

Discovery of unfixed endogenous retrovirus insertions in diverse human populations

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Endogenous retroviruses (ERVs) have contributed to more than 8% of the human genome. The majority of these elements lack function due to accumulated mutations or internal recombination resulting in a solitary (solo) LTR, although members of one group of human ERVs (HERVs), HERV-K, were recently active with members that remain nearly intact, a subset of which is present as insertionally polymorphic loci that include approximately full-length (2-LTR) and solo-LTR alleles in addition to the unoccupied site. Several 2-LTR insertions have intact reading frames in some or all genes that are expressed as functional proteins. These properties reflect the activity of HERV-K and suggest the existence of additional unique loci within humans. We sought to determine the extent to which other polymorphic insertions are present in humans, using sequenced genomes from the 1000 Genomes Project and a subset of the Human Genome Diversity Project panel. We report analysis of a total of 36 nonreference polymorphic HERV-K proviruses, including 19 newly reported loci, with insertion frequencies ranging from <0.0005 to >0.75 that varied by population. Targeted screening of individual loci identified three new unfixed 2-LTR proviruses within our set, including an intact provirus present at Xq21.33 in some individuals, with the potential for retained infectivity.

HERV-K | HML-2 | human endogenous retrovirus |
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During a retrovirus infection, a DNA copy of the viral RNA genome is permanently integrated into the nuclear DNA of the host cell as a provirus. The provirus is flanked by short target site duplications (TSDs), and consists of an internal region encoding the genes for replication that is flanked by identical LTRs. Infection of cells contributing to the germ line may result in a provirus that is transmitted to progeny as an endogenous retrovirus (ERV), and may reach population fixation (1). Indeed, more than 8% of the human genome is recognizably of retroviral origin (2). The majority of human ERVs (HERVs) represent ancient events and lack function due to accumulated mutations or deletions, or from recombination leading to the formation of a solitary (solo) LTR; however, several HERVs have been coopted for physiological functions to the host (3).

The HERV-K (HML-2) proviruses (4–9), so-named for their use of a Lys tRNA primer and similarity to the mouse mammary tumor virus (human MMTV like) (10), represent an exception to the antiquity of most HERVs. HML-2 has contributed to at least 120 human-specific insertions, and population-based surveys indicate as many as 15 unfixed sites, including 11 loci with more or less full-length proviruses (5, 6, 8, 9). To distinguish the latter from recombinant solo-LTRs, we refer to these elements as “2-LTR” insertions throughout this study. The majority of these insertions are estimated to have occurred within the past ~2 My, the youngest after the appearance of anatomically modern humans (4, 8, 11). Population modeling has implied a relatively constant rate of HML-2 accumulation since the *Homo-Pan* divergence (5, 12, 13). All known insertionally polymorphic HML-2 proviruses have signatures of purifying selection, implying ongoing exogenous replication, and retain one or more ORFs

(8, 13–15). HML-2 expression has been observed in tumor-derived tissues as well as normal placenta in the form of RNAs, proteins, and noninfectious retrovirus-like particles (3, 16–19). These unique properties raise the possibility that some HML-2 group members are still capable of replication by exogenous transmission from rare intact proviruses, from the generation of infectious recombinants via copackaged viral RNAs, or from rare viruses still in circulation in some populations. A naturally occurring infectious provirus has yet to be observed, although the well-studied “K113” provirus, which is not in the GRCh37 (hg19) reference genome but maps to chr19:21,841,544, has intact ORFs (9) and engineered recombinant HML-2 proviruses are infectious in cell types, including human cells (20, 21). The goal of this study was to enhance our understanding of such elements by identifying and characterizing additional polymorphic HML-2 insertions in the population.

The wealth of available human whole-genome sequence (WGS) data should, in principle, provide the information needed to identify transposable elements (TEs), including proviruses, in the sequenced population. However, algorithms for routine analysis of short-read (e.g., Illumina) paired-end sequence data exclude reads that do not match the reference genome. Based on read

Significance

The human endogenous retrovirus (HERV) group HERV-K contains nearly intact and insertionally polymorphic integrations among humans, many of which code for viral proteins. Expression of such HERV-K proviruses occurs in tissues associated with cancers and autoimmune diseases, and in HIV-infected individuals, suggesting possible pathogenic effects. Proper characterization of these elements necessitates the discrimination of individual HERV-K loci; such studies are hampered by our incomplete catalog of HERV-K insertions, motivating the identification of additional HERV-K copies in humans. By examining >2,500 sequenced genomes, we have discovered 19 previously unidentified HERV-K insertions, including an intact provirus without apparent substitutions that would alter viral function, only the second such provirus described. Our results provide a basis for future studies of HERV evolution and implication for disease.

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signatures stemming from such read pairs, specialized algorithms have been developed to detect TEs present within sequenced whole genomes. These methods seek to identify read pairs for which one read is mapped to a reference genome and the mate is aligned to the TE of interest (22). Additional criteria (e.g., read support, depth, presence of reads that cross the insertion junction) are then assessed to identify a confident call set. Recent applications of this general method to Illumina WGS data have indicated the presence of additional nonreference HML-2 insertions (12, 23), although validation and further characterization of these sites have been limited. Also, given the comparably short fragment lengths of typical Illumina libraries, it is not possible to distinguish between solo-LTR insertions and the presence of a 2-LTR provirus using these data alone, and experimentation is required to exclude sequencing artifacts.

To date, the number of human genomes analyzed for unfixed HML-2 proviruses is fairly small, limiting discovery of elements not present in the human reference genome, or “nonreference” elements, to those elements that are present in a relatively high proportion of individuals. Here, we build on existing detection methods to improve the efficiency of nonreference HML-2 identification from WGS data and assess the alleles present at each site. From analysis of more than 2,500 sequenced genomes, we have identified and characterized 36 nonreference insertions. We detected unique HML-2 insertions that were present in <0.05% to >75% of all samples and displayed variable presence across populations. Validation by locus-specific PCR confirmed three newly unreported 2-LTR proviruses within our dataset; one of these proviruses contains full ORFs for the viral *gag*, *pro*, *pol*, and *env* genes and lacks any obvious substitutions that would alter conserved sequence motifs, implying a potential for infectivity.

Materials and Methods

Data Analyzed. Illumina WGS data were obtained from 1000 Genomes Project (1KGP) samples, including a total of 2,484 individuals from 26 populations (24), and 53 individuals in seven populations from the Human Genome Diversity Project (HGDP) (25, 26). The 1KGP data were downloaded in aligned Binary Alignment/Map (BAM) format (<ftp://ftp.ncbi.nlm.nih.gov/1000genomes/ftp/data/>). HGDP data were processed as described (26), and are available at the National Center for Biotechnology Information (NCBI) Sequence Read Archive under accession SRP036155. Individual BAMs were merged using the Genome Analysis Toolkit (27) by population (1KGP) or dataset (HGDP). The 1KGP populations ranged from 66 to 113 individuals and had an effective coverage of $\sim 1,067x \pm 207.4x$ per pooled BAM; 53 HGDP samples were pooled to a single BAM of $\sim 429x$.

HML-2 Discovery from Read Pair Data. Candidate nonreference HML-2 LTRs were identified using RetroSeq (28). LTR-supporting read pairs were identified by running “discover” on individual BAM files, with read alignment to the HML-2 LTR5Hs consensus elements from RepBase (29) and previous reports (20, 21). RepeatMasker (30) HERV coordinates from the GRCh37/hg19 reference were used for exclusion of previously annotated sites. RetroSeq “call” was applied to the merged BAMs (above), requiring a read support of ≥ 2 for a call. A maximum read depth per call of 10,000 was applied for the increased coverage of the BAMs. To capture only novel insertions, calls within 500 bp of an annotated HML-2 LTR were excluded. Other RetroSeq options were kept at default values.

Reconstruction of Viral-Genome Junctions. For each RetroSeq candidate call, supporting read pairs and split reads within 200 bp of the assigned break were extracted from each sample and subjected to de novo assembly using CAP3 (31, 32). Assembled contigs were subjected to RepeatMasker analysis to confirm the LTR presence and type (i.e., LTR5Hs) (30), and then filtered to identify the most likely candidates, requiring separate contigs that contained the respective 5' and 3' HML-2 LTR edges, and the presence of ≥ 30 bp of both the LTR-derived and genomic sequence at each breakpoint. We examined contig pairs for the presence of 4-bp to 6-bp putative TSDs, but did not require their presence for a call. Output assemblies were aligned to the hg19 reference to confirm the position of the preintegration, or empty, site per call.

Analysis of Unmapped Reads for LTR Junction Discovery. Unmapped reads were retrieved from BAM files with Samtools (samtools.sourceforge.net/) from all 53 HGDP samples and 825 1KGP samples (≥ 10 samples per 1KGP population) and searched for a sequence that matched the 5' HML-2 LTR edge (TGTTGGGGAAAAGCAAGAGA), 3' LTR edge (GGGGCAACCCACCCCA-TACA), or 3' LTR variant (GGGGCAACCCACCCATTCA) that is observed in a subset of human-specific elements, requiring ≥ 10 bp of non-LTR sequence per read. Reads matching reference HML-2 junctions were removed. Candidate reads were then aligned to the hg19 reference to identify genomic position. Sequences with no match to hg19, with <90% identity, or that aligned to gaps or multiple genomic positions were searched against the chimpanzee (panTro4) and gorilla (gorGor3) references, and available human WGS data from the NCBI Trace Archive to identify insertions in structurally variable regions.

Validation and Sequencing. DNA from samples yielding positive reads was obtained from Coriell or the Foundation Jean Dausset-Centre d'Étude du Polymorphisme Humain. Coordinates for each insertion were based on mapping of assembled contigs or read-captured flanking sequence to the hg19 reference. PCR was performed with 100 ng of genomic DNA using primers flanking each site to detect either the empty site or solo-LTR alleles. A separate PCR was run to infer a 2-LTR allele with a primer situated in the HML-2 5' UTR paired with a flanking primer (6, 8, 33). Capillary sequencing was performed on at least one positive sample. The 2-LTR alleles were amplified in overlapping fragments from a single sample and sequenced to $\geq 4x$ (8, 9), and a consensus then constructed with the read traces from each site. Complete sequences are available in the NCBI GenBank under accession nos. KU054242–KU054309.

Phylogenetic Analysis. Full-length sequences representing either solo-LTRs or proviral 5' and 3' LTRs were aligned to the consensus HML-2 using ClustalX (34); the alignment was then edited, and truncated LTRs were removed (8). A single neighbor-joining tree was generated from the remaining 68 insertions (90 total LTRs) using MEGA6 (35). The Kimura 2 parameter model was used for branch length estimation, with α of 2.5 and deletions treated pairwise. Support for the tree was assessed using 1,000 bootstrap replicates.

Age Estimations. LTR divergence was used to infer the time since insertion, normalized to a neutral mutation rate of 0.24–0.45% per My as measured by calculating the divergence between orthologous human and chimpanzee HML-2 proviruses (6, 8). Alternatively, using mutation rates directly obtained from pedigrees (36) results in estimated times of insertion fourfold to ninefold as old and implausible integration times for many proviruses. Briefly, the nucleotide differences were totaled between proviral 5' and 3' LTRs, and the total was divided by the LTR nucleotide length. The percentage of divergence was then divided by the upper and lower bound mutation rates for age range estimation (million years) (6, 8).

In Silico Genotyping. Genotyping was performed for both reference (hg19) and nonreference unfixed insertions using a read-based method that has been used for genotyping Alu TEs in humans (32, 37). Briefly, discrete reference (e.g., the empty state) and alternate (e.g., the insertion state) alleles were recreated for each locus, including ± 600 bp upstream and downstream of the insertion point based on hg19 coordinates. Within those coordinates, Illumina read pairs that had at least one read mapped to the empty allele were extracted for each site (requiring a mapping quality score > 20). Treating the reconstructed alleles as the target genome, genotype likelihoods were then determined based on remapping of those reads to either allele, with error probabilities based on read mapping quality as described previously (32, 38). Samples without reads aligning to the reconstructed reference and alternate alleles for a particular site were not genotyped at that site. Insertion allele frequencies were estimated per site for all genotyped samples as the total number of insertion alleles divided by the total number of alleles. Detection frequencies (the proportion of individuals carrying the insertion) were calculated as the number of individuals with the insertion divided by the total number of individuals genotyped at each locus. We note that the reference insertion at 7p22.1, which is present as a tandem duplication of two proviruses that share a central LTR (6, 15), was treated as a single insertion (chr7:4,622,057–4,640,031). Nine of the 36 nonreference loci could not be aligned to the hg19 reference and were excluded from genotyping: insertions within duplicated segments (we refer to these as dup1 through dup4), insertions of unusual assembled structure (10q24.2b and 15q13.1), or insertions that could not be mapped to the hg19 assembly (10q26.3, 12q24.32, and 22q11.23b).

Proportion of Provirus Carriers. Unique 30-mers were identified from a set of 51 reference and nonreference elements from the HML2 subgroup using Jellyfish (39). Candidate 30-mers were further mapped against the GRCh37 genome reference using mrsFAST and mrFAST (40), and k-mers with >100 matches within an edit distance of two were omitted, resulting in a set of 83,343 k-mers. The position of each 30-mer in each HML2 element was determined, and 1,445 k-mers that crossed LTR-internal proviral junctions were omitted, leaving 5,698 k-mers that were unique to an LTR and 76,200 k-mers that were unique to the internal sequence. Total observed counts for each k-mer were determined in WGS sequence data from 53 HGDP and 2,453 1KP samples. The median k-mer depth for each element in each sample was determined. Median depths were normalized per sample by dividing by the maximum median depth observed for a proviral sequence. Elements with a normalized median k-mer depth ≥ 0.25 were considered to be present in a sample. The proportion of individuals for which an element was present was then determined for each population.

Results

HERV-K (HML-2) Insertions Discovered from WGS Data. The goal of this study was to use the extensive available WGS data in the 1KGP and HGDP collections to identify relatively rare polymorphic nonreference HML-2 insertions. To make the fullest use of all sequence information available within these data, we applied two approaches to identify candidate nonreference HML-2 insertions in the raw reads for these collections (Fig. 1).

First, we identified insertions based on read pair signatures using the program RetroSeq (28) (Fig. 1, *Left*). To improve the detection of insertions present in multiple samples, we combined reads within a population (1KGP) or study (HGDP) (32). Excluding calls within ± 500 bp of a reference HML-2 sequence, we obtained 140.3 ± 56.1 candidate calls per pool. Next, we applied

a de novo assembly approach to insertion-supporting reads to reconstruct the LTR-genome junction for as many sites as possible (32). Given the size of HML-2 LTRs (~968 bp per LTR), we inferred the presence of an insertion based on the presence of separately assembled 5' and 3' breakpoints. This requirement reduced false-positive calls, for example, as caused by SVA elements (SINE-VNTR-Alu), which have high identity to bases 1–329 of the HML-2 LTR. A total of 29 candidate HML-2 insertions with a flanking sequence were assembled, including K113 (19p12b; also see Fig. S1 *A* and *B* and Table 1).

As a second approach, we mined unmapped reads for evidence of LTR-genome junctions captured in reads that could not be placed on the human reference (Fig. 1, *Right*) and would therefore be missed using current read-based detection methods, such as RetroSeq. Using this approach permitted the identification of insertions in regions absent from the human reference. Excluding reads that could be aligned to annotated HML-2 junctions, we obtained overlap for the 29 candidate sites identified above, as well for as seven loci not found in assembled RetroSeq calls (Fig. S1 *C* and Table 1). Our final call set includes 17 insertions identified in recent reports from Marchi et al. (12) and Lee et al. (41). The nomenclature for all sites is as maintained in those studies and in other previous reports (8, 33).

Validation and Sequencing. We validated the presence of 34 of the 36 candidate insertions in at least one individual predicted to have the insertion (Table 1 and Dataset S1). The remaining two sites (at 10q24.2 and 15q13.1) were predicted to have an unusual inverted repeat structure based on assemblies of supporting reads at either site (Fig. S2), and could not be conclusively confirmed by sequencing, possibly due to hairpin formation. For the 34 validated nonreference sites, we confirmed 29 sites as having solo-LTRs and five sites with 2-LTR proviruses (at 8q24.3c, 19p12d, 19p12e, Xq21.33, and the published K113 provirus at 19p12b; also see Table 1). Four of the solo-LTRs were situated within duplicated segments and could not be mapped to unique positions in the hg19 reference (dup 1–dup 4), and two insertions, at 12q24.32 and 10q26.3, were located within structurally variable regions that are absent from the hg19 reference (Fig. S3). One insertion was initially mapped to the reported 9q34.11 locus (12, 41); however, comparison of the Sanger reads from its validated LTR-genome junctions revealed unexpectedly low identity in the extended flanking sequence. Our reexamination of this site indicates it maps instead to a region that is not in hg19 but is present in an alternate scaffold in the GRCh38 assembly at 22q11.23 (Table 1). This discrepancy may explain why this particular site has only been previously inferred by reads supporting only the 5' breakpoint of the integration (12, 41, 42).

We obtained full sequences for 30 of the 36 candidate insertions in at least one individual predicted to have the insertion (Dataset S1); these sequences included the full-length insertion at Xq21.33 that was found to have intact viral ORFs (NCBI GenBank accession no. KU054272). The remaining six insertions were extracted or reconstructed from public sequence databases for subsequent analysis as follows. The full sequence from one locus identified within a duplicated segment was reconstructed from Sanger reads corresponding to that site from the NCBI Trace Archive (dup1). The sequence flanking the insertion at 12q24.32 could be mapped to a previously sequenced fosmid clone in a region corresponding to an encompassing deletion of ~14.3 kb in the hg19 reference (43) (Fig. S3*4*). Another insertion, corresponding to a 2-LTR provirus, was also from a sequenced fosmid clone (19p12d) as reported (41, 44). The complete sequence of the K113 provirus (19p12b) was from the GenBank (accession no. AY037928). One solo-LTR, 1p31.1c, was detected and validated as a solo-LTR in a single individual of the 1KGP Yoruba. We searched for, but did not find evidence of, this site in subsequent PCR screens of other samples.

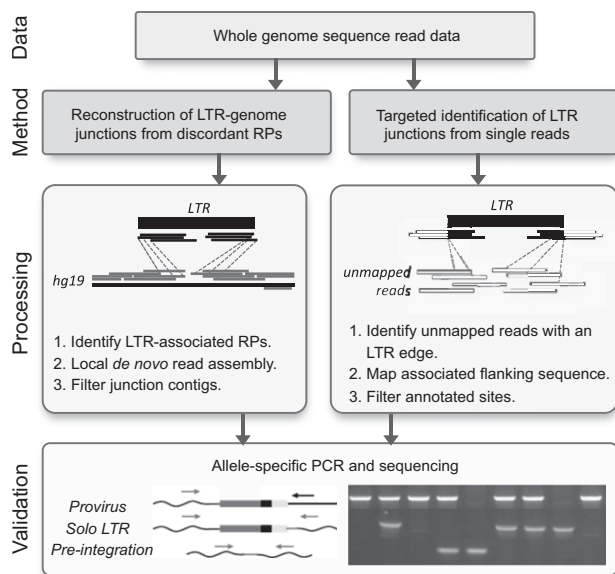


Fig. 1. Approaches for the detection of nonreference HML-2 insertions from WGS read data. Illumina short reads were processed by one of two methods. (*Left*) Read pairs (RPs) were identified that have one read mapped to the genome (gray) and mate to reads that map to the sequence matching the HML-2 LTR consensus (black). Supporting reads from each site were extracted and subjected to local assembly, and the resulting contigs were analyzed for the presence of LTR-genome junctions. (*Right*) Unmapped reads from each sample were identified that contained a sequence corresponding to the LTR edge, and the cognate sequence was then used to determine candidate integration positions from genomic data. (*Bottom*) PCR and capillary sequencing were used to validate candidate insertions in reactions that used flanking primers (gray arrows) to detect the presence of a solo-LTR or empty site, or a flanking primer paired with an internal proviral primer (black arrow) to infer the presence of a full-length allele. Representative products are shown in a genotyping gel to the right.

Table 1. Nonreference HML-2 insertions in human genomes

Locus	Coordinate GRCh37/hg19	Alias*	Alleles [†]	Flanking region and other properties	First report in humans (source)
1p13.2 [‡]	chr1:111,802,592	De5;K1	LTR, pre	L1 (L1PA6)	(41)
1p21.1 [‡]	chr1:106,015,875		LTR, pre		(12)
1p31.1c	chr1:79,792,629		LTR, pre	AluSz	This study
1q41	chr1:223,578,304	K2	LTR, pre	L1 (L1MDa)	(12)
3q11.2	chr3:94,943,488		LTR, pre	L1 (L1PA10)	This study
4p16c	chr4:9,603,240	K6	LTR, pre	ERV1 (HERVS71)	(12)
4p16d	chr4:9,981,605		LTR, pre	L2 L2b; SLCA29 intron 5/6	This study
5p15.32	chr5:4,537,604		LTR, pre	ERV1 (LTR1C)	This study
5q12.3	chr5:64,388,440	Ne7;K12	LTR, pre	L1 L1M6	(12)
5q14.1	chr5:80,442,266	De6/Ne1;K10	LTR, pre	RASGRF2 intron 17	(12)
6p21.32	chr6:32,648,036		LTR, pre	L1 (L1PA10)	(12)
6p22.3	chr6:16,004,859		LTR, pre	AluSx	This study
6q26	chr6:161,270,899	De2;K12	LTR, pre		(12)
7q36.3	chr7:158,773,385		LTR, pre		This study
8q24.3c	chr8:146,086,169		pro, pre	ERV1 (LTR46); COMMD5 intron 9 of transcript variant 2; <i>gag</i> and <i>pro</i> ORFs	This study
10q24.2b	chr10:101,016,122	De12		ERL MalR (MSTD); unexpected structure	This study
10q26.3 [§]	chr10:134,444,012		LTR, pre	INPP5A intron 2	This study
11q12.2	chr11:60,449,890	De4;K18	LTR, pre	L1 (L1M4); LINC00301 intron 6	(12)
12q12	chr12:44,313,657	Ne6;K20	LTR, pre	L1 (L1MB1); TMEM117 intron 2	(12)
12q24.31	chr12:124,066,477	K21	LTR, pre	AluSx1; LOC101927415 exon 3	(12)
12q24.32 [§]	chr12:127,638,080–127,639,871		LTR, pre	ERV1 (MER57); deleted in hg19; from fosmid CloneDB: AC195745.1 bases 17648–18615	(43)
13q31.3	chr13:90,743,183	Ne2;K22	LTR, pre	SINE (FLAM_A); LINC0559 intron 3	(12)
15q13.1	chr15:28,430,088			HERC2 intron 56; unexpected structure	This study
15q22.2	chr15:63,374,594	K24	LTR, pre		(12)
19p12b	chr19:21,841,536	De1;K113	pro, pre		(9)
19p12d	chr19:22,414,379		pro, pre	Deletion in 5' LTR; <i>pro</i> ORF; insertion within fosmid clone accession AC245253.1	(41)
19p12e	chr19:22,457,244	De11	pro, pre	AluSq	This study
19q12 [§]	chr19:29,855,781	De3;K28	LTR, pre	LOC284395 intron 9	(12)
19q13.43	chr19:57,996,939	Ne5	LTR, pre	2 kb upstream of ZNF419	This study
20p12.1	chr20:12,402,387	De14*;K30	LTR, pre		(12)
22q11.23b [§]	chr22:23852639–23852640	De7;K16	LTR, pre	ERVL-MaIR (MLT1C); maps to Hg38 alt locus scaffold 22_KI270878v1_alt:156355–180653	This study
Xq21.33	chrX:93,606,603	De9	pro, pre	L1 (L1MD1); <i>gag</i> , <i>pro</i> , <i>pol</i> , <i>env</i> ORFs	This study
Dup 1 [§]	Not determined		LTR	Flank maps to centromere associated duplications on multiple chromosomes	This study
Dup 2 [§]	Not determined		LTR, pre	Flank maps to duplicated regions within predicted FAM86 and ALG1L2 exonic variants	This study
Dup 3 [§]	Not determined		LTR, pre	Flank maps to 3 segmental duplications on chr1	This study
Dup 4 [§]	Not determined		LTR, pre	Deletion in hg19 reference; putative empty site on chr19 within fosmid CloneDB: AC232224.2	This study

*Reported originally in the sequenced Neandertal (Ne) or Denisovan (De) by Agoni et al. (42) or Lee et al. (51), or in modern humans (K) by Marchi et al. (12) or Lee et al. (41).

[†]Alleles detected. LTR, solo LTR; pre, preinsertion site; pro, 2-LTR provirus.

[‡]Previously PCR validated as solo-LTR by Lee et al. (41).

[§]Insertion is located within an encompassing structural variant not present in the hg19 reference.

Estimated Frequencies of Unfixed HML-2 Loci. We performed *in silico* read-based genotyping to obtain estimations of the allele frequencies of 27 nonreference insertions with clear integration coordinates, and extended the analysis to include 13 annotated polymorphic HML-2 loci from the hg19 human reference (5, 8) (Dataset S2). Briefly, reference and alternate alleles representing each HML-2 locus were recreated, and individual genotypes were then inferred based on the remapping of proximal Illumina reads to the reconstructed alleles per site per sample (*Materials and Methods*). Given the larger size of the HML-2 LTR (~968 bp) and relatively short reads in these data, 2-LTR and solo-LTR insertions are indistinguishable in read-based genotyping alone, such that genotypes were based on the presence or absence of

the HML-2 insertion at each locus. Values reported below correspond to allele frequencies unless otherwise noted.

Estimated frequencies of the variable HML-2 insertions present in the reference genome ranged from ~0.25 to >0.99 in genotyped samples (Fig. 2, *Upper*). Sites with the highest estimated frequencies corresponded to those loci previously reported with a solo-LTR or provirus present, but not a preinsertion site, based on limited PCR screens of those sites (6) (at 1p31.1, 3q13.2, 7p22.1, 12q14.1, and 6q14.1 in Fig. 2). This pattern is consistent with variability at these sites based predominantly on the 2-LTR and solo-LTR states. Genotyping of the insertions at 11q22.1 and 8p23.1a (K115) implied the presence of both insertion and preinsertion alleles, also consistent with PCR screens in other reports (6, 9, 33, 45),

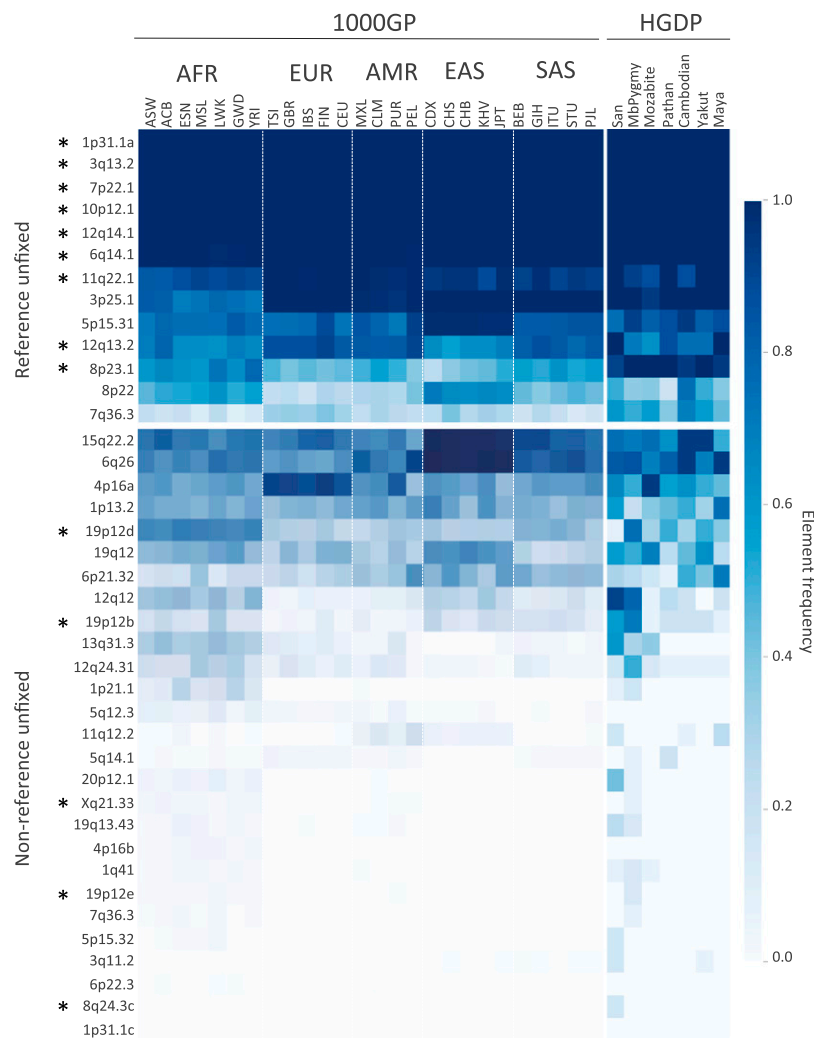


Fig. 2. Estimated insertion allele frequencies of unfixed HML-2 insertions in humans. A total of 40 HML-2 loci were subjected to *in silico* genotyping: 13 sites represented the unfixed HML-2 loci from the hg19 reference, and 27 sites corresponded to nonreference polymorphic HML-2 reported here. Genotypes were inferred for each unfixed HML-2 locus across samples based on remapping of Illumina reads to reconstructed insertion or empty alleles corresponding to each site. Samples lacking remapped reads at a particular site were excluded from genotyping at that site. Allele frequencies were then calculated for each population as the total number of insertion alleles divided by total alleles. Allele frequencies are depicted as a heat map according to the color legend to the right. The 1KGP (1000GP) and HGDP populations are labeled above (also refer to [Dataset S1](#) for population descriptors and other information). The locus of each of the unfixed HML-2 loci is labeled to the left according to its cytoband position. An asterisk is used to indicate insertions that have confirmed full-length copies. (*Upper*) Estimated distribution of reference unfixed HML-2 [from loci reported by Subramanian et al. (11) and Belshaw et al. (5)]. (*Lower*) Estimated distribution of nonreference HML-2 insertions. AFR, African; AMR, Admixed American; EAS, East Asian; EUR, European; SAS, South Asian.

noting the higher frequency of K115 within our samples ($\sim 53\%$) than in those reports (up to $\sim 34\%$ depending on ancestry). Four unfixed reference solo-LTRs ranged in frequencies from ~ 0.25 to as high as ~ 0.93 , also consistent with previous analysis of these sites (5). Extending the analysis to the 85 remaining human-specific HML-2 insertions that are suitable for genotyping in the human reference (81 solo-LTRs and four full-length proviruses) (5, 8) was consistent with sample-wide fixation among the vast majority of these loci; just eight loci had evidence of the nonreference allele among genotyped samples (Fig. S4 and Dataset S3).

Estimated frequencies of the nonreference HML-2 insertions were inferred to be from <0.0005 (the insertion having clear support in one or few individuals) to >0.75 of genotyped samples (Fig. 2, Lower). More than half of the nonreference HML-2 insertions were rare, with 15 insertions detected at frequencies of $<5\%$ and six insertions in $<1\%$ of all samples; just four of these loci have been previously reported (12). Sites with the lowest allele frequencies were predominantly in individuals of African

ancestry, with nine of 13 loci inferred in $<5\%$ of all samples but mostly limited to African populations, although insertions were also detected in non-African samples at ~ 0.005 to ~ 0.016 in those populations (e.g., at 5q14.1 and Xq21.33 in Fig. 3). The solo-LTR insertion at 1p31.1c was only identified in a single sample and was not detected in any other sample by genotyping; however, this observation does not exclude the possibility of its presence in some individuals, given the variability in read coverage between samples (*Discussion*). Nine of the 10 common insertions (detected in $>10\%$ of all samples), including the K113 provirus, have been previously reported in searches of WGS data (12, 41). A comparison of the overall presence of each HML-2 insertion, calculated as the proportion of individuals with evidence of the insertion, was generally in agreement with those reports (Fig. S5). The presence of K113 was estimated at a higher prevalence across samples here than in previous reports, in $\sim 27\%$ of all samples and as high as $\sim 52\%$ in African

populations, consistent with the prevalence of this insertion varying with ancestry (45).

Because the above analysis cannot distinguish between solo-LTR and 2-LTR alleles, we used a k-mer counting approach to infer the presence of 2-LTR HML-2 sequences in each sample. WGS data from each sample were queried using a catalog of 30-mers unique to each proviral sequence. Elements with a normalized depth ≥ 0.25 were inferred to be present in a sample. We find large variation in the proportion of individuals estimated to carry 2-LTR alleles for different elements, with 2-LTR alleles for elements present in the reference genome detected in a high proportion of individuals, whereas the nonreference 2-LTR insertions discovered in this study are extremely rare (Fig. S6 and Dataset S3). For example, we identified 44 samples with k-mer counts consistent with the presence of a 2-LTR allele for the Xq21.33 insertion.

LTR-Based Analysis of Unfixed HML-2 Proviruses. Using sequence information obtained for each insertion, we performed an LTR-based phylogenetic analysis (Fig. 3). Because proviral LTRs are identical at integration, the two LTRs on the same provirus will always pair in a phylogenetic tree, barring recombination between elements (46). Their unique source is further supported by the presence of TSDs, which are preserved during solo-LTR formation (6, 46, 47). To create the most informative tree, we added the LTRs from 21 human-specific proviruses, including 11 polymorphic 2-LTR insertions as reported by Subramanian et al. (8) and four unfixed solo-LTRs as reported by Belshaw et al. (5), to our validated set of LTRs from three 2-LTR proviruses (insertions at 8q24.3c, 19p12e, and Xq21.33), the 3' LTR of a truncated provirus (19p12d), and 30 solo-LTRs.

The analysis revealed a major lineage leading to a well-supported clade that contained all human-specific and polymorphic HML-2 sequences (Fig. 3A, boxed); a minor lineage included HML-2 elements that are fixed in humans (8) and did not contain any newly identified insertions. This phylogeny is consistent with previous analyses (8, 13, 46); however, the addition of our 34 unfixed loci permitted a more detailed examination of variable insertions (Fig. 3B). The majority of unfixed insertions clustered with the HML-2 consensus LTR (● in Fig. 3B) in a clade (* in Fig. 3B) whose members tended to have the shortest branches, consistent with their relatively recent integration and insertionally variable presence within humans (also refer to filled boxes in Fig. 3B). The human-specific reference elements were also distributed within this clade, consistent with previous observations (8). The remaining insertions were on branches with longer lengths, reflecting changes that accrued before insertion as well as during their longer existence as endogenous elements. We searched for, but did not observe, mispaired LTRs from the 2-LTR proviruses reported here. Subsequent examination of the TSDs from these proviruses confirmed all were intact, indicating these elements have not seeded past rearrangements (46). Extending this comparison with the TSDs of the identified solo-LTRs also verified their intact state.

Properties of Nonreference 2-LTR HML-2 Integrations. Assuming a constant mutation rate, the nucleotide divergence between cognate 5'-3' LTR pairs may be used to estimate the time since integration (47). Using this method, we previously estimated the average age of the human-specific 2-LTR insertions to within ~ 2.7 (± 1.1) My (8). Applying this method to the 2-LTR proviruses identified here suggests these insertions were formed within ~ 0.67 – 1.8 Mya (Table 1). Further refinement for the youngest elements is limited, because the variance for age estimates increases significantly for insertions with very little or no LTR divergence. The 19p12d provirus was excluded from this analysis due to deletion of the majority of its 5' LTR (Fig. 4). This truncation has also been observed in a few reference LTR5Hs

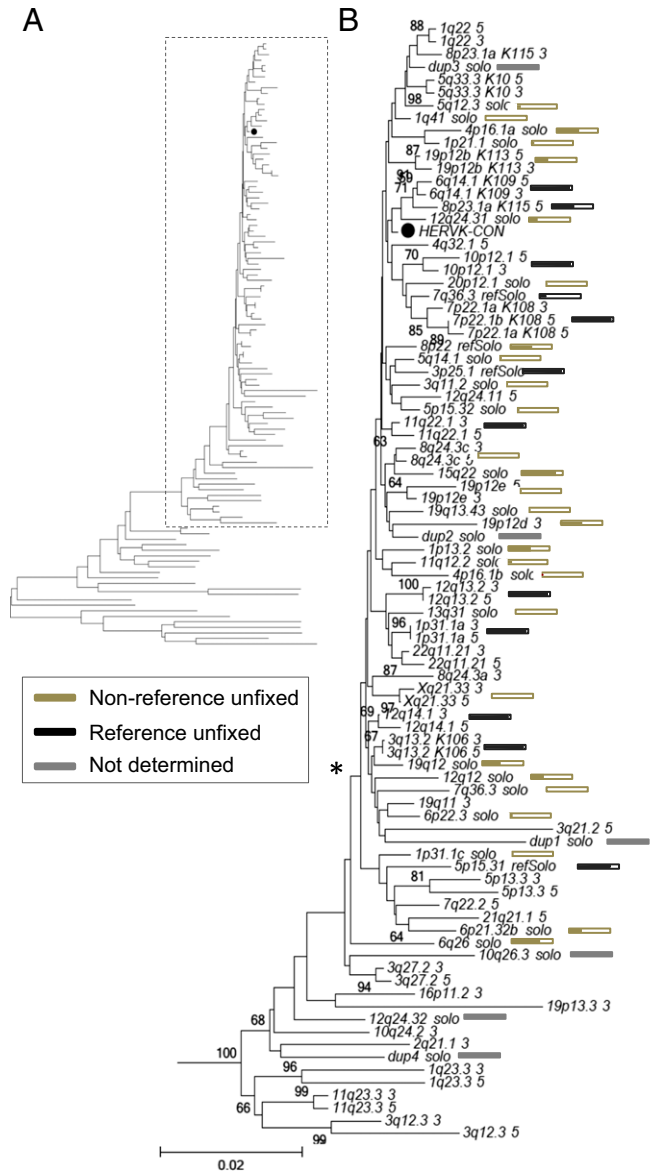


Fig. 3. Phylogenetic construction of HML-2 LTRs within humans. (A) Neighbor-joining tree was constructed based on the aligned nucleotide sequences corresponding to HML-2 LTRs from the LTR5Hs group, specifically including those nucleotide sequences considered to be human-specific and/or polymorphic. The LTRs were extracted from (i) all reference HML-2 proviruses previously inferred as belonging to the LTR5Hs HML-2 subgroup [as reported by Subramanian et al. (11)], (ii) unfixed reference solo-LTRs [as reported by Belshaw et al. (5)], and (iii) unfixed nonreference insertions as reported here. Both 5' and 3' LTRs were used for full-length insertions, when present. The closed circle (●) indicates the taxon corresponding to the HERV-K_{CON} LTR within the tree. Classic nomenclature has been included in taxon names for the better studied insertions: K113 (19p12b), K108 (7p22.1), K115 (8p23.1a), K106 (3q13.2), and K109 (6q14.1). (B) Detailed view of branches representing unfixed HML-2 insertions. Individual HML-2 loci are indicated for each branch as follows: the cytoband followed by a 5' or 3' for the 5' or 3' LTRs from full-length insertions, solo for nonreference unfixed solo-LTR insertions, or refSolo for reference unfixed loci. An asterisk is used to indicate the position of the clade containing the majority of unfixed insertions. Boxes are used to indicate estimated allele frequencies for each unfixed insertion at the end of each respective branch. The filled area within each box is shown as proportional to the estimated frequency of the insertion in all samples; the derived values are provided in Dataset S3. Gold and black boxes are used to represent nonreference and reference unfixed insertions, and gray bars indicate the elements for which the frequency could not be determined.

2-LTR insertions (8q24.3a, 10q24.2a, and 19q11) (8), which all possess unique, intact TSDs and do not share any flanking sequence, supporting their classification as independent integrations. It is likely that this common rearrangement occurred as a result of aberrant strand transfer during RT, as has been discussed (48).

HML-2 proviruses are classified by the presence or absence of a 292-bp deletion at the *pol-env* boundary, designated “type 1” or “type 2,” respectively. Deletion of a splice site in type 1 elements obliterates *env* and *rec* expression and results in mRNA encoding an ~9-kD protein, Np9, of possible cellular function (17, 19, 49). Sequence comparison of the 2-LTR proviruses identified here classifies the 19p12d and 19p12e elements as type 1 and 8q24.3c and Xq21.33 as type 2. The 19p12d and 19p12e insertions were found to have intact *pro* and *gag* ORFs, respectively, and the 8q24.3c insertion had both *gag* and *pro* ORFs. The Xq21.33 2-LTR element was found to be intact with ORFs for all HML-2 encoded genes (i.e., *gag*, *pro*, *pol*, *env*, *rec*) (Fig. 4). Indeed, it differs by only 39 of a total of 2,820 amino acids (98.6% amino acid identity) in all genes from the infectious consensus provirus HERV-K_{CON} (21). We searched for, but did not observe, any substitutions that would alter conserved sequence motifs (including the YIDD motif in reverse transcriptase), making this element a candidate for activity. This provirus is only the second naturally occurring intact HERV to be described, with the other being the noninfectious K113 (19p12b) that shares 98.9% amino acid identity to HERV-K_{CON} (9). Its potential for generation of infectious virus is currently under investigation.

Discussion

We report 36 nonreference HML-2 insertions, including 19 previously identified loci, from analysis of WGS read data from more than 2,500 globally sampled individuals. Seventeen of the 36 sites were recently reported in humans (12, 41), although with limited validation or element characterization. Here, we take full advantage of the 1KGP and HGDP WGS read data to identify nonreference viral-genome junctions from assembled anchored read pairs and individual unmapped reads, and use these data to estimate the presence of each of these elements within our sampled populations. We validated the presence of 34 of the 36 loci, including five loci with 2-LTR proviruses (including K113) and 29 solo-LTRs, and report the complete sequences for 30 of these insertions, including a 2-LTR provirus at Xq21.33 that appears to be intact. We provide a thorough analysis of unfixed HML-2 insertions that complements and builds on previous studies, and should enable future examination of the HML-2 group.

We used the available reads from each sample for in silico genotyping of a subset of sites to infer the population-wide frequencies of unfixed HML-2 elements, which is impractical on this scale in standard PCR-based screens. The inferred allele frequencies of the nonreference insertions ranged from 0.05 to >75% of genotyped samples and varied between populations, generally with the highest presence in African populations. With the exception of two previously sequenced sites in our set (dup1 and 12q24.11), all nonreference insertions were validated in samples of African ancestry, as has been observed for all HERV-K loci characterized to date, implying their insertion before the human migration out of Africa ~45,000–60,000 y ago (50). These two insertions could not be confidently mapped to the hg19 reference, and were therefore excluded from genotyping. All but one nonreference insertion was identified in more than one individual, with the exception being the 1p31.1c solo-LTR validated in NA18867. Genotyping of that site failed to reveal (but does rule out) its presence in other individuals. Analysis of the surrounding region revealed the presence of several SNPs that were unique to NA18867 within the 1KGP panel, suggesting that 1p31.1c may be associated with a very rare haplotype, rather than a de novo event, in the absence of comprehensive screening. These observations support the utility of short read data for

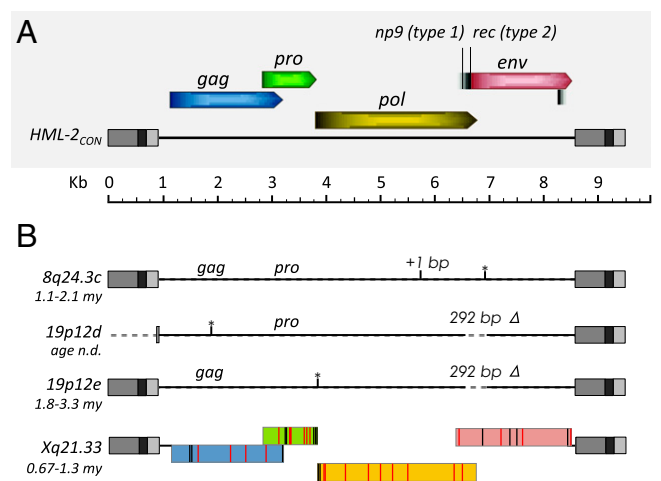


Fig. 4. Features of newly identified HML-2 proviruses in humans. (A) Schematic representation of the consensus HML-2 provirus, including the viral gene positions and frames to scale. Splice sites for np9 (type 1 insertion, 292 bp Δ) and *rec* (type 2) are indicated. Regions within the LTRs are colored in gray: U3, medium; R, dark; U5, light. (B) Features of nonreference identified proviruses are shown to scale. The region of 292 bp is labeled for type 1 insertions. Age estimations are shown for each site. n.d., not determined. The black vertical line indicates a frameshift mutation (as indicated “+1 bp”); black lines with asterisks are used to indicate positions of stop codons where present. Reading frames are shown for the Xq21.33 2-LTR provirus as colored as in A. Black vertical lines within the frames indicate the positions of base changes that are observed in other full-length HML-2 proviruses. Red vertical lines are used to indicate base changes that are unique to the sequenced Xq21.33 provirus.

element discoveries and sequence-based analysis, but also underscore the necessity of additional experimental validation steps and characterization of candidate proviruses.

Eight of our validated loci have been recently reported in the genomes of two sequenced archaic samples (42, 51) in addition to modern humans (12). We confirmed an additional three reported “archaic” sites in our data [19p12e and 10q24.2b, respectively: “De11” and “De12” in the study by Agoni et al. (42); 19q13.43: “Ne5” in the study by Lee et al. (51)], but found no evidence of the remaining eight reported archaic events. Properties of these 11 HML-2 loci are more consistent with insertion before the most recent ancestor with modern humans ~0.6–0.8 Mya (52) than with introgression. For example, the 2-LTR insertions at 19p12e and Xq21.33 are most prevalent in samples of African ancestry, and LTR divergences indicate their respective insertions to have been ~1.8–3.3 Mya and ~0.67–1.3 Mya, consistent with this time frame. Both sites are rare, with sample-wide allele frequencies estimated at 0.0103 and 0.0157 (~0.026–0.069 in the African sample) in our data (Dataset S3). Of the remaining genotyped loci also in archaic genomes, each was also most represented in African ancestry, with exception of the insertions at 11q12.2 and 5q14.1 (sample-wide allele frequencies estimated at ~0.046 and 0.026) that appeared most frequently in populations from the Americas or of East Asian ancestries but are also present in African populations, again implying ancient events (50). Given their overall distribution, it is likely these insertions are also older, although our ability to estimate insertion times is limited, given their presence as solo-LTRs.

We confirmed the presence of full-length proviruses at four loci, including the Xq21.33 provirus, which appears to be intact and without obvious defects, which implies the potential for replication competence and is now under further investigation. Given a genomic mutation rate of $\sim 2.2 \times 10^{-9}$ changes per site per year (53), an ERV could maintain infectivity over very long periods, and a number of infectious ERVs are known in other

species, including mice, cats, and some birds (3). Although such elements are likely to be regulated by silencing and downstream host mechanisms, disease states causing prolonged reactivation of HERVs could drive expression of such proviruses or the generation of recombinant infectious chimeras, as has been shown to occur in Ab-deficient mice (54) and as claimed for HML-2–derived transcripts in the blood of HIV-infected individuals (44). Indeed, a “recombinant” HML-2 provirus engineered from just three well-studied defective reference loci is infectious (derived from portions of K109, K115, and K108, respectively, at 6q14.1, 8p23.1a, and 7p22.1) (20), as are the HML-2 consensus genomes (20, 21). We anticipate that sequence-based comparisons and future experimental interrogation of intact proviruses, such as Xq21.33 and others that have yet to be discovered, will give insight into the functionality of similar HML-2 members.

Expression of HML-2 elements has been studied foremost in the context of disease, particularly from tumor-derived tissues (reviewed in 3, 16, 17), but also in otherwise normal human tissues (19). This expression has been shown to exhibit tissue specificity in the form of proviral RNAs from both type 1 and 2 proviruses that have no apparent match to annotated loci (16, 17), implying the presence of transcriptionally active but not yet characterized loci. Although the consequence of such expression is not fully understood, the correlation of HML-2 expression with certain disease states suggests that such variable expression may provide a useful biomarker. Because RNAs corresponding to all previously known polymorphic HML-2 proviruses (and the majority of human-specific elements) have been identified in such assays, additional polymorphic HML-2 copies are likely to be transcribed under certain conditions, justifying the continued characterization of “new” loci and the regulation of their expression. Our analysis of four additional nonreference HML-2 proviruses indicates the presence of discriminatory nucleotide positions within each of these elements (Fig. S6); such sites should aid their assignment in future experimental assays.

Several HML-2 insertions were nearby or within genic regions (Table 1). For example, the 8q24.3c provirus is situated within *COMMD5*, a gene involved in hypertension-related renal repair whose expression is elevated in the kidneys of hypertension-resistant rat models (55). The 4p16d LTR, in ~1.5% of all samples (with highest prevalence in African ancestry), was within the *SLCA29* (*GLUT9*) gene; the *GLUT9* uric transporter is a direct target of p53 and is implicated in antioxidant functions (56). Also, the 5q14.1 LTR lies within the *RASGFR* gene associated with regulation of dopamine neuron activity and reward sensitivity in alcohol use (57). Previous mining of cohort-based WGS data has inferred the 5q14.1 LTR in ~14 to ~30% of those samples (12, 41), although it was detected in ~2.7% of all samples here, possibly explained by the global survey in the data used. Genotyping of this site did permit population distribution estimates, in which we found the highest prevalence in European and American ancestry (up to ~7% of samples), with apparent absence in East Asian individuals (Fig. 2 and Dataset S3). The

relationship between a particular LTR and a biological effect requires further investigations, but these observations serve as a reminder that such insertions may be associated with phenotypic effects in some individuals.

Low levels of replication have been suggested based on the presence of unfixed HML-2 loci (13). Coalescent analysis of globally sequenced LTRs from the K106 provirus (the reference insertion at 3q13.2) has produced an age estimation of 0.15 My based on sequence conservation of that site across sampled individuals (11). Other studies suggest replication until at least 0.25 Mya (12) based on modeling estimations of an expected number of loci, given the number of observed unfixed sites and the proportion of individuals predicted to carry those insertions (12). Although we cannot rule out the possibility of ongoing replication, our comparison of 2-LTR sites suggests a most recent time of insertion at least ~0.67 Mya for those proviruses and we find no evidence of insertions with evidence of more recent formation, noting limitations in properly estimating integration times for recombinant solo-LTRs. The number of rare insertions in our data (15 insertions in <5% and six in <1% of all samples), including the 1p31.1c LTR detected in a single individual and the 2-LTR provirus at 8q24.3c in just a few samples (from the HGDP San and 1KGP Yoruba populations), suggests that additional remaining HML-2 loci are likely to be very rare, specific to groups not yet surveyed, or within low coverage regions of the genome. The rarity of such proviruses, however, is likely to reflect more recently integrated proviruses as well as less time for selective removal of deleterious, pathogenic ones. Continued efforts to analyze additional genome sequences, particularly from previously unstudied populations (particularly of African ancestry), will contribute to the identification of intact and potentially replication-competent proviruses, as is supported by this study.

Our approach shares limitations common to all read-based discovery methods. Given the variability in per-sample coverage, we have likely missed other sites that may be present in one or a few samples or insertions located in otherwise inaccessible regions of the genome. Likewise, other read-based analyses, such as genotyping and derived frequency estimates of each site, must be interpreted with caution, given requirements for read support over each site. Continued improvements in sequencing technologies (longer read lengths) and costs will ameliorate such issues in the future. Such changes will also increase the feasibility of assembly-based approaches, permitting the direct reconstruction of full insertion, ultimately contributing to a more complete picture of all types of genomic variation.

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