## Re-evaluating inheritance in genome evolution: widespread transfer of LINEs between species

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Transposable elements (TEs) are mobile DNA sequences, colloquially known as

'jumping genes' because of their ability to replicate to new genomic locations. Given

a vector of transfer (e.g. tick or virus), TEs can jump further: between organisms or

species in a process known as horizontal transfer (HT). Here we show that LINE-1

(L1) and Bovine-B (BovB), the two most abundant TE families in mammals, were

initially introduced as foreign DNA via ancient HT events. Using a 503-genome

dataset, we identify multiple ancient L1 HT events in plants and show that L1s

infiltrated the mammalian lineage after the monotreme-therian split. We also extend

the BovB paradigm by identifying more than twice the number of estimated transfer

events compared to previous studies, new potential blood-sucking parasite vectors

and occurrences in new lineages (e.g. bats, frog). Given that these TEs make up

nearly half of the genome sequence in today's mammals, our results provide the first

evidence that HT can have drastic and long-term effects on the new host genomes.

This revolutionizes our perception of genome evolution to consider external factors,

such as the natural introduction of foreign DNA. With the advancement of genome

sequencing technologies and bioinformatics tools, we anticipate our study to be the

first of many large-scale phylogenomic analyses exploring the role of HT in genome

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evolution.

**Keywords** 

Genome evolution, horizontal transfer, transposon, eukaryote, mammal

Significance statement

LINE-1 (L1) elements occupy about half of most mammalian genomes (1), and they are believed to be strictly vertically inherited (8). Mutagenic L1 insertions are thought to account for approximately 1 of every 1000 random, disease-causing insertions in humans (4-7). Our research indicates that the very presence of L1s in humans, and other therian mammals, is due to an ancient transfer event – which has drastic implications for our perception of genome evolution. Using *a machina* analyses over 503 genomes, we trace the origins of L1 and BovB retrotransposons across the tree of life, and provide evidence

Introduction

of their long-term impact on eukaryotic evolution.

Transposable elements (TEs) are mobile segments of DNA which occupy large portions of eukaryotic genomes, including more than half of the human genome (1). Long interspersed element (LINE) retrotransposons are TEs which move from site to site using a "copy and paste" mechanism, facilitating their amplification throughout the genome (2, 3). The insertion of retrotransposons can interrupt existing genetic structures, resulting in gene disruptions, chromosomal breaks and rearrangements, and numerous diseases such as cancer (4-7). Two of the most abundant retrotransposon families in eukaryotes are LINE-1 (L1) and Bovine-B (BovB) (8, 9).

Horizontal transfer (HT) is the transmission of genetic material by means other than parent-to-offspring: a phenomenon primarily associated with prokaryotes. However, given a vector of transfer (e.g. virus, parasite), retrotransposons have the innate ability to jump between species as they do within genomes (2, 10). Studies investigating the possibility of

HT in retrotransposons are limited, mainly including CR1s and RTEs (9, 11-13). Given

the limited evidence to date, we tested the hypothesis that horizontal transfer is a

ubiquitous process not restricted to certain species or retrotransposons. We used L1 and

BovB elements as exemplars because of their contrasting dynamics and predominance in

mammalian genomes. BovB retrotransposons provide an excellent example of horizontal

transfer: divergent species contain highly similar BovB sequences and the analysis of

various tick species reveals a plausible vector of transfer (9). In contrast, L1 elements are

believed to be only vertically inherited, based on knowledge gained primarily on

mammalian organisms (8). We hypothesise that the presence of L1s in therian mammals,

and absence in monotremes, is due to an ancient HT event. In this study, we use BovBs as

a comparison to identify common characteristics of horizontally transferred elements in

contemporary eukaryotic species.

Three criteria are typically used to detect HT candidates: 1) a patchy distribution of the TE

across the tree of life; 2) unusually high TE sequence similarity between divergent taxa;

and 3) phylogenetic inconsistencies between TE tree topology and species relationships

(14). To comprehensively test these criteria, we performed large-scale phylogenomic

analyses of over 500 eukaryotic genomes (plants and animals) using iterative similarity

searches of BovB and L1 sequences.

**Results and Discussion** 

Our findings show that there are two phases in HT: effective insertion of the TE, followed

by expansion throughout the genome. Figure 1 shows that both BovB and L1 elements

have been horizontally transferred because of their patchy distribution across eukaryotes.

Both are absent from most arthropod genomes yet appear in relatively primitive species

such as sea urchins and sea squirts. Furthermore, both TEs are present in a diverse array of species including mammals, reptiles, fish and amphibians. The main difference between BovB and L1 lies in the number of colonised species. BovBs are only present in 60 of the 503 species analysed, so it is easy to trace their horizontal transfer between the distinct clades (e.g. squamates, ruminants). In contrast, L1s encompass a total of 407 species, within plants and animals, and they are ubiquitous across the well-studied therian mammals. However they are surprisingly absent from platypus and echidna (monotremes). There are only two possible explanations for this; either L1s were expunged shortly after the monotreme-therian split but before they had a chance to accumulate, or monotremes never had L1s. The first scenario is unlikely in the context of L1 distributions in other eukaryotes. Consider the 60 currently available bird genomes: full-length L1s have all but been eradicated from the avian lineage, but every bird species bears evidence of ancient/ancestral L1 activity through the presence of fragments (15). In contrast, there are no L1 fragments in monotremes. We therefore conclude that L1s were inserted into a common ancestor of therian mammals, between 160 and 191 Million Years Ago (MYA), and have since been vertically inherited (see below).

The abundance of TEs differs greatly between species. As shown in Fig. 1, mammalian genomes are incredibly susceptible to BovB and L1 expansion. More than 15% of the cow genome is formed by these TEs (12% BovB, 3% L1). This is without considering the contribution of TE fragments (16) or derived Short Interspersed Elements (SINEs), boosting retrotransposon coverage to almost 50% in mammalian genomes (1). Even within mammals there are noticeable differences in copy number; for example, bats and equids have a very low number of full-length BovBs (<50 per genome) compared to the thousands found in ruminants and Afrotherian mammals. The low copy number here is

TE-specific rather than species-specific; there are many L1s in bats and equids. Hence, the rate of TE propagation is determined both by the genome environment (e.g. mammal

versus non-mammal) and the type of retrotransposon (e.g. BovB versus L1).

To develop a method for identifying horizontal transfer events, we used BovB, a TE known to undergo HT. We clustered and aligned BovB sequences (both full-length

nucleotide sequences and amino acid reverse-transcriptase domains) to generate a

representative consensus for each species, and infer a phylogeny (Fig 2a shows the

nucleotide-based tree). The phylogeny supports previous results (9) — with the topology

noticeably different from the tree of life (Fig. 1) — although we were able to refine our

estimates for the times of insertion. For example, the cluster of equids includes the white

rhino, Ceratotherium simum, suggesting that BovBs were introduced into the most recent

common ancestor before these species diverged. The low copy number in equids and

rhino, observed in Fig. 1, is not because of a recent insertion event. The most likely

explanation is that the donor BovB inserted into an ancestral genome, was briefly active,

lost its ability to retrotranspose and was subsequently inherited by its descendants.

The placement of arthropods is intriguing, revealing potential HT vectors and the origin of

BovB retrotransposons. For example, BovBs from butterflies, moths and ants appear as a

basal monophyletic group, sister to sea squirt Ciona savignyi BovB. The presence of

BovB in all these species suggests that BovB TEs may have arisen as a subclass of ancient

RTEs, countering the belief that they originated in squamates (13). The next grouping

consists of two scorpion species (Mesobuthus martensii and Centruroides exilicauda)

nestled among the snakes, fish, sea urchin and leech — a possible vector. But the most

interesting arthropod species is *Cimex lectularius*, the common bed bug, known to feed on

animal blood. The full-length BovB sequence from *Cimex* shares over 80% identity to viper and cobra BovBs; their reverse transcriptase domains share over 90% identity at the amino acid level. Together, the bed bug and leech support the idea (9) that blood-sucking parasites can transfer retrotransposons between the animals they feed on.

We extended the BovB paradigm to include 10 bat species and one frog (*Xenopus tropicalis*). The bats were not included in the phylogenetic analysis because their BovB sequences were too divergent to construct an accurate consensus. Instead, we clustered all individual BovB sequences to identify two distinct subfamilies (Fig. 2b); one containing all the horse and rhino BovBs as well as eight bat sequences, and the other containing the remaining bat BovBs as well as the single BovB from *Xenopus*. We also included three annotated sequences from a public database (17) to resolve an apparent discrepancy between the naming of BovB/RTE elements. Our results have several implications: first, bat BovBs can be separated into two completely distinct clades, suggesting bat BovBs arose from independent insertion events; second, the BovBa-1-EF bat clade may have arisen from an amphibian species, or vice versa; and third, the naming conventions used in RepBase (17) need updating to better distinguish BovB and RTE sequences. This third point is discussed in the Supplementary Information (see Supp. Fig. 1).

In order to exhaustively search for all cases of BovB HT, we replicated the all-against-all BLAST (18) approach used in El Baidouri *et al.* (2014) (19) to detect individual HT candidate sequences. Briefly, this compares all sequences within a database to generate BovB clusters or families. We identified 215 HT candidate families which contained BovBs belonging to at least two different eukaryotic species. Many of these were closely related species; so to find the HT families most likely to be true events we restricted the

analysis to families that linked species in different eukaryotic Orders (e.g. Afrotheria and Monotremata). We performed *a machina* validation for each candidate HT family: pairwise alignments of the flanking regions to rule out possible contamination or orthologous regions, and phylogenetic reconstructions to confirm discordant relationships. A total of 22 HT families passed all of the tests, indicating at least 22 cross-Order HT events. Many HT families included one or two reptile BovBs, and numerous mammalian BovBs (see Supp. Table 6). This is important for determining the direction of transfer. BovBs are thought to have entered ruminants after squamates (13). The single reptile element in a family is therefore likely to be the original transferred sequence, supporting the theory that retrotransposons undergo HT to escape host suppression or elimination (19). Altogether, our results demonstrate that the horizontal transfer of BovB elements is even more widespread than previously reported, providing one of the most compelling examples of eukaryotic horizontal transfer to date.

We carried out the same exhaustive search in L1s, which presented a challenge because of greater divergence and a strong vertical background. Producing a consensus for each species was impractical as most species contained a divergent mixture of old, degraded L1s and young, intact L1s. Instead, we used the all-against-all clustering strategy on the collated dataset of L1 nucleotide sequences over 3kb in length (>1 million sequences total). 2815 clusters contained L1s from at least two different species; these were our HT candidates. As with BovBs, to improve recognition of HT we looked for families displaying cross-Order transfer. Most non-mammalian L1s (insects, reptiles, amphibians) had already been excluded because they definitively grouped into species-specific clusters, even at low (50%) clustering identity. The remaining families were from plants and mammals. After the validation tests, we found that all the mammalian candidate families

were very small (e.g. one L1 element per species), and located in repeat-dense,

orthologous regions in the genome most likely explained by vertical inheritance (see

Supp. Fig. 3). Thus, we found no evidence for recent L1 transfer since their insertion into

the therian mammal lineage and subsequent shaping of modern therian genomes.

Nevertheless, four plant families presented a strong case for L1 horizontal transfer (Fig.

3a). High sequence identity was restricted to the elements themselves, there were more

than two L1 elements in each family, the sequences encoded open reading frames or had

intact reverse-transcriptase domains, and the phylogenetic reconstructions showed

evolutionary discordance. The number of elements in each family mimicked the patterns

seen with BovBs: very few elements from the 'donor species', and a noticeable expansion

of L1s in the 'host species'. This indicates that transferred L1s can retain activity and

expand within their new host. Moreover, it contradicts the belief that L1s are exclusively

vertically inherited, and supports our conclusion that a similar event introduced L1s to

mammals. At this stage, we do not know the vector of transfer since none of the analysed

arthropods showed similarity to plant L1 sequences.

During our mining of candidate L1 HT families, we serendipitously discovered a chimeric

L1-BovB element present in cattle genomes (Bos taurus and Bos indicus), shown in Fig.

3b. This particular element most likely arose from a recently active L1 element (98%

identical to the canonical Bos L1-BT (17)) inserting into an active BovB (97% identical to

Bos BovB (17)). In fact, L1s and BovBs have accumulated to such extents in these two

genomes that they have created the ideal environment for chimeric repetitive elements.

With two reverse-transcriptase domains and high similarity to currently active L1/BovB

elements, this chimeric element has the potential to still be functional - presenting the

possibility for L1 elements to be horizontally transferred throughout mammals by being

transduced by BovBs.

In conclusion, both BovB and L1 retrotransposons can undergo HT, albeit at different

rates. We extracted millions of retrotransposon sequences from a 503-genome dataset,

demonstrating the similarly patchy distributions of these two LINE classes across the

eukaryotic tree of life. We further extended the analysis of BovBs to include blood-

sucking arthropods capable of parasitising mammals and squamates, as well as two

distinct clades of bat BovBs and the first report of BovB in an amphibian. Contrary to the

belief of exclusive vertical inheritance, our results with L1s suggest multiple ancient HT

events in plants and, strikingly, HT into the early therian mammal lineage. The rapid

speciation following the split of theria and australosphenids (monotremes), between 160-

220 MYA, coincides with the invasion of L1 elements into therian genomes. We therefore

speculate that the speciation of therian mammals was driven in part by the effect of L1

retrotransposition on genome structure and function (including regulatory effects on

transcriptional networks). This ancient transfer event allowed expansion of L1s and

associated SINEs, transformation of genome structure and regulation in mammals (7) and

potentially catalysed the therian radiation.

**Materials and Methods** 

Detailed description of the methods, including tables and figures, are available in the

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Supplementary Information.

Extraction of L1 and BovB retrotransposons from genome data

To extract the retrotransposons of interest, we used the methods and genomes previously

described in Ivancevic et al. (2016) (15). Briefly, this involved downloading 499 publicly

available genomes (and acquiring four more from collaborations), then using two

independent searching strategies (LASTZ (20) and TBLASTN (18)) to identify and

characterise L1 and BovB elements. A third program, CENSOR (21), was used with the

RepBase library of known repeats (17) to verify hits with a reciprocal best-hit check. The

raw L1 results have been previously published in Ivancevic et al. (15) (Supplementary

Material); the BovB results are included in the Supplementary Information.

Extraction and clustering of conserved amino acid residues

Starting with BovBs, USEARCH (22) was used to find open readings frames (ORFs),

with function -fastx\_findorfs and parameters -aaout (for amino acid output) and -orfstyle 7

(to allow non-standard start codons). HMMer (23) was used to identify reverse

transcriptase (RT) domains within the ORFs. RT domains were extracted using the

envelope coordinates from the HMMer domain hits table (-domtblout), with minimum

length 200 amino acid residues. The BovB RT domains from all species were collated into

one file and clustered with UCLUST (22). This was done as an initial screening to detect

potential horizontal transfer candidates. The process was then repeated with L1 elements.

Clustering of nucleotide sequences to build one consensus per species

The canonical BovB retrotransposon is 3.2 kb in length (9, 17), although this varies

slightly between species. In this study, we classified BovB nucleotide sequences ≥2.4kb

and ≤4kb as full-length. We wanted to construct a BovB representative for each species.

Accordingly, for each species, UCLUST (22) was used to cluster full-length BovB

sequences at varying identities between 65-95%. A consensus sequence of each cluster

was generated using the UCLUST -consout option.

The ideal cluster identity was chosen based on the number and divergence of sequences in

a cluster. E.g. for species with few BovBs, a lower identity was allowed; whereas for

species with thousands of BovBs, a higher identity was needed to produce an alignable

cluster. The final clustering identity and cluster size for each species are given in Supp.

Table 1. Note that the bat species are not included in this table - they were clustered

separately, due to the high level of divergence between BovBs.

This method was tested on L1 retrotransposons, but the results were not ideal; most

species simply had too many L1 sequences. Other methods tested on both BovBs and L1s

included using centroids instead of consensus sequences (this gave better alignments but

was less representative of the cluster), and using the same clustering identity for all

species (e.g. 80% - this did not work well for species with less than 100 elements in the

genome).

Inferring a phylogeny from consensus sequences

Consensus sequences were aligned with MUSCLE (24). The multiple alignment was

processed with Gblocks (25) to extract conserved blocks, with default parameters except

min block size: 5, allowed gaps: all. FastTree (26) was used to infer a maximum

likelihood phylogeny using a general time reversible (GTR) model and gamma

approximation on substitution rates. Geneious Tree Builder (27) was used to infer a

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second tree using the neighbour-joining method with 1000 bootstrap replicates.

Distinguishing RTEs from BovBs

All sequences which identified as BovB or RTE were kept and labelled accordingly to

their closest RepBase classification (17). However, there appeared to be numerous

discrepancies with the naming: e.g. some RTE sequences shared >90% identity to BovBs,

and vice versa. BovB retrotransposons are a subclass of RTE, and they were only

discovered relatively recently. It is likely that several so-called RTE sequences are

actually BovBs.

To determine which species had BovB sequences, and which only had RTEs, we used the

species consensus approach to build a BovB/RTE phylogeny (see Supp. Figure 1). This

effectively separated BovB-containing species from RTE-containing species. The RTE

sequences were not included in further analyses.

Clustering of nucleotide BovB sequences from bats and *Xenopus* 

A reliable BovB consensus could not be generated for any of the ten bat species because

the sequences were too divergent and degraded. Some bat BovBs seemed similar to equid

BovBs; others did not. Likewise, the single full-length BovB from frog *Xenopus tropicalis* 

was very different to canonical BovBs, sharing highest identity with the bats.

In an effort to characterise these BovBs into families, we grouped all full-length BovB

sequences from the bats, frog, equids and white rhino into a single file. We also added two

RepBase equid sequences (RTE-1\_EC and BovB\_Ec) and 1 RepBase bat sequence

(BovBa-1 EF) (17). After clustering, we expected to find one family of equid BovBs, the

equid RTE sequence as an outlier, and numerous families containing bat and frog BovBs.

The actual findings are described in the text (Fig. 2b). We used UCLUST (22) to cluster

the sequences (function -cluster\_fast with parameters -id, -uc, -clusters). The highest

identity at which there were only 2 clusters/families was 40%. At higher identities, the

equid BovBs stayed together but the bat and frog BovBs were lost as singletons.

To confirm the clustering, we also used MUSCLE to align all the sequences and FastTree

to infer a phylogeny (see Supp. Figure 2).

HT candidate identification - BovBs and L1s

We compiled all confirmed BovB and L1 sequences into separate multi-fasta databases

(316,017 and 1,048,478 sequences respectively). The length cut-off for BovBs was ≤2.4kb

and ≥4kb; for L1s, ≥3kb. BovBs were analysed first to identify characteristics of

horizontal transfer events.

To detect HT candidates, we used the all-against-all clustering strategy described in El

Baidouri et al. (2014) (19). Briefly, this method used a nucleotide BLAST (18) to

compare every individual sequence in a database against every other sequence; hence the

term all-against-all. BLAST parameters were as follows: -r 2, -e 1e-10, -F F, -m 8 (for

tabular output). The SiLiX program (28) was then used to filter the BLAST output and

produce clusters or families that met the designated identity threshold.

For BovB sequences, we tested identities of 40-90%. High identity thresholds were useful

for finding very recent HT events (e.g. over 90% identity between the bed bug and

snakes). However, the majority of clusters contained several copies of the same BovB

family from a single species - indicative of vertical inheritance. Using a lower identity

threshold was more informative for capturing ancient HT events. At 50% identity, the

clustering preserved the recent, high-identity HT events while also finding the ancient,

lower-%identity HT events. We concluded that this was the best %identity to use for our

particular dataset, considering it includes widely divergent branches of Eukaryota.

Clusters were deemed HT candidates if they contained BovB elements belonging to at

least two different species. To reduce the number of possible HT clusters, we went one

step further and kept only the clusters which demonstrated cross-Order transfer (e.g.

BovBs from Monotremata and Afrotheria in the same cluster). All potential HT candidates

were validated by checking that they were not located on short, isolated scaffolds or

contigs in the genome. The flanking regions of each HT candidate pair were extracted and

checked (via pairwise alignment) to ensure that the high sequence identity was restricted

to the BovB region. This was done to check for contamination or orthologous regions.

Phylogenies of HT candidate clusters were inferred using maximum likelihood and

neighbour-joining methods (1000 bootstraps).

The same procedure was performed to screen for nucleotide L1 HT candidates. As an

extra step for L1s, we also used all ORF1 and ORF2 amino acid sequences from a

previous analysis (15) to conduct similar all-against-all BLAST searches. However, the

amino acid clusterings did not produce any possible HT candidates.

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**Author contributions** 

A.M.I. performed the analysis, interpreted the results and wrote the manuscript. R.D.K.

T.B. and D.L.A. supervised the development of work and assisted in analysing the results

and writing the manuscript. T.B. provided access to DNA samples and performed

laboratory validation experiments.

The authors declare no conflict of interest.

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**Supplementary information** 

Additional supplementary information is provided in the attached PDF.

**Data availability** 

Data generated or analysed during this study are included in the main text and

Supplementary Information. Raw sequences are provided upon request.

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## Figure legends

Figure 1: **Presence and coverage of L1 and BovB elements across eukaryotes.** The Tree of Life (29) was used to infer a tree of the 503 species used in this study; iTOL (30) was used to generate the bar graph and final graphic. The red arrow marks the L1 horizontal transfer event into therian mammals between 163-191 MYA. Branches are coloured to indicate which species have both BovB and L1 (green), only BovB (orange), only L1 (blue), or neither (black). Bar graph colours correspond to BovB (orange) and L1 (blue). An interactive version of this figure is available at: <a href="http://itol.embl.de/shared/atma">http://itol.embl.de/shared/atma</a>.

Figure 2: **HT of BovB retrotransposons**. (2a) Neighbour-joining tree (1000 bootstrap replicates) inferred using full-length nucleotide BovB consensus sequences, representing the dominant BovB family in each species. Nodes with confidence values over 50% are labelled and branches are coloured taxonomically. RTE sequence from *Schistosoma* 

mansoni was used as the outgroup. (2b) Network diagram representing the two distinct

BovB clades in bats. Nodes are coloured taxonomically apart from the RepBase (17)

sequences (light brown). RTE-1 EC and BovB Ec are shown to belong to a single family,

while BovBa-1 EF-like bat sequences form a separate family containing a single full-

length BovB from the frog *Xenopus*.

Figure 3: HT of L1 in plants and newfound chimeric L1-BovB element. (3a) TimeTree

(31) illustrating the putative L1 horizontal transfer events between plant species. Shows

only the species involved in HTs, and Amborella trichopoda as the outgroup. Background

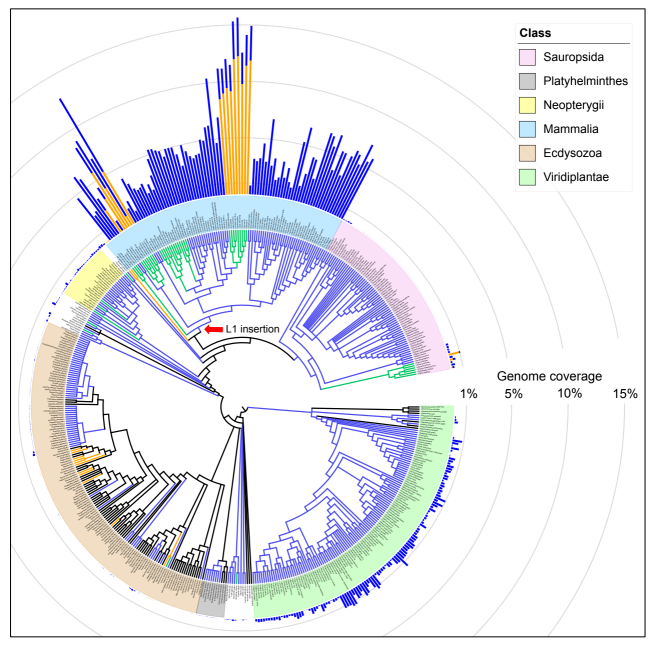
is coloured to match the ages in the geological timescale. (3b) Chimeric L1-BovB

retrotransposon found in cattle genomes (Bos taurus and Bos indicus). L1-BT and BovB

correspond to RepBase names (17), representing repeats which are known to have been

recently active. RVT 1 = reverse-transcriptase, EN = endonuclease domain. The orange

bar is the length of the entire open reading frame.



A

