HERV-F, a new group of human endogenous retrovirus sequences

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Using primers from a conserved region of the XA34 human endogenous retrovirus (HERV) family, four pol fragments originating from new members of the family were amplified from human genomic DNA. Southern blot analysis demonstrated similar hybridization patterns in human, chimpanzee and orangutan and distinct hybridization to macaque DNA. The probes also exhibited weaker hybridization to squirrel monkey DNA. Using large genomic clones, two full-length XA34-related HERVs have been identified. One of the HERVs is located downstream of a human Krüppel-related zinc finger protein gene, ZNF195. Both of the newly identified long terminal repeats have potential TATA boxes, poly(A) signals and transcription factor-binding sites but they differ to a high degree, especially in the U3 region. The primer-binding sites were found to be homologous to tRNAPhe (TTC), and therefore these new HERVs have been given the name HERV-F. The closest relatives to the HERV-Fs are the RTVLH-RGH family. Phylogenetic analyses of the Gag, Pol and Env regions are discussed. Both of the newly identified HERV-Fs were shown to contain protease, reverse transcriptase, integrase and env regions and had characteristic deletions in the integrase and env regions. In addition, the capsid protein gene of gag and two conserved zinc-binding motifs that are characteristic of a potential nucleic acid-binding protein were also identified. Apart from an ORF spanning the protease of one HERV-F, no other longer ORFs were found.

Introduction

The human genome, in common with all mammalian genomes, carries a high load of inheritable retrovirus sequences. About 0.1–1% of the human genome consists of human endogenous retroviruses (HERVs) (Lower et al., 1996; Patience et al., 1997), and an even larger portion of the genome consists of reverse-transcribed and transposed sequences (Smit, 1996). HERVs can be divided into different families based on their sequence similarities. Class I families, e.g. HERV-E (4-1) (Repaske et al., 1983, 1985), HERV-R (ERV-3) (O'Connell et al., 1984), HERV-I (RTVL-Ia) (Maeda, 1985) and HERV-H (RTLV-H2) (Mager & Freeman, 1987), are similar to mammalian type C retroviruses, whereas class II families, e.g. HERV-K (C4) (Dangel et al., 1994) and HTDV/HERV-K (HERV-K10) (Ono et al., 1986), share similarities with mammalian type B and D retroviruses.

The copy number of the HERV families varies from single copies to up to 10^4 copies per genome. These different copy numbers could either represent multiple integration events of the same or a closely related virus, or proviruses amplified after the original integration by, for example, retrotransposition or chromosomal rearrangements. There are indications of a negative correlation between the presence of the env gene in the provirus and the copy number, such that proviruses without env have a more retrotransposon-like structure and thereby are more amplified (Lower et al., 1996). Even though the copy number of complete proviruses is sometimes rather limited, solitary long terminal repeats (LTRs) are more frequent. This excision of protein-coding regions from HERVs might be the result of it being harmful for an organism to carry and express functional HERVs.

The majority of the HERVs that are fixed in the population are ancient from an evolutionary point of view (at least 10–100 million years old). All known HERVs are defective and cannot produce functional infectious retrovirus particles. However, all
HERVs are not transcriptionally silent and different proviruses with ORFs have been reported (Lower et al., 1995; O’Connell et al., 1984). Moreover, retrovirus particles of HERV origin have also been reported (Lower et al., 1993).

Using low-stringency hybridization with an ERV-9 env probe, a new HERV was identified from a glioma cDNA library (Widegren et al., 1996). All of the endogenous retrovirus (ERV) cDNA clones isolated were polyadenylated, the longest clone (Widegren probe, a new HERV was identified from a glioma cDNA library (only a short fragment of a

The reaction mixture (20 µl) contained 1 µg of each of primers 5′GGAATTCACTATGATCTGG 3′ and 2793 5′TGTTGA-GAGCTAGACTGCTGC 3′, 100 ng total DNA, 2.5 mM MgCl2, 2 µM dNTPs, 0.5 µM α-32P-labelled dATP, standard buffer and 1 U Ampli-Taq polymerase (Perkin-Elmer Cetus). The PCR products were separated on an 8% wedge-shaped denaturing sequencing gel and detected by autoradiography. To recover the DNA from the dried sequencing gel, a gel slice containing the DNA fragment was excised and rehydrated in TE buffer. The sample was boiled for 10 min and used for re-amplification with the same primers under the PCR conditions described above. The amplification efficiency was increased by using 20 µM instead of 2 µM dNTPs and the re-amplified fragments were cloned into the pT7blue vector (Novagen) and sequenced.

**Computer analysis.** The Wisconsin package version 9.1-unix (Genetics Computer Group, Madison, WI, USA) (Devereux et al., 1984) was used for DNA sequence analysis. The database used for NetBlast homology searches was a continuously updated version of the GenBank nucleotide database. Alignments were made primarily with the PIEPUL and LINEUP programs of the GCG package. Phylogeny was analysed with a UNIX version of PAUP (Swofford, 1992). Trees were constructed with a heuristic tree-search and bootstrap analysis for 1000 replications with the parsimony optimality criterion by using PAUP and the trees were displayed with DRAWGRAM of the PHYLIP software package (Felsenstein, 1993). Percentage identity was determined by using FASTA of the GCG package. The TFSEARCH and MOTIF programs were used with a threshold score of 85° to search the on-line TRANSFAC database (Heinemeyer et al., 1998) for potential transcription-factor-binding sites.

**Southern blot analysis.** The pol fragments of 155–156 bp from XA39, XA40, XA41 and XA42 were used for Southern blot analysis. These pol probes were labelled with [α-32P]dCTP by means of PCR from cloned material. The labelling reactions (20 µl) contained 3 ng template, 10 ng primer 593, 50 µM each of dATP, dTTP and dGTP, 15 µM [α-32P]dCTP, 2.5 mM MgCl2, PCR buffer (Perkin-Elmer Cetus) and 1 U Ampli-Taq polymerase (Perkin-Elmer Cetus). The reactions were run for six cycles of 94 °C for 60 s, 54 °C for 60 s and 72 °C for 4 min.

Total genomic DNA from human, chimpanzee (*Pan troglodytes*), orangutan (*Pongo pygmaeus*), squirrel monkey (*Saimiri sciureus*), macaque (*Macaca fascicularis*) and rat (*Rattus norvegicus*) was digested with *Pst*I and separated on a 1% agarose gel and the resultant DNA was vacuum-blotted to a Biodyne B membrane (PALL). Hybridization was carried out overnight in Rapid Hybridization solution (Amersham) at 65 °C. The filter was washed in several steps with decreasing concentrations of SSC/SDS and an increasing temperature. The final washing conditions were 1× SSC–0.5% SDS at 65 °C for 90 min.

**Results**

**Isolation and characterization of XA34-related elements**

In a previous study, we reported sequence information from five HERVs isolated either as cDNA clones (XA34), genomic clones (XA38) or by PCR (XA35–XA37) (Widegren et al., 1996). Using this information, PCR primers were constructed from a conserved portion of the RT region. Amplification was performed under low-stringency conditions from human genomic DNA and the PCR products were separated by PAGE into two size groups of approximately 153–158 and 140–146 bp (Fig. 1). DNA from two gel fragments (A and B; Fig. 1) was recovered by PCR and cloned and 25 individual clones were sequenced and analysed. Nine different XA34-related clones were identified (Table 1) and four of these clones, derived from the longer fragment (B; Fig. 1), are previously unidentified XA34-related elements. These four elements were named XA39, XA40, XA41 and XA42.

Phylogenetic distance
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Fig. 1. Denaturing sequencing gel of PCR products from human total DNA with primers preferentially amplifying a pol fragment from HERVs related to XA34. The PCR primers were 100% identical to XA34. The PCR products were labelled with 35S-dATP during the amplification (50 °C, 2·5 mM MgCl2). The regions corresponding to A and B were cut from the gel and the DNA was re-amplified, cloned and sequenced. On separation under denaturing conditions, each homogenous PCR product appeared as four bands, representing the two separated DNA strands, with or without the extra base added by Taq polymerase. A sequencing reaction was used as a molecular mass marker.

Table 1. PCR products related to XA34

<table>
<thead>
<tr>
<th>HERV</th>
<th>Region*</th>
<th>Size (bp)</th>
<th>Identity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>XA34</td>
<td>A</td>
<td>142</td>
<td>81</td>
</tr>
<tr>
<td>XA35</td>
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<td>XA38</td>
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<td>92</td>
</tr>
<tr>
<td>HERV-Fa†</td>
<td>–</td>
<td>(155)</td>
<td>88</td>
</tr>
<tr>
<td>HERV-Fb†</td>
<td>–</td>
<td>(155)</td>
<td>91</td>
</tr>
</tbody>
</table>

* The region in Fig. 1 from which the element was isolated.
† The HERV-Fa and HERV-Fb sequences are from the genomic clones.

Table 1. PCR products related to XA34

PCR products were amplified by using primers from the conserved RT region of XA34. Sizes of the different amplicons include the primer sequence. Percentage nucleotide identity, excluding primer sequences, to the consensus sequence is shown.

Analysis of genomic clones

The sequencing of the human genome is continuously adding the sequences of large genomic clones to the databases. XA34-related DNA sequences were used to screen databases for similar elements to this family. A NetBlast search revealed the existence of two large genomic clones that contained sequences with similarities to the XA34 pol region. Since we originally reported the XA34 family, we believe that it is our responsibility to characterize these more complete proviruses further.

The first of the two XA34-related elements identified is contained within a 177 kb genomic clone (accession number Z95126; bases 44530–51154) isolated from human chromosome Xq21.1–Xq21.3. The newly identified HERVs, XA42, contains the most similar pol region, exhibiting 90% identity over the region analysed. When the genomic clone was mapped by using PsI, the predicted fragment corresponding to the pol probe shown in Fig. 2 was 2224 bp in length. As this virus is located on the X chromosome, it would be expected that this probe would result in a stronger band when hybridizing to female rather than male DNA. Such a band can be observed at 2-2 kb, as shown in Fig. 2. When referring to base positions in this HERV, base 44530 in the genomic clone (Z95126) is denoted as base 1.

The second genomic clone (accession number AC000378), of 133 kb, was isolated from human chromosome 11p15.5. This clone contains an XA34-like HERV situated between bases 89430 and 95587, with a pol sequence that is 100% identical to XA41 over the region analysed. Base 89430 in clone AC000378 is denoted as base 1 in this HERV. On mapping the genomic clone with PsI, the predicted fragment corresponding to the pol probe shown in Fig. 2 was found to be 3485 bp in length. A human Krüppel-related zinc finger gene, ZNF195 (accession number AF003540; Hussey et al., 1997), was found to be located upstream of the HERV; the longest ZNF195 expressed sequence tag (EST) identified from GenBank (accession number AA505095) was found to terminate 243 bp upstream of this HERV element with a poly(A) tail. ESTs terminating within the HERV sequence, and thereby representing possible read-through transcripts from ZNF195, have also been identified (C. Kjellman & B. Widegren, unpublished results).

A schematic alignment of the two XA34-like HERVs, XA34

monkey, macaque (male only) and rat (male only) demonstrated that XA39–XA42 all exhibited similar patterns of hybridization to those of the previously identified XA34-related elements (Widegren et al., 1996). The hybridization patterns of XA39 and XA42 are shown in Fig. 2 (a, b). The probes detected similar patterns in human, chimpanzee and orangutan. There was also hybridization to macaque DNA and a rather weak but distinct hybridization to DNA from squirrel monkey, a New World monkey (Fig. 2 b).

A schematic alignment of the two XA34-like HERVs, XA34
and XA38 is shown in Fig. 3. Further analyses of different regions in the two XA34-related HERVs are described below. Within the genomic clone Z95126, it was possible to identify two similar (89% identical) 450 bp potential LTR regions separated by 5.7 kb and organized as direct repeats (positions 1–453 and 6167–6614). Similarly, in the second genomic clone, AC000378, it was also possible to identify two direct repeats (435 bp) sharing 90% identity and separated by 5.3 kb (positions 1–436 and 5709–6146). In clone Z95126, a direct repeat of 5 bp was located directly upstream of the potential 5' LTR and downstream of the potential 3' LTR, thus delineating the boundaries of the HERV (Fig. 4). The alignment of these LTR regions in Fig. 4 demonstrates the loss of the dinucleotides AA at the 5' end and TT at the 3' end from the HERVs as a result of processing during the integration event (Temin, 1981). There is also a short (5 bp) inverted repeat starting with the nucleotides TG at the ends of the potential LTRs, as underlined in Fig. 4. Because of the low sequence similarity to other characterized LTRs, it is not feasible to use homology alone to define different regions of the potential LTRs. However, it is possible to identify the TATA box (at base 340 in Fig. 4), the poly(A) signal (at base 410 in Fig. 4) and clustered potential sites for binding of transcription factors such as Sp1, GATA, AML-1a, STATx, CREB, cEBP, AP-4 and AP-1 (data not shown) in the LTR-like regions. The junction between the R and U₅ sequences is located at the polyadenylation site 16–25 bp downstream of the poly(A) signal and the U₅–R junction is found at the CAP site 20–30 bp downstream of the TATA box (Guntaka, 1993). The potential U₅ regions from these two HERVs share very little sequence similarity, whereas the potential R and U₅ regions exhibit a high degree of identity (75%) (Fig. 4).

The primer-binding site has previously been used in the classification of endogenous retroviruses (Larsson et al., 1989). We have been able to identify a region located just downstream of the 5' LTR in both Z95126-HERV and AC000378-HERV...
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Fig. 3. Schematic alignment of HERV-Fa, HERV-Fb, XA34 & XA38. The LTRs, CA, NC, PR, RT, IN, SU and TM regions are indicated. Angled lines indicate regions presumably deleted from the HERVs as compared with e.g. the RTVLH-RGH family. XA34 was isolated as a cDNA and not a genomic sequence [indicated in the figure by the poly(A) tail]. The ORF of the PR gene in HERV-Fb is indicated. The regions used for phylogenetic analyses are indicated by lines marked I–IV above the figure. Dotted lines indicate that the elements are probably longer; however these regions are not yet cloned.

that is similar to the 18 bp region at the 3’ end of a human tRNA\textsuperscript{Phe} (TTC) (accession number K00350). It was found that the Z95126-HERV shares 100% identity to tRNA\textsuperscript{Phe} (TTC), whilst AC000378-HERV deviated by only one mismatch in the 18 bp region (Fig. 4). By applying the taxonomic nomenclature of Larsson et al. (1989), we suggest that the Z95126-HERV and the AC000378-HERV are named HERV-Fa and HERV-Fb, respectively.

Downstream from the 5’ LTRs, both HERV-Fa and HERV-Fb have potential leader/gag regions (HERV-Fa, bases 460–2408; HERV-Fb, bases 434–2435) of 2 kb. HERV-Fa and HERV-Fb share similarity (70% nucleotide identity) over the complete gag region. However, at the nucleotide level, it is very difficult to align accurately the first 1·5 kb after the LTR of HERV-Fa or HERV-Fb with the gag regions of known retroviruses or retrovirus sequences. The final 0·5 kb of the gag region is more similar (55% nucleotide identity) to RTVLH-RGH2. However, the deduced amino acid sequences of the complete capsid (CA) region (region I in Fig. 3) of HERV-Fa (bases 1440–2199) and HERV-Fb (bases 1464–2220) were aligned over 267 amino acids with the corresponding regions of known HERVs. Only a single maximum-parsimony tree was reconstructed (Fig. 5a) following 1000 bootstrap replications using the heuristic tree search. This tree groups the two HERV-Fs together and demonstrates that the HERV-H1 family, as represented by RTVLH-H2 and RTVLH-RGH2, is the most closely related ERV family. Over the CA region, the amino acid identity between HERV-Fa and HERV-Fb is 63% and it is 35% between RTVLH-H2 and HERV-Fa. Within the CA region in both HERV-Fa and HERV-Fb, a conserved major homology region was identified with 50% (HERV-Fa) and 35% (HERV-Fb) amino acid identity to the MuLV major homology region (Strambio-de-Castillia & Hunter, 1992; Zlotnick et al., 1998).

Immediately after the CA region, both HERV-Fa and HERV-Fb contain lysine-rich basic sequences with two conserved zinc-binding motifs (CX\textsubscript{2}CX\textsubscript{2}H\textsubscript{2}C). These are characteristic of potential nucleic acid-binding protein (NC) regions. The coding potential of the HERV-Fa and HERV-Fb gag regions is disrupted by stop codons and introduced frame-shift mutations (Fig. 6).

The genes of the pol region encoding protease (PR), RT and IN of HERV-Fa and HERV-Fb were identified by aligning the pol regions to RTVL-H2 and RTVLH-RGH2. The PR gene (bases 2409–2739) and the RT gene (bases 2740–4117) of HERV-Fa have an approximately 780 bp deletion that has removed the C-terminal portion of PR and the N-terminal portion of RT. In contrast, HERV-Fb contains the complete PR (bases 2436–2820) and RT genes (bases 2821–4802). HERV-Fb has an ORF encoding a complete PR gene product, which terminates at the beginning of RT (Figs 4 and 6). The maximum-parsimony tree (Fig. 5b) reconstructed from the deduced and aligned amino acids of the PR region (region II in Fig. 3) demonstrates that HERV-Fb should be assigned to the same branch as, but separate from, RTVLH-RGH2 and RTVL-H2. Over the PR region, the amino acid identity between HERV-Fb and RTVLH-RGH2 was 39%.

HERV-Fa and HERV-Fb share 80% sequence identity (at the DNA level) in the RT gene; this represents the most conserved genetic region within the retroviruses. A maximum-parsimony tree (Fig. 5c) was reconstructed after a heuristic tree search and 1000 bootstrap replications with the aligned sequences of 651 amino acids spanning part of the RT region.
and part of the IN region (region III in Fig. 3), and this tree clearly groups HERV-Fa, HERV-Fb and XA34 together and separates them from the RTVLH-RGH family. Over ~205 amino acids in the C terminus of the conserved RT, there is approximately 38–42% amino acid identity between the HERV-Fs and XA34 and RTVLH-RGH1 (Table 2). The amino acid identity between the HERV-Fs, HUMER4-I, ERV9, HERV-W and RTVLH-RGH1 is summarized in Table 2. HERV-Fa, HERV-Fb, XA34 and XA38 all carry large deletions that result in somewhat truncated C-terminal portions of the IN gene and a truncation (HERV-Fa) or deletion (HERV-Fb, XA34 and XA38) of the surface protein (SLU) gene in the env region. However, given that the truncations observed in the four different HERVs are not in exactly the same position within the IN gene (300 bp variation) (Fig. 3), these are probably due to independent events. HERV-Fa has the deletion at position 5034 and HERV-Fb has the deletion at position 5150. There are no ORFs that have the potential to encode functional proteins within the RT or truncated IN regions (Fig. 6).

The env region of HERV-Fa (bases 5034–6018) contains a large C-terminal portion of the SU gene and a complete TM gene, as identified by alignment to RTVLH-RGH2. In HERV-Fb, the SU gene and the N-terminal portion of the TM gene are deleted. The sequences of HERV-Fa and HERV-Fb are similar (about 60% nucleotide identity) along the entire length of the 3’ end of the virus until the beginning of the 3’ LTR, with the similarity to RTVLH-RGH2 ending just downstream of the TM gene. Phylogenetic analysis was performed on the conserved TM region (region IV, Fig. 3) spanning 121 amino acids that also includes the CKS-17 motif, which has been reported to have immunosuppressive activity (Cianciolo et al., 1985). Only a single maximum-parsimony phylogenetic tree was reconstructed (Fig. 5d) following bootstrap analysis (1000 replications) with a heuristic search. The analysis groups HERV-Fa, HERV-Fb and XA34 together whilst separating them from the RTVLH-RGHs, ERV-9 and HERV-W. The env region of XA34 that was used for the phylogenetic analyses includes a region downstream of Alu that was obtained from a genomic clone (accession number AC005281) (C. Kjellman & B. Widegren, unpublished results). The env regions of the HERV-Fs do not have the potential to encode functional peptides. The sequence AGGAGGTTTGAA is found 163 bp downstream of the env region and immediately upstream of the 3′ LTR in HERV-Fa. This is most likely to be a conserved potential polypurine tract. A potential polypurine tract with the sequence AAAAGGCTAAAA is also found in HERV-Fb, and there is no ORF that has the potential to encode functional peptides in the SU gene and the N-terminal portion of the TM gene (Fig. 6).

The Southern blot analysis indicates the presence of HERV-F-related elements in the genome of a New World primate (squirrel monkey) (Fig. 2b). One means of identifying the time at which the ERV was integrated is to analyse the two LTRs of a specific provirus and compare the divergence between these two LTRs and relate it to the divergence of other cellular genes from different species (Dangel et al., 1995). In order to be able to do this, one has to assume that the LTRs of a particular ERV have evolved independently since the time of integration. One also has to assume that the LTRs were identical when the retrovirus was originally integrated. Analysis of the LTRs of HERV-Fa shows that there are 36 base pair exchanges in 2 × 442 bp and 40 base pair exchanges in 2 × 433 bp in HERV-Fb. This gives an approximate divergence of 0.041 base pair exchanges per position for HERV-Fa and 0.046 base pair exchanges per position for HERV-Fb. If the same calculations are made for the LTRs of HERV-K (C4) (Dangel et al., 1994, 1995), a divergence of 0.039 base pair exchanges per position...
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Fig. 5. Phylogenetic analyses of the aligned deduced amino acid sequences of CA of Gag (a), PR (b), the partial RT/IN region of Pol (c) and the partial TM region of Env (d) from the HERV-Fs and different retrovirus sequences. These regions are indicated as I–IV in Fig. 4. The sequences were aligned using PILEUP and LINEUP and the phylogenetic analyses were made using PAUP. Only single maximum-parsimony trees were reconstructed by using the heuristic tree-search for 1000 bootstrap replications. The bootstrap values are indicated on the trees, which were displayed using DRAWGRAM.

Fig. 6. Start (above the line) and stop (below the line) codons in frames 1–3 (indicated by numbers in parentheses) of the HERV-F sequences illustrate the disrupted reading frames of HERV-Fa and HERV-Fb. The analysis was made by using FRAMES (GCG).
Table 2. Percentage amino acid identity of a conserved ~205 amino acid region of RT

<table>
<thead>
<tr>
<th>HUMER-4-1</th>
<th>HERV-W</th>
<th>ERV9</th>
<th>RGH1</th>
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<th>HERV-Fb</th>
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(42 bp exchanges in 2 × 538 bp) is obtained. The time for the integration of HERV-K has been estimated to be just after the split between the New World and Old World primates, as based on the divergence of the short intron 9 of the C4 gene (Dangel et al., 1995). The divergence of this marker between human and macaque (Macaca mulatta) is approximately 0.029 base pair exchanges per position (24 bp exchanges in 2 × 412 bp) and that between human and cotton top tamarin (Saguinus oedipus), a New World monkey, is approximately 0.053 base pair exchanges per position (43 bp exchanges in 2 × 409 bp). If this approximation was widened to include the HERV-Fs, it would indicate that HERV-K (C4) and the HERV-Fs were integrated at approximately the same time, i.e. after the split of the New and Old World monkeys. However, it should be stressed that there are no guarantees of equal mutation rates in different HERV integrations.

Discussion

We have isolated nine separate pol sequences demonstrating homology to class I HERVs and have suggested previously that these HERVs should be regarded as a separate family of retroviruses (Widegren et al., 1996). Two complete proviruses have now been characterized that demonstrate strong sequence similarity to XA34. On the basis of their primer-binding sites, they have been designated HERV-Fa and HERV-Fb. The related elements XA34–XA42 share strong similarity to these HERV-Fs, but the primer-binding sites of these elements have not yet been identified. However, on the basis of sequence homology, we suggest that XA34–XA42 should also be classified in this HERV-F family of endogenous retroviruses.

The LTRs of the HERV-Fs were identified and found to share no sequence similarity to LTRs of other known HERVs. The LTRs of HERV-Fa differ strongly from HERV-Fb within the 5′ region (potential U3) but are more similar within the 3′ region (R and U6). A high degree of variation in the U3 region, particularly in the enhancer sequences, has been demonstrated in other retrovirus families (Majors, 1990). Given that the LTRs of HERV-Fa and HERV-Fb share very little sequence similarity in U3, these two proviruses are likely to be the result of two separate germline infections of distinct but related retroviruses. This is also supported by the fact that there is no sequence similarity between the flanking regions of the two HERV-Fs, clearly demonstrating that these proviruses are not the result of a gene amplification.

The HERV-F gag genes differ strongly from other known gag genes; however, the potential CA gene and conserved zinc-binding motifs of a lysine-rich potential NC gene were identified in both HERV-Fs. These gene products usually represent the most conserved proteins of the gag region (McClure et al., 1988). It was found that the HERV-Fs and the related XA34 and XA38 all have deletions within the C-terminal portion of the IN gene and the N-terminal portion of the SU gene. The fact that these deletions have not taken place at exactly the same location in each gene indicates that these deletions have occurred as separate events that for some reason have been directed towards this particular region. One can speculate that these deletions took place before the integration into the germline, thus facilitating the retroviruses in becoming endogenous. It is also possible that the deletions took place after integration, thereby silencing both the IN and the env genes. The truncation of the env gene gives the HERV a more retrotranspon-like structure. Except for an ORF spanning the PR region of HERV-Fb, the HERV-Fs were not found to contain any longer ORFs within the gag, pol or env regions.

HERV-Fa and HERV-Fb can be grouped together by phylogenetic analysis of their capsid, protease, polymerase and transmembrane protein regions. These analyses and the rather low degree of similarity in RT demonstrate that they are clearly separated from other known HERVs, the HERV-H family being their closest relative. The phylogeny of the different regions follows a similar pattern, in that neither of the two HERV-Fs shows any sign of recombination with other HERV families. We have discussed previously a possible recombination between XA34 and ERV-9, since the region upstream of the XA34 Alu is rather similar to the ERV-9 env, but here we also included the env region located downstream of the Alu in the analyses and we cannot conclude that such a recombination has occurred.

Southern blot hybridization using pol probes from different HERV-F-related proviruses demonstrates the presence of approximately 16 HERV-F family members in the human genome. Database analysis of genomic sequences also revealed a number of elements that contain just HERV-F-like LTRs and gag regions (e.g. accessions Z86001, Z83745 and AC000216). Therefore, it is probably correct to assume that the HERV-F family, in common with many other ERV families, contains a large number of truncated HERV remnants dispersed over the genome. The 16 members identified by Southern blot hybridization define the number of pol-containing HERV-F-related elements in the genome.

Southern blot hybridization of XA34-related elements demonstrated the presence of these HERVs in all Old World
primates analysed. However, it was also possible to detect HERV-F bands in New World primates, e.g. XA42 (Fig. 2b), therefore suggesting that the first integration took place more than 60 million years ago (Arnason et al., 1996). Of course, it is also possible that the hybridization to the New World primate DNA reflects cross-hybridization to the more conserved pol regions of unrelated proviruses. However, the absence of signal from the hybridization to rat DNA, which carries many C-type ERVs, is an indication of a rather high specificity of the hybridization. The specificity is also supported by the fact that cross-hybridization to the HERV-H family, which is the most closely related family, with approximately 1000 members (Mager & Henthorn, 1984), would be expected to give a much larger number of bands than the relatively small number detected with the XA39 and XA42 probes. We also tried to relate the divergence between the 5’ and 3’ LTRs of the HERV-F family with the divergence of the HERV-K (C4) LTRs and that of the short intron 9 of the C4 gene (Dangel et al., 1995). The time of integration for HERV-K was estimated to be just after the split between the New World and Old World primates (Dangel et al., 1995). Our analyses indicated that the HERV-F LTRs and HERV-K (C4) LTRs have the same degree of divergence, indicating that these HERVs were integrated at about the same time. This calculation contradicts the interpretation of the Southern blot analysis, as this indicates that the HERV-Fs were integrated before the split between the New World and Old World primates. However, it should be remembered that there is an inevitable degree of uncertainty when calculating rates of divergence, particularly over such short genetic regions, and that there are no guarantees of equal mutational rates in the different HERV elements. There is of course also the possibility of cross-hybridization between conserved pol regions of different retrovirus origin in Southern blot analysis. In order to be able to determine with absolute certainty whether the HERV-F family is present in New World primates, the potential elements should be cloned and sequenced.

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References


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