**REVIEW**

Demystified . . . Human endogenous retroviruses


Human endogenous retroviruses (HERVs) are a family of viruses within our genome with similarities to present day exogenous retroviruses. HERVs have been inherited by successive generations and it is possible that some have conferred biological benefits. However, several HERVs have been implicated in certain cancers and autoimmune diseases. This article demystifies these retroviruses by providing an insight into HERVs, their means of classification, and a synopsis of HERVs implicated in cancer and autoimmunity. Furthermore, the biological roles of HERVs are explored.

Human endogenous retroviruses (HERV's) represent footprints of previous retroviral infection and have been termed “fossil viruses”. They are transmitted vertically through the germline and are thus inherited by successive generations in a Mendelian manner. Over time, HERVs have been subjected to repeated amplification and transposition events giving rise to multi-copy and single copy proviruses that are distributed within the DNA of all cells. Overall, HERVs constitute about 1% of the human genome. HERVs possess a similar genomic organisation to present day exogenous retroviruses such as human immunodeficiency virus (HIV) and human T cell leukaemia virus (HTLV), and are composed of gag, pol, and env regions sandwiched between two long terminal repeats (LTRs) (fig 1). The LTRs possess nucleotide sequence motifs that are fundamental to the regulation of retroviral gene expression. In brief, the gag and env genes encode retroviral capsid and envelope proteins, respectively, whereas the pol gene encodes enzymes for viral replication, integration, and protein cleavage. Retroviruses in effect are retrograde, because the flow of genetic information is reversed compared with the normal pathway of molecular biosynthesis—DNA → RNA → protein. Indeed, all retroviruses necessitate the conversion of viral RNA into a cDNA intermediary, which is catalysed by the enzyme reverse transcriptase.

“Overall, human endogenous retroviruses constitute about 1% of the human genome”

Over 20 HERV families have been identified during the past two decades. Although many are defective through the accumulation of mutations, deletions, and termination signals within coding sequences, a limited number of HERVs have the potential to produce viral products and, indeed, to produce viral-like particles. Furthermore, some HERVs have been implicated in certain autoimmune diseases and cancers and might have a role in the aetiology and pathology of disease. However, many HERVs have been present in our genome for a considerable period of time so that their presence may also be of benefit to the human host.

**MOLECULAR BIOLOGY AND HERVS**

The focus of the human genome project is to sequence the 3 billion DNA bases that compose our human genome. Interestingly, only a small proportion (about 3%) appears to constitute the estimated 24 179 to 87 720 genes that are translated into the proteins necessary to life. The function of non-coding DNA (maligned as “junk DNA”) is not readily obvious and it may have important roles in packaging and influencing gene expression. DNA is composed of two helical strands orientated in an antiparallel fashion (fig 2), where each strand consists of a deoxyribose phosphate backbone and a series of purine and pyrimidine bases denoted A (adenine), G (guanine), T (thymine), and C (cytosine). A unit of sugar, phosphate, and base is strictly termed a nucleotide, although casually represented by its base. The two DNA strands associate through the complementary pairing of bases—for example, A–T and C–G—which is stabilised through hydrogen bonding. Stretches of DNA, measured in base pairs, constitute the numerous genes that are found on the 23 pairs of human chromosomes. Genes vary in size, with some in the order of 20–40 kilobase pairs (kbp), whereas others extend to millions of base pairs. In addition, human genes are composed of exons, which are transcribed and translated into amino acids, and introns, which are interspersed between exons and represent non-translated regions that contribute to the large size of some genes. Retroviruses are remarkably small (in the order of 9–10 kbp) compared with other pathogenic viruses, such as cytomegalovirus, which is 200 kbp.

**Abbreviations:** CFTR, cystic fibrosis gene; CTL, cytotoxic T lymphocyte; EBV, Epstein-Barr virus; HERV, human endogenous retrovirus; HIV, human immunodeficiency virus; HTLV, human T cell leukaemia virus; IDDM, insulin dependent diabetes mellitus; ISP, immunosuppressive peptide; LINES, long stretches of related sequences; LTR, long terminal repeat; MHC, major histocompatibility complex; MMTV, mouse mammary tumour virus; PCR, polymerase chain reaction; PTN, pleiotrophin gene; RA, rheumatoid arthritis; SLE, systemic lupus erythematosus; SS, Sjogren’s syndrome; TGGT, testicular germ cell tumour
Nevertheless, retroviruses capitalise on their limited size by using frame shifts to retrieve genetic information from multiple overlapping open reading frames (fig 3). Interestingly, the open reading frames of many defunct HERVs are interrupted by premature stop codons that interrupt the “read through” of genetic information. A previous demystified article highlights the use of the polymerase chain reaction (PCR) to detect certain nucleotide regions of HERVs that are common to all retroviruses.

“Retroviruses capitalise on their limited size by using frame shifts to retrieve genetic information from multiple overlapping open reading frames”

Before protein synthesis, mRNA is transcribed from relevant stretches of DNA (reliant on a triplet base code or codon) that specifies a start signal (for example, ATG), an amino acid, or a termination signal (TAA, TAG, or TGA). The primary mRNA transcript is refined by splicing out unwanted introns and then by adding a series of adenine bases to its 3′ end. The process of polyadenylation is considered crucial to the stability of the mRNA that leaves the nucleus to attach to cytoplasmic ribosomes. Here, individual tRNA molecules attached to a specific amino acid are aligned with mRNA that dictates the assembly of polypeptide chains (fig 4). Strictly speaking, tRNAs use anticodons for complimentary binding, although a codon, such as AAU for lysine (denoted by K in the single amino acid code), is often used with reference to the tRNA “primer binding site” of a HERV—for example, HERV-K. Interestingly, tRNAs may use the “wobble” phenomenon in that the first two bases are essential for binding but the third base is less crucial. Therefore, it is possible that tRNAs may over-ride stop codons and allow the growth of a polypeptide chain. Ultimately, both exogenous and endogenous retroviruses exploit the cell’s genetic machinery and comply with the same generic principles of protein biosynthesis to produce further viral progeny. However HERVs stop short of viral budding (fig 3) and consequently are non-infectious.

CLASSIFICATION OF HERVS

Historically, retroviruses were shown to cause malignant tumours and leukemias in chickens and mammals—for example, the Rous sarcoma virus and mouse mammary tumour virus (MMTV), respectively—and were subdivided into three subfamilies: oncovirinae (oncoviruses), lentivirinae (slow viruses), and spumavirinae (foamy viruses). Retroviruses were also classified according to morphological and biological criteria: in particular by the observance of retroviral particles in infected cells. In brief, type A retroviruses were only visible inside cells, devoid of envelope, and referred to as intracisternal particles (A type particles), whereas types B, C, and D were enveloped and produced extracellular particles that varied in size and appearance. Subsequently, in the 1980s, the retroviruses HTLV-I and HIV-1 were reported and a new classification formulated by the International Committee for Taxonomy of Viruses. This taxonomy reflected the similarity of a virus to an established retroviral genus, such as murine leukemia related virus. Set against this background, the first HERV was reported in 1981, and many others have subsequently been identified (table 1) using several different methodologies. These have included the screening of human...
Human endogenous retroviruses

Molecular biosynthesis: human endogenous retroviruses

![Diagram of molecular biosynthesis]

- S-LTR
- GAG
- POL
- ENV
- LTR-3'

Open reading frames of HERV-K: transcript generated from “frame shifts of sequence” information (denoted at Frames 1, 2, 3)

LTR control: e.g., hormone responsive element

Potential translation of 3 reading frames

- UUCUCCCAAAAAGGACGGAUGAGCCGUUUA

Figure 3

Diagram illustrating the influence of the long terminal repeat (LTR) on the production of endogenous viral peptides. Of particular importance is that a human endogenous retrovirus (HERV) product may be generated using different open reading frames.

Figure 4

A schematic diagram illustrating that a premature stop codon on mRNA may be over-ridden at the ribosome, thus allowing a potential to produce full length or truncated viral products.

Molecular biosynthesis

- UCAAGG
- LYS
- ASP
- GLU

Potential translation of 3 reading frames

- PHE-PRO-LYS-LYS-ASP-GLU
- STOP
- ALA-VAL-TYR

- PHE-PRO-LYS-ARG-THR-ASP-GLU
- PRO-PHE

- SER-GLN-LYS-GLY-ARG-MET-SER-ARG-LEU

Hermans and cancer

Although the precise role(s) of HERVs in the carcinogenic process has not been fully elucidated there are several studies that, if taken together, put forward a convincing argument for the possible involvement of HERVs in malignancy. HERVs may be involved in carcinogenesis by virtue of the expression of HERV mRNA, functional proteins, or retroviral-like particles. They may also be associated with the generation of new promoters or the activation of proto-oncogenes. The expression of HERV-R mRNA is increased in some cases of small cell lung carcinoma. In addition, a teratocarcinoma cell line has been shown to possess a HERV-K sequence and to secrete retroviral-like particles. Testicular germ cell tumours (TGCTs) have been shown to contain proteins of the HERV-K family and patients with TGCT often exhibit a specific immune response to gag and env proteins. It has been suggested that HERV-K may be important in the progression of TGCT through inhibition of an effective immune response, and the HERV env genes have been shown to encode immunosuppressive proteins. It is clear that overexpressed HERV proteins can elicit high titre IgG responses in some settings (for example, HERV-K10 in patients with renal cancer).
reverse transcriptase activity.

Conversely, activation of proto-oncogenes by which proto-oncogenes become activated, leading to subsequent neoplastic transformation. In particular, activation of proto-oncogenes of the ras family is common in many tumour types, and some studies have suggested a potential role for HERV's in ras activation. It was shown a methylnitrosourea induced rat mammary carcinoma cell line produces retroviral particles into the intron of c-Ha-ras was responsible for its more than 10 fold overexpression.51

Recently, it has been shown that lymphotropic herpesvirus, Epstein-Barr virus (EBV), itself a potent transforming agent, can transcriptionally activate the env gene of HERV-K18, which possesses superantigen activity (as demonstrated by major histocompatibility complex (MHC) class II dependent preferential activation of TCRVB13 T cells in response to murine B cells transfected with the HERV-K18 env gene).52

The authors suggested that this phenomenon accounts for the previously described EBV associated superantigen activity, which might in turn be crucial to T cell activation by EBV. Although the exact function of HERVs in the carcinogenic process is still under investigation, the evidence implicating HERVs in the carcinogenic process is substantial and further investigation will be required to elucidate the contribution of HERVs to the development of malignancy.

HERVS AND AUTOIMMUNITY

In 1990, an article appeared in the Times newspaper (24 November) with the title “AIDS-like virus may cause arthritis”. The report focused on Robert Garry’s research that identified retroviral particles in lip biopsies taken from patients with primary Sjogren’s syndrome (SS).53 Similarly, in other autoimmune rheumatic diseases, such as rheumatoid arthritis (RA) and systemic lupus erythematosus (SLE), a plethora of articles added to this intriguing observation by providing evidence of retroviral antigens at the site of disease, or the presence of antiretroviral antibodies in the sera of patients.54–60 A novel report in 1994 used both PCR (using consensus primers) and serological tests to investigate the presence of retroviruses in a cross section of patients with rheumatoid diseases, including RA, SS, and SLE.54–60 Interestingly, PCR failed to amplify products relating to HTLV-I or HIV-1, although antibodies to retroviral antigens were detected in the sera of patients. Consequently, there appeared to be a conundrum: antibodies to retroviral particles were present but no evidence to implicate exogenous retroviruses could be found. Between 1996 and 1999, some research groups used so called “degenerate” retroviral primers in their PCR reactions.54–60 These primers cater for modest variations within two segments common to all retroviruses within the reverse transcriptase encoding pol region and provide an intervening “fingerprint region”, which permits DNA sequencing. In brief, these studies,54–60 revealed nucleotide homologies to endogenous retroviral families, including viruses with similarity to known exogenous retroviruses. Thus, it was plausible that the presence of HERVs could provide an explanation for the presence of antiretroviral antibodies in certain rheumatoid diseases.54–60 HERVs have also been implicated in other autoimmune diseases, such as multiple sclerosis (HERV-W, HERV-H) and insulin dependent diabetes mellitus (IDDM) (HERV-K, IDDM22), in addition to inflammatory vascular diseases.62–66 However, in the case of IDDM, subsequent studies55–56 have not been able to confirm this association. Mechanisms whereby HERVs could influence autoimmunity include molecular mimicry (HERV’s sharing amino acids common to host proteins), superantigen motifs that bypass the normal MHC restrictive process of T cell stimulation, aberrant expression of antigens, and the presence normally expressed only at very low amounts in a few normal adult tissues) appears to be responsible for the aggressive and invasive growth of human choriocarcinoma.64

Overexpression is a common mechanism by which proto-oncogenes become activated, leading to subsequent neoplastic transformation. In particular, activation of proto-oncogenes of the ras family is common in many tumour types, and some studies have suggested a potential role for HERV's in ras activation. It was shown a methylnitrosourea induced rat mammary carcinoma cell line produces retroviral particles into the intron of c-Ha-ras was responsible for its more than 10 fold overexpression.51

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Table 1 Classification of human endogenous retroviruses (HERVs)

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<thead>
<tr>
<th>HERV family</th>
<th>Representative accession number</th>
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Class II HERVs (type A, B, and D related HERVs)

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as detected by the SEREX method (serological identification of expressed genes),53 suggesting that HERV proteins may in the future provide targets for antiviral immunotherapy.

“It has been suggested that HERV-K may be important in the progression of testicular germ cell tumours through inhibition of an effective immune response”.

HERV-K might be important in the pathogenesis of human breast cancer. It has been shown that the T47D human mammary carcinoma cell line produces retroviral particles54 with reverse transcriptase activity.55 Both the HERV-K10 related sequences of T47D cells56 and the reverse transcriptase activity57 are increased by steroid hormone treatment, which is thought to be the result of transcriptional activation via binding of the progesterone receptor to regions on the HERV-K genome that correspond to progesterone and glucocorticoid response elements.

In choriocarcinoma, it has been shown that a HERV type C is inserted into the human growth factor gene, pleiotrophin (PTN). This results in the generation of a novel tissue specific promoter, which results in the expression of HERV–PTN fusion transcripts, leading to the production of biologically active PTN protein. Expression of the PTN protein (which is
of neo-antigens, perhaps as a result of HERV and/or exogenous viral combinations. The use of animal models has also served to enhance our understanding of endogenous retroviruses. In a lupus model, an 8.4 kbp endogenous retroviral transcript is expressed in affected mice. Furthermore, a retroviral element in one of the introns of the fas apoptosis gene appears to alter the splicing of fas transcripts, resulting in a lupus-like autoimmune disease in MRL-lpr/lpr mice.

Further investigations using animal models and multicentre patient studies are needed to establish links between specific HERVs and autoimmune diseases because many HERVs are also expressed in varying amounts, or in a coordinated fashion, in normal tissues.

**Immunosuppressive peptide**
The product derived from the env gene of mammalian type C retroviruses possesses motifs—for example, the fusion peptide, leucine zipper protein, and immunosuppressive peptide (ISP)—that are essential for fusion and the infection of cells. In brief, the precursor env product is cleaved into two components: a surface protein (gp70) and a transmembrane protein (p15E) that contains an immunosuppressive region. Of interest is an ISP, termed CKS-17, that suppresses lymphocytes and alters cytokine profiles in animal models. The ISP sequence from the murine leukaemia virus LQNRRGLD-LLFLKEGGGC (single amino acid code) is reasonably well conserved in HERV-H19 (LQNRRGLDLTAEGGLGLC), HERV-R (ERV3) (YQNRLALDYLLAQEEGVC), and HERV-E(4–1) (YQNRLALDYLLAAEGGVC), but less so in HERV-K10 (FEASKAHNLNPGTEAIA). The presence of an ISP could be advantageous to a virally infected cell—in terms of shielding or “cloaking” itself from immunological attack—but may equally be important to a host. This is perhaps exemplified by HERV-R (ERV3), which is highly expressed in trophoblastic cells and results in high concentrations of env protein (~65 kDa) in syncytiotrophoblasts. The immunosuppressive potential of this HERV and the fusogenic nature of placenta tissue suggests a possible involvement in normal placental function, in protecting the developing fetus from maternal immunological attack.

**BIOLOGICAL IMPORTANCE OF HERVS**
Phylogenetic studies have shown that some HERVs emerged over 3 million years ago, whereas others appeared after the divergence of hominoid and ape lineages. Consequently, HERVs have been present in our genome for a considerable period of time and perhaps have been retained because they performed a useful biological function. Alternatively, some HERVs may have been difficult to eliminate and thus persisted during evolution. To confer a selective advantage, a premise remains that HERVs produce products and/or augment mechanisms that benefit host survival. Several possibilities are highlighted below.

**Figure 5** Expression of endogenous viruses HERV-K and RTVL-H in human B cells as demonstrated by reverse transcriptase polymerase chain reaction.
immune responses. Furthermore, it is possible that HERVs may change the pattern of gene expression during embryo development by altering different rates of development of different parts of the embryo. Another endogenous retrovirus, HERV-W, has also been shown to encode a protein termed syncytin, which may have a role in placental morphogenesis.

Antiviral resistance
Mechanisms of conferring protection against a related viral agent may include retroviral receptor blockade (by HERV products) and interference of replication by antisense mRNA. However, it is also possible that HERV peptides could prime an immune response to an undesirable agent. Intriguingly, cohorts of female sex workers in the Gambia and Kenya have been identified who remain uninfected and seronegative despite being repetitively exposed to HIV. The apparent “resistance” to HIV infection is suggested to result from MHC class I restricted cytotoxic T lymphocytes (CTLs)—HLA-B35 and HLA-A2 (A*6802)/HLA-B18—which are found in Gambians and Kenyans, respectively. The immune response to HIV infection is characterised by a vigorous HIV specific CTL response where virus-specific CTLs recognise antigen in the form of processed peptides (eight to 10 amino acids in length), which are bound in the cleft of MHC class I molecules on the surface of antigen presenting cells. In this context, recent data have highlighted similarities between HIV CTL peptide sequences and regions of HERV-K10. Consequently, in some cases previous exposure to HERV peptides could potentially immunise certain individuals, although this argument does not explain why some females seroconvert after a reduction in sex work.

Long terminal repeats
Retroviral genes that have been integrated into the genome are bordered by short direct repeats of host DNA and LTR sequences of about 500–600 nucleotides. These LTRs can influence neighbouring genes because they may contain transcriptional regulatory elements such as enhancers, promoters, hormone responsive elements, and polyadenylation signals. The influence on protein expression is demonstrated by proteins such as salivary amylase, ZNF80, cytochrome C1, Kruppel-like H-plk, and phospholipase A2-L, whose genes are linked to LTRs. Numerous solitary LTRs are also present within our genome. These sequences, which include about 10 000 copies of solitary LTRs pertaining to HERV-K, have arisen through recombination events that could have “deleted” harmful retroviral genes. It is noteworthy that LTRs are located near MHC genes and thus could modulate antigen expression, particularly in individuals who possess identical MHC genes but different LTRs. With reference to autoimmune diseases, the presence of an HERV-K LTR in the HLA-DQ region (DQ-LTR3) has been shown to have a strong influence on the occurrence of IDDM. Furthermore, the presence of DQ-LTR on HLA-DQB1*0302 and its absence on DQB1*0201 are independent risk markers for IDDM. The presence of DQ-LTR has also been shown to be a marker for increased susceptibility to RA in subjects with the HLA-DR4-DQB1*0301 and HLA-DR4-DQB1*0302 haplotypes. In addition, polymorphisms within LTRs could be important because changes in sequence could greatly affect promoter activity. Interestingly, three allelic forms of the LTR of HERV-R (ERV3) changes in sequence could greatly affect promoter activity. In addition, polymorphisms within LTRs could be important because enhancers and promoters to new sites in the genome and thus alter the expression of existing adjacent genes. Of course, there remains a potential to disrupt genes by insertion mutation and produce defective and/or truncated products, as has been highlighted by Kazazian, who reported cases of human genetic disease caused by the random insertion of LINES.

PREMATURE STOP CODONS
It is known that many HERVs possess premature stop codons and of course the RNA products of truncated genes may well be eliminated before protein translation. However, examples of premature stop codons that result in human disease by the production of a truncated protein product have been demonstrated in cases of the DNA repair disorder xeroderma pigmentosum and also in leukaemia. In addition, approximately 5% of cystic fibrosis cases are caused by premature termination codons. Interestingly, some aminoglycoside antibiotics appear to suppress stop codons in several organisms. Howard and colleagues demonstrated this effect in cell cultures that expressed constructs carrying two different premature stop codons within the cystic fibrosis gene (CFTR). Treatment with antibiotic produced a full length CFTR: the physiological suppression of the premature stop codon was thought to result from a mispairing of an amino-acyl tRNA that successfully bound to a stop codon. The importance of this observation is that HERVs, despite possessing premature stop codons, may have the potential to produce truncated or full length products that are fundamental to the mechanisms of molecular mimicry, aberrant expression of products, and the stimulation of T cells through superantigen motifs.

CONCLUSIONS
This article has provided an overview of a complex topic that may have ramifications in host protection, cancer, and autoimmunity. Ultimately, are HERVs friends or foes? In conferring a biological advantage, HERVs (and solitary LTRs) may indeed be beneficial. Their role in immunological homeostasis and perhaps protection against exogenous retroviruses is intriguing. Alternatively, HERV insertion mutation, molecular mimicry, superantigen motifs, and recombination with other viruses could be responsible for the development and pathology of disease. An additional aspect is whether the presence of HERV peptides during ontogeny culminates with a role in the immune repertoire. As a result, peptides with similarity to HIV-CTL sequences could be more dangerous to a given individual.
Human endogenous retroviruses

Take home messages

- Human endogenous retroviruses (HERVs) make up part of
  our genome and represent footprints of previous retroviral
  infection
- HERVs, although present, possess a low risk of being infectious
- The HERV-K superfamily represents one of the most active
  HERVs and is capable of producing retroviral particles
- HERVs may be beneficial to the host but could also be
  harmful, and may be involved in certain autoimmune
  diseases and cancers

"An additional aspect is whether the presence of HERV
peptides during ontogeny culminates with a hole in the
immune repertoire"

Clearly, there is a need for multicentre studies to ascertain
firm associations between HERVs and autoimmune disease
states and certain cancers. In particular "gene chip" technolo-
gies will no doubt relate HERV expression with disease and
pathological progression. Transcription of individual HERVs
or the coordinated expression of HERVs, although important,
must be balanced against expression found in normal tissues.
Consequently, studies of HERV/LTR polymorphisms, transacti-
vation by helper viruses (or other triggers), and the role of full
length or spliced transcripts may provide further knowledge of
these viruses. In addition, there is a requirement for a panel of
readily available antibody reagents (for example, monoclonal
antibodies, recombinant phage antibodies) to determine
retroviral products at the site of disease. No doubt the field of
HERV research will continue to accelerate so that we can
fully ascertain the consequences of renegade endogenous retro-
viruses and their transfer in xenotransplantation. 84–86

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REFERENCES

1 Tristem M. Identification and characterisation of novel human
endogenous retrovirus families by phylogenetic screening of the human
eds. Molecular biology of tumour viruses. 2nd ed. Cold Spring Spring
4 Boller K, König H, Sauter M, et al. Evidence that HERVK is the
endogenous retrovirus sequence that codes for the human
retroviruses with complex mRNA expression and particle formation.
6 Nelson PN. Retroviruses in rheumatic diseases. Ann Rheum Dis
7 Nelson PN, Lever AM, Smith S, et al. Molecular investigations implicate
human endogenous retroviruses as mediators of antiretroviral antibodies
8 International Human Genome Sequencing Consortium. Initial
sequencing and analysis of the human genome. Nature

9 Baumforth KRN, Nelson PN, Digby JE, et al. The polymerase chain
10 Lower R, Lower J, Kurth R. Review. The viruses in all of us: characteristics
and biological significance of human endogenous retrovirus sequences.
11 Bernhard W. Electron microscopy of tumour cells and virus tumours.
12 Dalton AJ. Further analysis of detailed structure of types B and C
13 Coffin J. Genetic diversity and evolution of retroviruses. Curr Top
14 Martin MA, Bryan T, Rasheed S, et al. Identification and cloning of
endogenous retroviral sequences present in human DNA. Proc Natl Acad
15 Larsson E, Nato N, Cohen M. Human endogenous proviruses. Curr Top
16 Urmovits HB, Murphy WH. Human endogenous retroviruses: nature,
occurance, and clinical implications in human disease. Clin Microbiol
18 Franklin GC, Christen S, Hansson IM, et al. Expression of human
retrovirus sequences related to those of mouse mammary tumor virus.
endogenous retrovirus class-like sequences. J Gen Virol
20 Medstrand P, Blomberg J. Characterization of novel reverse
transcriptases encoding human endogenous retroviruses similar to
type A and B retroviruses: differential transcription in normal human tissues.
21 Ono M, Yasunaga T, Miyata T, et al. Nucleotide sequence of human
endogenous retrovirus genome related to the mouse mammary tumor
gene intron 9 as a phylogenetic marker for primates: terminal length
repeats of the endogenous retrovirus ERVK(C4) are a molecular clock of
23 Tonjes RR, Lower R, Boller K, et al. HERVK: the biologically most active
human endogenous retrovirus family. J Acquir Immune Defic Syndr
retroviruses K (HERVK) proviruses are unique to humans. Curr Biol
HIIV-related endogenous sequences in man. Nucleic Acids Res
endogenous retrovirus ERV3 (HERV-K) mRNA in normal and neoplastic
K10: expression of gag protein and detection of antibodies in patients
identification of transcribed retrovirus sequences (KUS PCR) reveals the
expression of the human endogenous retrovirus loci HER/HERV and
29 Schulte AM, Shoupeng L, Kurtz A, et al. Human trophoblast and
choriocarcinoma expression of the growth factor pleiotrophin attributable
to germ-line insertion of an endogenous retrovirus. Proc Natl Acad Sci
U S A 1996;93:14759–64.
30 Bera TK, Tsukamoto T, Panda DK, et al. Defective retrovirus insertion
activates c-Ha-ras proto-oncogene in an MNU-induced rat mammary
31 Sauter M, Roemer K, Best B, et al. Specificity of antibodies directed
against Env protein of human endogenous retroviruses in patients
32 Larsson E, Andersson G. Beneficial role of human endogenous
33 Lindeskog M, Medstrand P, Blomberg J. Sequence variation of human
endogenous retrovirus ERV-R related elements in an env region
corresponding to an immunosuppressive peptide: transcription in normal
34 Sahin U, Torricelli A, Pfenninger M. Serological identification of human
produced by a human breast carcinoma cell line: immunological
relationship with mouse mammary tumour virus proteins. Proc Natl Acad
36 Patience C, Simpson GR, Colletta AA, et al. Human endogenous
retrovirus expression and reverse transcriptase activity in the T4D
endogenous retrovirus genome by female steroid hormones in
38 Kurts A, Schulte AM, Welte A. Pleiotrophin and multiple normal
39 Rhim JS, Yang JH, Lee IH, et al. Oncogene and transgenes correlates of
cancer and risk assessments. In: Zarzer C, editors. Oncogenes and
transactivates the human endogenous retrovirus HERV-K18 that encodes a
Di Giovine F, Murphy VJ
Wu J, Nakagawa K
Krieg AM
Katsumata K, Herrman M
Perl A, Garson JA
Antibodies and HLA class II alleles. Lupus erythematosus and overlap syndrome: correlation with antinuclear family.
Rheumatological diseases although a minority of sera cross-react with MSRV-RNA in serum of patients with multiple sclerosis.
Cell diabetes mellitus does not depend on specific expression of the human peripheral blood mononuclear cells.
Integration of an endogenous retrovirus in an apoptosis gene.
Strains.
Full-length endogenous MCF-related transcript in autoimmune mouse, Zhou T, He J, Mountz JD. Autoimmune disease in mice due to integration of an endogenous retrovirus in an apoptosis gene.
Development of insulin-dependent diabetes mellitus does not depend on specific expression of the human peripheral blood mononuclear cells.
UV light and HIV gene activation.
Expression of human endogenous retrovirus in placental trophoblasts suggests involvement in human placental morphogenesis.
Ralphs DW, Konkel ME, Sanker TI, et al. Known and novel patient sequences belonging to the HERV-W family were found in primates.