6	7	8	1	2	3	4	5	6	7	8
All	82	82	82	82	82	82	82	82	116	116	116	116	116	116	116	116
A	<i>40</i>	10	23	34	14	41	6	2	<i>54</i>	49	30	72	73	77	71	67
C	16	27	31	20	13	1	17	9	7	11	21	9	7	6	9	10
G	15	6	26	20	8	20	30	6	11	14	16	16	15	14	19	12
T	11	39	2	8	47	20	29	65	43	41	49	19	21	19	17	27
N									1	1						
Others																

The highest nucleotide frequencies defining the consensus insertion site motif are in italics.

Phylogenetic Analysis

Two parts of the retroviral genome were selected for the phylogenetic analysis: the *gag* region (consensus sequence nucleotides 2718–3923) and a 5' part of the *pol* region (consensus sequence nucleotides 4450–6789; see Methods). Both phylogenetic trees (Fig. 3A and Supplementary Figure 2Sa, available on line at <http://genome.org>) show a complex pattern of HERV-W copies, in which poly(A) elements are dispersed within the retro copies. Due to a high similarity at non-CpG sites, there are few polymorphic sites, and thus the reliability of the obtained topology is weak (see low bootstrap values in Fig. 3A, Supplementary Figure 2Sa). We have selected only a small subset of elements, and the obtained tree (Fig. 3B) shows a better supported branching pattern, in which, again, poly(A) elements are dispersed within retroviral copies.

Genomic Distribution

The genomic distribution of HERV-W elements was calculated for 100-kb nonoverlapping segments of the genome (see Methods). Figure 4A shows the HERV-W frequency in genomic segments classified according to four ranges of the GC content. Both retro and poly(A) copies show distribution biased toward the GC-poor part of the genome. Especially, soloLTRs have a strong tendency to localize in GC-poor regions.

The intrachromosomal distribution of HERV-W (Fig. 4B) shows several interesting features. First, the intrachromosomal distribution varies among all poly(A), retro, and soloLTR groups. Poly(A) elements are over-represented on chromosomes 3, 6, 19, and X and less abundant on chromosomes 16, 17, 21, 22, and Y. Retro copies are more frequent on chromosomes 4, 5, 7, 13, X, and especially chromosome Y, where both internal sequence-containing copies and soloLTRs are clearly over-represented. On the other hand, retroviral copies are under-represented on smaller chromosomes 16, 17, 19, 20, and 22. Generally, HERV-W elements show biased distribution toward chromosomes 3, 4, X, and particularly Y, and are less frequent on chromosomes 16, 17, 20, and 22, in agreement with experimental results (Voisset et al. 2000).

Comparison of the Length and the GC Level of HERV-W Elements

We selected 46 complete poly(A) copies and 29 complete retro copies for comparison of the length and the GC level. From all copies, only positions 256–9732 of the consensus, corre-

sponding to the full-length poly(A) sequence without the poly(A) tail, were compared. To eliminate the influence of various insertions and satellite expansions, only sequences homologous to the consensus were used; sequences were aligned to the consensus and nonhomologous parts were neglected as in Figure 1. No striking differences were found in the GC content, poly(A) sequences being slightly GC poorer than the retro copies (45.8% compared with 47.7%). The length was calculated as a percentage of gap positions in all available positions of the consensus ($100 \times \text{gaps length}/\text{total length}$). These numbers were used to compare splicing variation. Because the majority of internal deletions correspond to several splice variants (visible from the Supplementary Figure 1S, available on line at <http://www.genome.org>), these numbers estimate the percentage of sequences missing due to the splicing. For the retro group, we obtained, on average, 41.4% of gaps in comparison with the consensus, for the poly(A) group, 37.6% of gaps.

DISCUSSION

Processed Pseudogenes of HERV-Ws Display the Features of LINE-Mediated Transposition

Our analysis of HERV-W copies in the human genome revealed unusual endogenous retroviral structures. We found elements colinear with retroviral mRNAs followed by poly(A) tails, the poly(A) copies, resembling processed pseudogenes of cellular genes (Vanin 1985; Weiner et al. 1986). On the basis of the lack of complete LTRs, which regenerate during normal retroviral reverse transcription, we suggest that these poly(A) copies arose by virtue of heterologous reverse transcriptase, most probably through the LINE-mediated reverse transcription.

The majority of poly(A) copies is surrounded by direct repeats of variable length (6–21 bp, mean 9.51 bp), characteristic for insertions of LINES, Alus, and processed pseudogenes, which are believed to have arisen through double enzymatic nicking of host DNA by the LINE endonuclease (Feng et al. 1996; Jurka 1997; Cost and Boeke 1998). In contrast, standard members of the HERV-W family, the retro copies, have canonically short 4-bp direct repeats as already described (Repbase Update; Blond et al. 1999). Analysis of insertion sites of poly(A) (Table 2) elements showed a strong preference for (A/T)(A/T)T|AAAAA sequences, strongly resembling the preferential TT|AAAA cutting motif of the LINE endonuclease found in LINES, Alus, and processed pseudogenes (Feng et al.

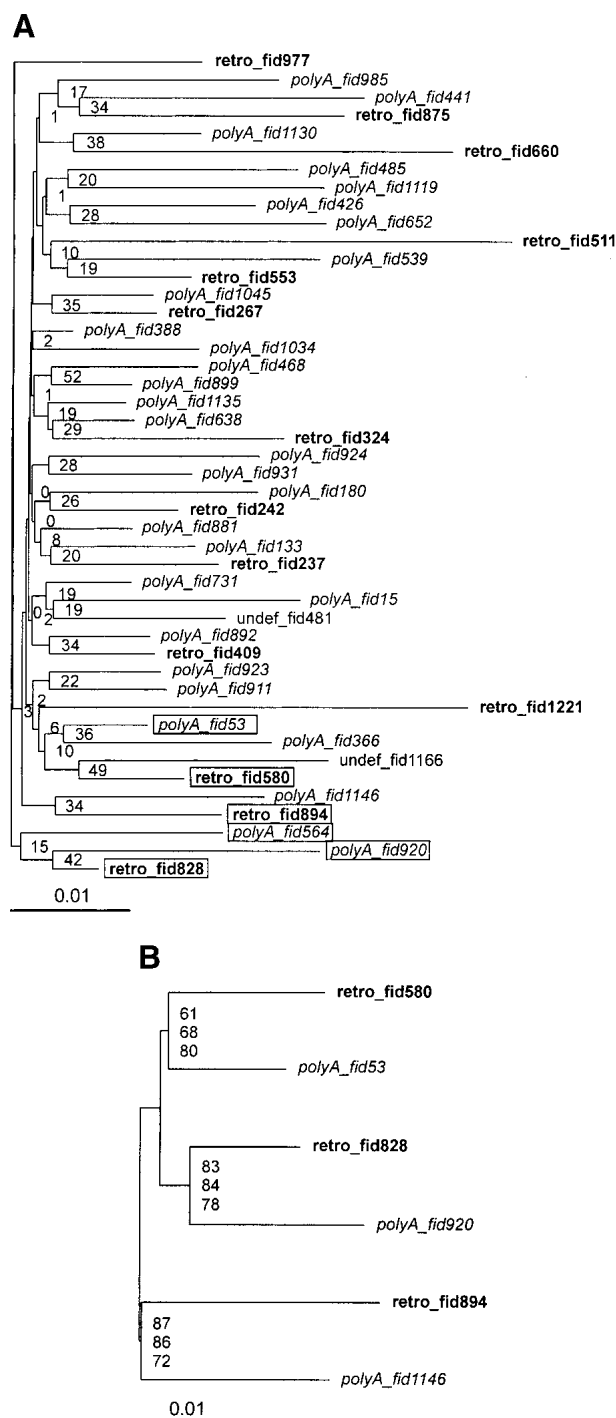


Figure 3 Phylogenetic trees of HERV-W elements. We selected multiple alignment of the proximal part of the *pol* region (nucleotides 4450–6789). All gap- and CpG-containing sites were excluded (see Methods). Poly(A) elements are in italics, retroviral copies are in bold to highlight dispersion of poly(A) elements within the retro group. (A) Neighbor-joining tree with 1000 bootstrap replicates. Forty-six elements at 581 non-gap, non-CpG sites. Rectangles show the elements selected for further phylogenetic analysis in Fig. 4B. (B) A subset of 6 elements at 1290 non-CpG, non-gap sites. Topology is based on a maximum likelihood tree, bootstrap values (1000 replicates) ordered from *top* to *bottom* show support of the topology for maximum likelihood, maximum parsimony, and neighbor-joining methods implemented in the *phylo_win* program (see Methods).

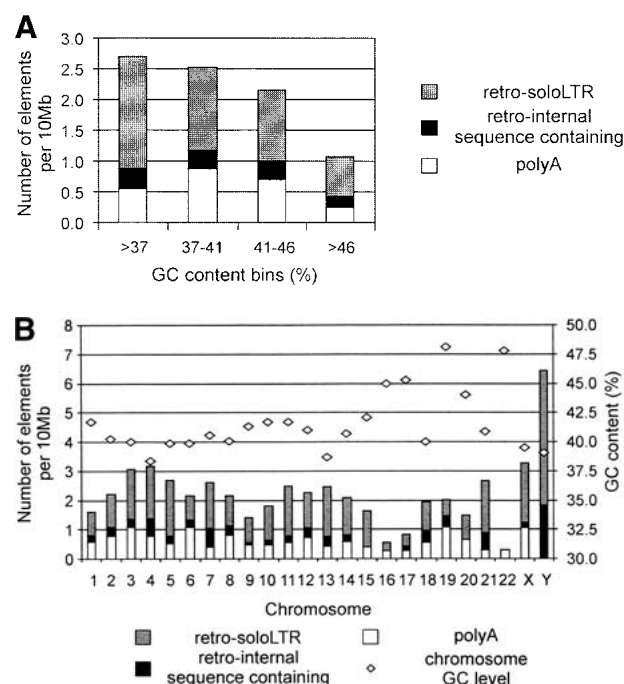


Figure 4 Genomic distribution of HERV-W elements. (A) Isochore distribution of poly(A), soloLTR, and other retroviral (internal sequence-containing) copies was calculated in 100-kb long, nonoverlapping segments. The distribution is shown in the number of independent elements per 10 Mb. Four categories of GC levels corresponding to four isochore families were used (see Methods). (B) The chromosome distribution is shown as a number of independent elements per 10 Mb (*left y axis*). The bars represent chromosomal densities of poly(A), soloLTR, as well as internal sequence-containing retro copies. The GC levels of chromosomes are shown as diamonds (*right y axis*).

1996; Jurka 1997; Cost and Boeke 1998; Wei et al. 2001). Poly(A) tails of poly(A) elements often contain microsatellites. Similar expansions were described for Alus, in which oligo-dA-rich tails have been shown to serve as nuclei for the genesis of simple repeats (Arcot et al. 1995).

Frequent 5' truncations in the poly(A) group (Fig. 1; Table 1) also described in LINES and processed pseudogenes (Voliva et al. 1983; Vanin 1985; Weiner et al. 1986; Smit 1999) the presence of spliced variants, the mRNA-derived structure, the poly(A) tail, and the insertion-site characteristics mentioned above, which altogether indicate that the LINE enzymatic machinery generated the pseudogene copies of the HERV-W family [poly(A) group]. The recent experimental demonstration of processed pseudogenes generated by LINE-encoded enzymes (Esnault et al. 2000; Wei et al. 2001) strengthens our conclusion that the poly(A) group is generated by nonretroviral, LINE-mediated mobilization. The complex pattern on phylogenetic trees, in which poly(A) copies are dispersed within retroviral copies (Fig. 3A,B), suggests multiple and independent origins of poly(A) copies.

Distribution and Stability of HERV-W Copies Depends on the Mechanism of Transposition

The genome-wide distribution of HERV-W poly(A) copies shows a bias toward the GC-poorer part of the genome than that of retro copies (Fig. 4A). This bias is typical for LINES,

young Alus, and processed pseudogenes, and is probably linked to the AT-rich insertion motif of these elements (International Human Genome Sequencing Consortium 2001; Pavlíček et al. 2001). Similarly, for the retro group, we also found a weak bias toward AT-rich insertion sites, and the genomic distribution is also biased toward the GC-poor isochores.

In contrast to LINES, there is no clear general insertion motif for retroviruses. Both retrovirus and LINE integration are influenced by the DNA bending (Pryciak and Varmus 1992; Muller and Varmus 1994; Jurka et al. 1998), the nucleosome structure at the site of integration (Pryciak and Varmus 1992; Cost et al. 2001), and the state of chromatin condensation (Leib-Mösch et al. 1993, Rynditch et al. 1998). The only sequence-specific insertion motif reported so far is the palindromic pentanucleotide GT(A/T)AC found in nonautonomous LTR retrotransposons MaLR (mammalian apparent LTR retrotransposon) (Smit 1996b) and HIV-1 integration (Stevens and Griffith 1996; Carreau et al. 1998). From the data in Table 2, we derived a weak preferential motif ATC|ATA(G/T) and a strong negative motif notTnotGnotCnotA. The described preferential motif is similar, but not the same as the CA(A/T)TG pentanucleotide, the complementary version of the MaLR and HIV-1 insertion motif. The studied MaLR and HIV-1 insertion sequences were different in the length of the TSD – 5 bp instead of 4 bp for HERV-W integration, suggesting that on the contrary to the high evolutionary conservation of the integrase region in retroviruses, insertions could differ in both TSD length and in the insertion motif. Despite the fact that the insertion motif obviously is not the only factor determining retrotransposon integration, the AT-rich LINE insertion motif seems to be responsible for the GC-poor bias in the processed pseudogene distribution.

The length of homology at 5' and 3' ends of poly(A) and retroviral elements is the determining factor for the element stability. The majority of retroviral sequences in the human genome are soloLTRs (Jurka 1998), which arose by homologous recombination between 5' and 3' LTR with excision of the central part and one LTR, leaving only a single LTR (Mager and Goodchild 1989). In our data set, most retro copies are soloLTRs (343 of 420 elements, Table 1). On the other hand, in the poly(A) group, only two elements begin close to the 5' start of the 3' LTR and there is no abundance of elements starting near the 5' end of the 3' LTR (Supplementary Figure 1S and Supplementary Table 1S, available on line at <http://www.genome.org>). Thus, apparently, there are no, or very few soloLTRs originated by homologous recombination in the poly(A) group. This observation is consistent with the requirement for the minimum length of homology needed for efficient intra- and extrachromosomal recombination

between closely linked homologous sequences, which ranges from 163 to 295 bp (Rubnitz and Subramani 1984; Liskay et al. 1987; Waldman and Liskay 1988). Retro copies have two complete LTRs, providing them with 780 bp of perfect homology at both ends, which is sufficient for effective homologous recombination. Poly(A) copies, due to the absence of retroviral reverse transcription, lack complete LTRs, and 5' and 3' ends share just the R region, yielding 71 bp of the homology at both ends, clearly below the limit for efficient homologous recombination. In accordance with their different propensity to recombination, both groups differ in their pattern of chromosomal distribution (Fig. 4B). HERV-W retro copies are over-represented on chromosomes 3, 4, X, and particularly Y in comparison with the rest of the genome. On the contrary, poly(A) copies show no bias toward Y. Because of the limited recombination, chromosome Y is known to possess a high concentration of HERVs (Kjellman et al. 1995). The increased density of LTR-retrotransposons on chromosome X, and even higher on Y, is mostly due to the larger ratio of complete elements over solitary LTRs (Smit 1999). Retro copies with internal sequences represent 28.6% (4/14) of all HERV-W retroviral copies on chromosome Y, whereas solely 18% (73/406) for the rest of the genome. The limited meiotic recombination seems to be the causative factor for both concentration of retro copies and increased proportion of retro copies containing the internal sequences on chromosome Y. Noteworthy, however, is the fact that the frequent presence of soloLTRs on chromosome Y implicates a possible role of some intrachromosomal (mitotic) recombination mechanisms such

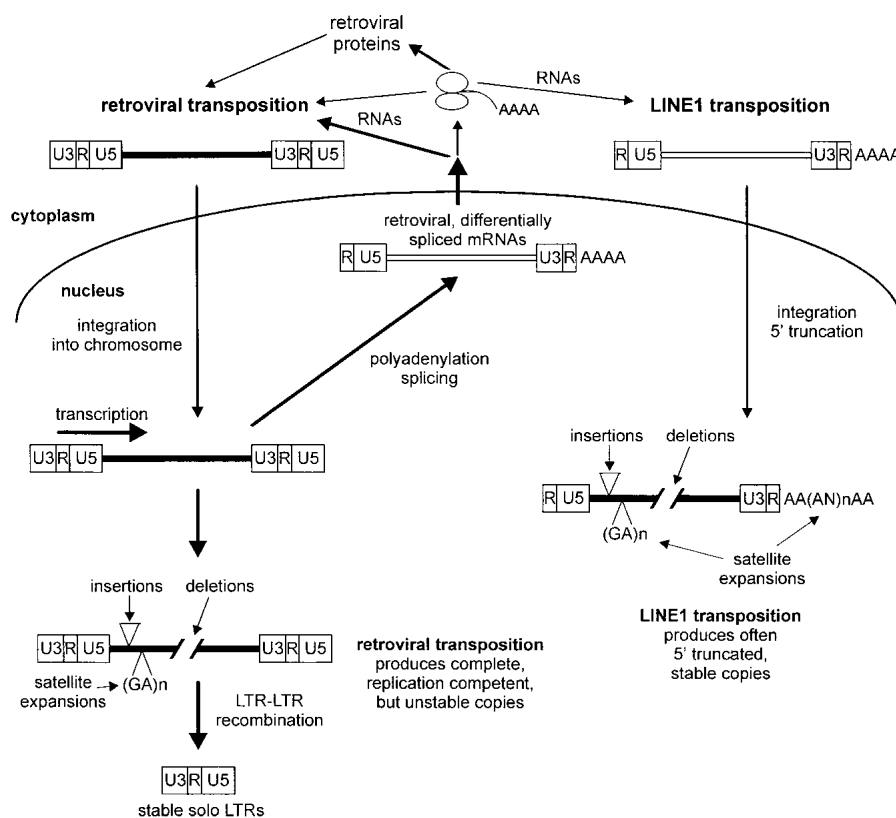


Figure 5 Schematic representation of amplification of poly(A) and retro copies, differences, and stability in the genome.

as intrachromatid single-strand annealing, reciprocal exchange, replication slippage, or abortive reciprocal recombination (for review, see Klein 1995; Lambert et al. 1999).

In conclusion, we could see that the retroviral and the LINE transposition differ in the integration, distribution and the stability of inserted copies (Fig. 5). The retroviral reverse transcription produces potentially expression- and replication-competent copies, which are unstable due to the homologous LTR–LTR recombination, and are frequently excluded during evolution. On the other hand, LINEs often produce 5' truncated copies, unable to transcribe and further mobilize due to the loss of promoter sequences in the U3 region of the 5' LTR. These copies have, however, a better chance to keep their original structure during evolution. Retro copies represent 64.2% (420/654) of all HERV-W elements, whereas poly(A) copies are just 26.9% (176/654). On the other hand, poly(A) copies account for 52.5% (149/284) of elements containing internal sequences, whereas the proportion of retro copies is only 27.11% (77/284). This implies that the majority of HERV-W *gag*, *pol*, and *env*-coding regions experimentally detected on human chromosomes (Voisset et al. 2000) are actually pseudogenes, probably nontranscribed promoterless copies, and the current search for possible causative agents of several human diseases (Perron et al. 1989, 1997; Gaudin et al. 1997, 2000; Komurian-Pradel et al. 1999; Kim and Crow 1999) could concentrate on a small subset of transcriptionally competent HERV-W elements.

Remarks on the Processed Pseudogene Formation

The LINE machinery is highly effective in the transposition of its own copies (Dombroski et al. 1991; Moran et al. 1996). The estimated frequency of cellular mRNA mobilization is only 0.01%–0.05% per LINE transposition (Wei et al. 2001). We found numerous processed pseudogenes within the HERV-W family, but not in the closely related HERV9 family. In the HERV database (Pačes et al. 2002), we have made a preliminary analysis on the basis of the presence of poly(A) tails and TSDs and we estimate that the HERV9 family has more than seven times less pseudogenes than the HERV-W family. In the class 1 HERVs, we have not found any retroviral family with >5% of processed pseudogenes (data not shown). In a direct in vitro test, the frequency of Moloney murine leukemia virus-based vector transposition was just 10^{-8} – 10^{-6} per cell per generation (Tchénio et al. 1993). Similarly, genomic library screening for the HERV-H family indicated very few (<1%) copies with the structure of processed pseudogenes (Goodchild et al. 1995). Hence, LINE-mediated transposition of the retroviral mRNA seems to be rather rare, with the exception of HERV-W mRNA.

What drives the selectivity of capturing the non-LINE mRNA by LINE-encoded enzymes? Only transpositions in the germ line, primordial germ line, or early embryonal cells have a chance to proceed into the next generation and eventually to be fixed. Both LINEs and Alus are known to be expressed in testicular tissues and they could be potentially transposed into germ-line cells (Branciforte and Martin 1994; Schmid 1998). It is noteworthy that expression of transcripts homologous to the syncytin gene have been detected not only in placenta, but also in testis (Mi et al. 2000); it is therefore conceivable that coexpression of LINEs and HERV-W elements facilitates the generation of processed pseudogenes in the HERV-W family. On the other hand, transcription up-regulation in testis is not limited to LINEs, Alus, and HERV-

Ws and could serve as an engine for generation of processed pseudogenes from various mRNAs (Schmidt 1996).

Virtually all types of mRNA are capable of retrotransposition (Brosius 1999), but the most efficient are genes connected with the translation machinery and ribosomes (Venter et al. 2001). A genome-wide analysis of processed pseudogenes suggested that the processed pseudogenes are preferentially generated from short, GC-poor RNAs (Goncalves et al. 2000). The complex splice pattern of the HERV-W family provides necessary GC and length variation and a good opportunity to test these predictions and compare the affinity of LINE and retroviral enzymes for various mRNAs. In our analysis, we have not found any difference between the retroviral and LINE replication. Also, in the analysis of the human genome sequence, no bias has been found in the GC content of genes generating processed pseudogenes (Venter et al. 2001).

METHODS

Identification and Localization of HERV-W Sequences in the Human Genome

The RepeatMasker program (Smit and Green RepeatMasker at <http://repeatmasker.genome.washington.edu>) was used to identify HERVs in the GoldenPath assembly of 87% of the human genome (Haussler et al. Human Genome Working Draft at <http://genome.ucsc.edu/>). The fragments of LTRs, as well as internal retroviral sequences of the HERV-W (HERV17) family and the related HERV9 family, were included in the following analysis. All selected retroviral fragments were compared against family profiles of HERV-W and HERV9 families using the HMMER package (HMMER v. 2.1.1; Eddy 1998; <http://hmmer.wustl.edu/>) to eliminate errors due to misidentifications of closely related families. Only elements with better matches to the HERV-W profile than to that of the HERV9 profile were used further as real HERV-W elements. Precise positions of starts and ends of elements were determined by recursive alignment with the Repbase Update consensus of the HERV family using the Dialign2 program (Dialign v. 2.0, Morgenstern et al. 1996).

Analysis of the Genomic Structure

All elements were aligned using the Dialign2 program and the multiple alignment was manually checked for errors in the Seaview editor (Galtier et al. 1996). Computing of a large alignment was approximated by aligning each element with a consensus sequence obtained from the Repbase Update, followed by assembly of pairwise alignments into multiple alignments. As a consensus, we used sequences of LTR17 and HERV17 consensus, joined in the order LTR17, HERV17, LTR17 into one sequence. In 3' LTR, we have introduced a 50-bp insertion of a poly(A) stretch in the place where the poly(A) tail of retroviral RNAs is located (from position 327 of the LTR17 consensus).

Characterization of Insertion Sites of HERV-W Copies

We have identified insertion sites of all copies, including the TSD and the insertion motif. First, the sequences were scanned for an exact match of 6 bp or longer at both ends of HERV-W elements in a region from 30 bp outside to 5 bp inside. Complete retroviral copies as well as complete soloLTRs were scanned for direct repeats of 4 bp or longer at both ends of the element.

Phylogenetic Analysis

For the phylogenetic analysis, we have selected two regions in the alignment, the *gag* region from 2718 to 3923 bp and a proximal part of the *pol* region, until the major *env* splicing

variant (nucleotides 4450–6789). Then, we excluded all incomplete and many gap-containing elements. We obtained 36 elements in the *gag* region and 46 sequences in the *pol* alignment. All CpG and gap-containing positions were excluded. The topology was obtained by neighbor joining (Tajima-Nei distance), maximum parsimony, and maximum likelihood methods implemented in the *phylo_win* program (Galtier et al. 1996).

Genomic Distribution of HERV-W Copies

Retrotransposon densities were calculated in 100-kb long, nonoverlapping fragments from the GoldenPath assembly (<http://genome.ucsc.edu>). The densities were calculated for the isochore families as given by Zoubak et al. (1996). Isochore family intervals were as follows: <37%, 37%–41%, 41%–46%, and >46% content of GC for L1, L2, H1, and H2+H3 families, respectively.

ACKNOWLEDGMENTS

We thank Oliver Clay, Jan Svoboda, and Giorgio Bernardi for critical reading of the manuscript. This work was supported by grant No. 204/01/0632 of the Grant Agency of the Czech Republic to J.H. A.P. is supported in part by a PhD fellowship of the French Government program Doctorat en cotutelle.

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Received September 28, 2001; accepted in revised form December 20, 2001.