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Highlights

RetroSnake is an end-toend pipeline for detection of HERV-K insertions

Modular and computationally efficient (~4 h per genome)

Easy setup and installation with Snakemake

Can be installed and used by users with limited computational experience

Kabiljo et al., iScience 25, 105289 November 18, 2022 © 2022 The Authors. https://doi.org/10.1016/ j.isci.2022.105289

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RetroSnake: A modular pipeline to detect human endogenous retroviruses in genome sequencing data

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SUMMARY

Human endogenous retroviruses (HERVs) integrated into the human genome as a result of ancient exogenous infections and currently comprise $\sim 8\%$ of our genome. The members of the most recently acquired HERV family, HERV-Ks, still retain the potential to produce viral molecules and have been linked to a wide range of diseases including cancer and neurodegeneration. Although a range of tools for HERV detection in NGS data exist, most of them lack wet lab validation and they do not cover all steps of the analysis. Here, we describe RetroSnake, an end-to-end, modular, computationally efficient, and customizable pipeline for the discovery of HERVs in short-read NGS data. RetroSnake is based on an extensively wet-lab validated protocol, it covers all steps of the analysis from raw data to the generation of annotated results presented as an interactive html file, and it is easy to use by life scientists without substantial computational training. Availability and implementation: The Pipeline and an extensive documentation are available on GitHub.

INTRODUCTION

Human endogenous retroviruses (HERVs) integrated into the human genome as a result of ancient exogenous infections which invaded the germ lines (Gifford and Tristem, 2003). Although they make up a striking portion of the human genome (8%), they are mostly inactive as a result of the accumulation of a large number of mutations, methylation, and histone modifications (Grandi and Tramontano, 2018). The most recent HERV family to integrate in this fashion was HERV-K, which has been linked to a wide range of diseases including cancer (Chen et al., 2019) and neurodegeneration (Dembny et al., 2020). However, characterizing the HERV genomic landscape is challenging: HERV sequences are thousands of bases long, they are polymorphic in humans, and present a high degree of sequence similarity. Currently, a number of bioinformatics tools for the identification of HERVs in next-generation sequencing (NGS) data exist, mostly based on the exploitation of split and discordant reads to reveal the presence of potential HERV insertions (Chen and Li, 2019; Gardner et al., 2017; Keane et al., 2013; Santander et al., 2017). However, most of these tools lack wet lab validation of their results and are not end-to-end as they do not cover all steps of the analysis. These factors greatly limit their use.

Running specialized tools for the detection of HERV insertions is usually only one step in the discovery process. Data commonly need to be pre-processed, while the output of tools is only a starting point for the downstream analysis necessary for a biological interpretation of the predicted insertions. One such tool, Retroseq (Keane et al., 2013), has been used as part of a custom protocol to characterize HERV-K insertions designed and adopted by Wildschutte and colleagues (Wildschutte et al., 2016). In their protocol, the authors refine the Retroseq results through insertion junction reconstruction and scanning of the reconstructed contigs for presence of repetitive elements of interest. Notably, their protocol has been extensively validated by PCR and capillary sequencing, and over the years their predicted insertions have become a *de facto* standard for non-reference HERV-K insertions. A custom implementation of that protocol has recently been benchmarked against similar tools and it produced the most reliable predictions (Bowles et al., 2022). However, an automatized implementation of this protocol is not available making its adoption in further studies challenging without substantial informatics development.



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1









Figure 1. Graphical overview of the full RetroSnake pipeline

CramToBam:conversion of cram files to bam files; retroseqDiscover and retroseqCall: two steps of retroSeq for detection of HERV-K insertions; *filterCalls*: quality filtering of predicted insertions based on flags produced by RetroSeq; verify: use RepeatMasker to verify the predicted HERV-K insertion in contigs assembled from reads surrounding the insertion site; markKnownFiltered, markKnownVerified: report all previously reported HERV-K predictions; *markNovelFiltered, markNovelFiltered*, to verified: report all HERV-K predictions which have not been reported previously; *annotateFiltered* and *annotateVerified*: use AnnotSV to annotate predictions with biologically meaningful information. See STAR Methods for more details about each step.

Here, we present RetroSnake, a comprehensive Snakemake pipeline for the detection of HERVs in shortread NGS data. RetroSnake covers all commonly needed steps, from the pre-processing of sequencing data in CRAM/BAM format, to the annotation of results with information from a range of biological databases, and includes Retroseq and the Wildshutte protocol, in a single, flexible, and modular pipeline that requires minimal setup and informatics skills.

RESULTS

RetroSnake pipeline architecture

RetroSnake uses a modern computational workflow management system, Snakemake (Koster and Rahmann, 2018), to combine all steps employed in the detection of transposable elements into a single, fast, and easy to use pipeline. The core steps of the RetroSnake pipeline are as follows: (i) file conversion to a BAM format required for Retroseq; (ii) running Retroseq; (iii) filtering the predicted insertions; (iv) further refining of the results through insertion junction reconstruction and scanning of the reconstructed contigs for presence of the target HERV; (v) comparing the predicted insertions with known insertions; (vi) annotation of the results with information from a range of biological databases (e.g. overlapping genes and regulatory elements, predicted functional effect); (vii) generation of an interactive html report.

The Snakemake workflows consist of rules that define how to create output files from input files. The workflow is inferred by dependencies between the rules that arise from one rule requiring an output file of another as input. Upon execution, Snakemake determines the combination of rules necessary to achieve a requested output. This combination of rules is a directed acyclic graph (DAG) of jobs. A graphical presentation of RetroSnake DAG is shown in Figure 1. Users can request many different levels and formats of output: e.g., a bed file with novel verified insertions, or an html file with annotated insertions that can

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chr1	79792629	79792630	chr1	38241392	38242392									
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chr6	32648035	32648041	chr17	44153867	44154867									
chr6	161270898	161270905	chr18	24507661	24508661	19_24011288_24011289_MEL_1	MEI	RPSAP58						
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All Filtered Retroseq Predictions

Annotated Predictions with Links to External Resources

Figure 2. Different output options

Predictions annotated with overlapping genes and regulatory elements with hyperlink. Genome version used is Hg19

be opened in any browser and displays hyperlink to other resources. The full list of options for RetroSnake output is in Table S1, and an illustrative figure of annotated insertions is shown in Figure 2.

RetroSnake utilizes the Conda package management system (https://conda.io) with its Bioconda channel (Gruning et al., 2018), which can be installed and updated easily with only a few shell commands. Through Conda, it is possible to define isolated software environments per rule. Upon execution of a workflow, Conda obtains and deploys the defined software packages. Many RetroSnake steps rely solely on software distributed via Conda, and as such, require no further installation. The only dependencies that require manual installations are RepeatMasker and AnnotSV. Detailed guidance for their deployment is provided on GitHub (https://github.com/KHP-Informatics/RetroSnake).

Results on 162 human short-read whole-genome sequencing samples

To illustrate the utility of RetroSnake, we applied it to the detection of HERV-Ks in a dataset of 162 human whole-genome sequencing samples from the MRC London Neurodegenerative Diseases Brain Bank based at the Institute of Psychiatry, Psychology & Neuroscience, King's College London. These samples were sequenced as part of Project MinE. The data are described in detail in previous publications (lacoangeli et al., 2019; lacoangeli et al., 2021; Project MinE ALS Sequencing Consortium, 2018). Out of 36 known non-reference HERV-K insertions (Table S2), our pipeline identified 19 in at least one subject. These insertions and their frequencies at different stringency levels of RetroSnake are in Table 1. For these 19 insertions, frequencies of the predictions were compared with frequencies previously reported for the same insertions in multiple publications, where available (Figure 3). Most HERV-K insertions of highest frequency have indeed been previously reported. RetroSnake also found 45 novel insertions when its most stringent verification level was used, 18 of which were found in at least two samples (Figure S1). Table S3 includes a list of all predicted novel insertions.

Lab validation

Nested PCR was used to validate one novel high frequency HERV-K detected by RetroSnake located approximately at chr6:153429801 of Hg19 (Figure S1), falling inside an intron of the *RGS17* gene. This was the novel insertion with the highest frequency in our dataset. It was detected in 24 samples, and we were able to retrieve a DNA sample for one of them. The lab validation protocol was applied to the sample predicted to contain the HERV-K insertion and two samples in which it was predicted to be absent. DNA was obtained from the MRC London Neurodegenerative Diseases Brain Bank at King's College London. The nested PCR protocol gave a specific product at the expected size in the sample predicted to have the HERV-K insertion (Figure 4). This product was not observed in the two samples in which RetroSnake did not predict the insertion. The PCR product sequence aligns to the correct chromosome 6 location

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Table 1. Known non-reference HERVK insertions and their prevalence in 162 subjects from Project MinE, as detected by our pipeline at three levels of stringency

	Filtorod	Filtered + verified	Filtered + verified	Gono (appointed	Overlapping Regulatory
Insertion	(% subjects)	(% subjects)	strict (% subjects)	by AnnotSV)	AnnotSV)
chr1:111802591-111802598	0.65	0.65	0.51		
chr4:9603239-9603245	0.93	0.93	0.67		DEFB131A; DRD5
chr5:64388439-64388446	0.06	0.06	0.06		
chr5:80442265-80442272	0.06	0.06	0.02	RASGRF2	
chr6:16004793-16004926	0.01	0.01	0.00		
chr6:32643383-32643537	0.03	0.02	0.02		
chr6:32648035-32648041	0.04	0.04	0.01		
chr6:161270898-161270905	0.72	0.72	0.66		
chr9:132205208-132205208	0.33	0.33	0.27		TOR1A (morbid); DOLK (morbid); NUP188 (morbid); NTMT1; LRRC8A; PTGES; ASB6; USP20; FNBP1; C9orf50; DOLPP1; TBC1D13; C9orf78; IER5L; PRRX2; SH3GLB2; GPR107; ENDOG; CRAT; TOR1B;
chr10:101016044-101016228	0.95	0.02	0.01		
chr12:44313656-44313662	0.12	0.12	0.08	TMEM117	
chr12:124066476-124066483	0.25	0.25	0.19	TMED2-DT	
chr13:90743182-90743189	0.14	0.14	0.09	LINC00559	
chr15:28430044-28430186	0.86	0.02	0.02	HERC2	
chr15:63374593-63374600	0.87	0.86	0.70		HERC1 (morbid); TPM1 (morbid); CA12 (morbid); USP3; FBXL22; RPS27L; RAB8B; APH1B; MIR190A; LACTB;
chr19:21841536-21841542	0.14	0.14	0.06		
chr19:22414303-22414381	0.40	0.04	0.02		
chr19:29855781-29855787	0.52	0.49	0.15	LOC284395	UQCRFS1 (morbid); C19orf12 (morbid); URI1; PLEKHF1; POP4; VSTM2B;
chr22:23852639-23852640	0.34	0.00	0.00		

Columns Gene and Overlapping Regulatory element are obtained by AnnotSV and KnotAnnotSV (Geoffroy et al., 2021; Geoffroy et al., 2018) step of the pipeline. Genome version used is Hg19.

up to base pair ~286 in the sequencing read; from this position, it then aligns well to LTR5_Hs (Figure S2), validating the predicted HERV-K integration site.

Computational efficiency

The whole pipeline took approximately 4 hours of CPU time per sample, with an additional 2 hours if CRAM to BAM conversion was needed. The CPUs used were Intel(R) Xeon(R) CPU E5-2660 v3 @ 2.60GHz. Full performance results for single sample processing are in Table 2. These statistics refer to the mean of three different samples.

DISCUSSION

RetroSnake is an efficient and comprehensive computational pipeline for the detection of transposable elements. The main advantages of RetroSnake over the other available tools are that it is end-to-end, based on an extensively wet-lab validated protocol for HERV-K detection, and developed within the Snakemake







Previously Reported HERVK Insertions found by RetroSnake Pipeline

Lee et al WGS500 TCGA #filter ed+verified strict filter ed+verified medium filter ed

Figure 3. Known non reference insertions and their frequency

Figures for Lee et al., WGS500 and TCGA are obtained from (Marchi et al., 2014). Filtered, filtered+verified medium and filtered+verified strict are frequencies obtained by running RetroSnake with different levels of stringency on 162 Project MinE samples. The genome version used was Hg19.

framework which provides computational efficiency, scalability, customizability, crash recovery features, and it is easy to use.

End-to-end

RetroSnake starts with an alignment file and encompasses all steps commonly needed for the characterization of mobile elements. It eliminates the need for postprocessing of predicted insertions that are commonly necessary for the other existing tools. These post processing steps include not only comparing the predictions with previously known ones but also annotation of the results with biologically relevant information from various databases.

Based on a wet-lab validated protocol

RetroSnake is based on a previously described and experimentally validated protocol (Keane et al., 2013; Wildschutte et al., 2016). The authors have used PCR and capillary sequencing and have validated the presence of 34 of the 36 candidate insertions in at least one individual predicted to have the insertion. Using the same protocol gives us confidence in the quality of predicted insertions. Furthermore, we have verified the presence or absence of one novel, high frequency insertion in one sample predicted to carry the insertion and two samples predicted not to carry it.

Efficiency

The Snakemake framework allows the analysis to scale from single samples on personal computers to multiple samples on HPC clusters. When run on a cluster, Snakemake takes care of submitting jobs in parallel in an optimized way. The full pipeline including CRAM to BAM conversion takes around 6 hours of CPU time.







Figure 4. Gel electrophoresis result for the nested PCR product There is a no-template control sample labeled N.T. Lanes labeled '-' are from samples not predicted to contain the insertion. The lane labeled '+' is predicted to have the insertion. The expected product size, given a HERV-K insertion, is denoted by the white arrow.

CRAM to BAM conversion is only performed if the alignment files in CRAM format are used. If data in BAM format are provided, the pipeline can process one sample in less than 4 hours, including verification and annotation; for the same task, other tools such as STEAK (Santander et al., 2017) and Ervcaller (Chen and Li, 2019) would take 7.5 and 14.5 CPU hours, respectively (Bowles et al., 2022).

Customizability

RetroSnake is a highly modular pipeline that allows for a fast integration of additional steps and the replacement of existing ones. For example, the HERV-K insertion detection step could be accomplished using another tool without affecting the rest of the pipeline. Other annotation rules can be added to complement the existing AnnotSV annotations. RetroSnake can be used for the detection of any transposable elements by providing their reference sequences.

Ease of use

RetroSnake can be installed and used even by users with limited computational experience. RetroSnake uses the package manager Conda and its Bioconda channel, which enables automatic installation of all tools provided by these channels. For example, in order to run Retroseq, filter the results and classify insertions into known and novel, no additional tools need to be manually installed as they are automatically obtained by Conda. For the verification and annotation steps, additional tools need to be installed and detailed instructions are available on GitHub. In order to run RetroSnake, only basic knowledge of the terminal usage is necessary. Detailed examples and instructions are provided on our GitHub.

Crash recovery

When Snakemake determines which rules are to be executed by building a DAG of jobs, it checks timestamps of input and output files. As long as a rule was properly executed and its input file is older than its output file, Snakemake will not attempt to regenerate the output. The rule will run only if the timestamp of the input file has changed, or if the output file is missing. If an error occurs during execution, the output up to the point of failure is preserved, and the user will not need to rerun the whole pipeline. Similarly, if the parameters need to be changed, only particular rules need to be rerun. Another advantage provided by the Snakemake framework is that in case of a failed rule, intermediate files created by that rule are removed by Snakemake before exiting. This prevents truncated files from being mistakenly used in downstream analysis, which is a problem occasionally encountered in bioinformatics pipelines.

Conclusions

In conclusion, we presented RetroSnake, a computationally efficient, customizable, easy to use, and comprehensive pipeline for detecting HERVs in short-read genome NGS data. RetroSnake presents important advantages with respect to other available tools. For example, it is the only pipeline based on an extensively wet-lab validated protocol (Wildschutte et al., 2016), and it is the most complete HERV detection pipeline, producing annotated insertions presented as an html file with hyperlink, easy enough to be used by life scientists without substantial computational training.

Limitations of the study

RetroSnake is able to predict insertion points of HERVs. However, the short-read sequencing technology does not enable it to characterise they sequence beyond the presence of the (LTR). HERV-K LTRs alone are almost 1kb long and a complete provirus can be up to 10 kb, while reads are 100–250 bp long. Therefore, unmapped parts of the insertion (e.g. viral protein coding genes) cannot be mapped uniquely, and our tool only detects the insertion point using the LTR reference. However, with the increased availability of

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Table 2. Running time on a single CPU, maximum resident set size of all tasks in job; a maximum virtual machine size, and the largest file written for the three steps of the pipeline applied to one genome

Step	Running time on 1 CPU	Maximum resident set size	Maximum virtual machine size	Largest file written
CRAM to BAM conversion	02:11:57	0.14G	0.72G	90G
Retroseq	03:33:37	0.17G	1.15G	3M
Verification	00:21:54	0.33G	1.43G	100M

All other steps have negligible CPU requirements. All jobs were run on A High-Performance Computing (HPC) cluster Rosalind, hosted by King's College London. A single CPU has been allocated for each step. Metrics was obtained using sacct command on slurm. These statistics are the mean of three different samples.

long-read sequencing data (read length >10,000 bp), it is possible to not only predict the insertion point but to also determine the entire sequence of the insertion. Another limitation is that RetroSnake does not provide the zygosity of predicted insertions.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- **RESOURCE AVAILABILITY**
 - Lead contact
 - Materials availability
 - O Data and code availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
 - Subject details
- METHOD DETAILS
 - Pipeline steps
 - Whole-genome sequencing
 - Lab validation

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.105289.

ACKNOWLEDGMENTS

UK Research and Innovation; Medical Research Council; South London and Maudsley NHS Foundation Trust; MND Scotland; Motor Neurone Disease Association; National Institute for Health Research; Spastic Paraplegia Foundation; Rosetrees Trust; Darby Rimmer MND Foundation. Funding for open access charge: UKRI. R.K. is funded by the MND Scotland. H.M. is supported by a GSK studentship and the KCL funded Centre for Doctoral Training (CDT) in Data-Driven Health. A.I. is funded by the Motor Neurone Disease Association and South London and Maudsley NHS Foundation Trust. This is an EU Joint Programme-Neurodegenerative Disease Research (JPND) project. The project is supported through the following funding organizations under the aegis of JPND-http://www.neurodegenerationresearch.eu/(United Kingdom, Medical Research Council MR/L501529/1 to A.A.-C., principal investigator [PI] and MR/R024804/1 to A.A.-C., PI]; Economic and Social Research Council ES/L008238/1 to A.A.-C. [co-PI]) and through the Motor Neurone Disease Association. This study represents independent research partly funded by the National Institute for Health Research (NIHR) Biomedical Research Center at South London and Maudsley NHS Foundation Trust and King's College London. The work leading up to this publication was funded by the European Community's Horizon 2020 Programme (H2020-PHC-2014-two-stage; grant 633413). A.A.K. is funded by ALS Association Milton Safenowitz Research Fellowship (grant number22-PDF-609.DOI:10.52546/ pc.gr.150909." title = "doi:DOI:10.52546/pc.gr.150909.">DOI:10.52546/pc.gr.150909.), The Motor Neurone Disease Association (MNDA) Fellowship (Al Khleifat/Oct21/975-799), The Darby Rimmer Foundation, and The NIHR Maudsley Biomedical Research Centre. H.B. his is funded by the National Institute for Health and Care Research (NIHR) Biomedical Research Center based at Guy's and St Thomas' NHS Foundation Trust and King's College London. C.R.B. is funded by MRC (MR/S000844/1). We acknowledge use of the research computing facility at King's College London, Rosalind (https://rosalind.kcl.ac.uk), which is delivered in partnership with the National Institute for Health Research (NIHR) Biomedical Research Centers at South London & Maudsley and Guy's & St. Thomas' NHS Foundation Trusts and part-funded by capital

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equipment grants from the Maudsley Charity (award 980) and Guy's and St Thomas' Charity (TR130505). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR, King's College London, or the Department of Health and Social Care.

AUTHOR CONTRIBUTIONS

R.K., H.B., and A.I. contributed to concept and design the study. R.K. implemented the software, run the analyses, and drafted the manuscript. H.M. and H.B. have tested the pipeline. H.B., C.R.B., and C.M.S. have designed and preformed the wet lab validation. A.I. supervised the study. A.A.-C. and A.I. raised the study funding. All authors contributed to the write-up.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: March 3, 2022 Revised: August 8, 2022 Accepted: October 4, 2022 Published: November 18, 2022

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and algorithms		
Snakemake	Koster and Rahmann (2018)	https://github.com/snakemake/snakemake
RetroSnake	This paper	https://doi.org/10.5281/zenodo.7050012, https://github.com/KHP-Informatics/RetroSnake
Oligonucleotides		
PCR primers CTCACTTCTCCCCCTTGTG TAACCAAATGTGCGGCTGCT TTTCAGAGAGCACGGGGTTG	This paper	N/A
Critical commercial assays		
Q5 High-fidelity 2X master-mix	New England Biolab	M0492S
pCR™Blunt II-TOPO™	Invitrogen	451,245
QIAprep Spin Miniprep	QIAGEN	27106X4

RESOURCE AVAILABILITY

Lead contact

Requests for further information should be directed to the lead contact, Alfredo lacoangeli (alfredo. iacoangeli@kcl.ac.uk).

Materials availability

This study did not generate any new materials.

Data and code availability

• Data

Sequencing data from ProjectMinE is available upon reasonable request to the lead contact. • Code

All original code has been deposited at https://github.com/KHP-Informatics/RetroSnake and is publicly available as of the date of publication. https://doi.org/10.5281/zenodo.7050012.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Subject details

RetroSnake has been applied for the detection of HERV-Ks in a dataset of 162 human whole genome sequencing samples from the MRC London Neurodegenerative Diseases Brain Bank based at the Institute of Psychiatry, Psychology & Neuroscience, King's College London. The dataset comprises 107 ALS cases (60 male, 47 female; age at death 64.49 ± 12.31 years), 54 controls (22 male and 32 female, age at death 76.31 ± 14.56 years), and one sample with missing phenotype information.

METHOD DETAILS

Pipeline steps

CRAMToBam

This step calls Samtools (Li et al., 2009) in order to convert CRAM files to BAM files. CRAM files are a compressed version of BAM files, and frequently, due to the large size of the files and memory constraints, files





are stored as CRAMs. This first, optional step does the conversion to BAM format which is necessary for the next step. If the BAM files are already present, this step is skipped.

Retroseq

Retroseq is executed with default parameters – 80 identity threshold which can adjusted. The fasta file containing the sequence of the repetitive element needed for the execution of Retroseq has been downloaded from RepeatMasker with query 'LTR5_Hs' and is available on our GitHub PAGE. Both Retroseq *discover* and *call* options have been used.

Filtering

Output of Retroseq call is a vcf file with predicted insertions and various quality flags characterising each prediction. For filtering of Retroseq results we have used the following combination of flags *fl* and *gq* from Retroseq output: For *fl* of 8, *gq* has to be 10 or more; for *fl* of 7 *gq* has to be 20 or more and for *fl* of 6, *gq* needs to be as high as 29. All predictions with *fl* lower than 6 are rejected.

Marking known and novel insertions

The compiled list we used as known non reference HERV-K insertions is in Table S2. This list has been assembled from literature (Lee et al., 2014; Marchi et al., 2014; Subramanian et al., 2011; Wildschutte et al., 2016).

We used BEDTools (Quinlan and Hall, 2010) to mark as known insertions all predictions within 500 bp of a known prediction. Novel insertions were defined as predicted insertions not found within 500bp of a previously reported insertion. All novel predicted insertions in all subjects were merged using BEDTools; the presence/absence of an insertion in each subject was then obtained by using BEDTools to check the overlap with any of the merged insertions.

Verification

The verification step is based on our interpretation of the verification described by (Wildschutte et al., 2016). All reads from the aligned BAM that overlap each predicted insertion point until 1000bp downstream of it are collected and assembled into contigs using CAP3 (Huang and Madan, 1999). The assembled contigs are then run through RepeatMasker using DFAM libraries of repetitive elements. Only hits with less than 10% substitutions in the matching region compared to the consensus, and with less than 3% of bases opposite a gap in either a query or a repeat sequence pass the initial filter. For medium level verification, we further request at least one contig to contain LTR5_Hs. For strict verification, we require the presence of multiple contigs with LTR5_Hs. If only a single contig has LTR5_Hs, we require that the contig has to be in the first half of list of contigs, as the contigs assembled by CAP3 have been sorted by quality. The verification level is easily controlled by parameter *verificationLevel*.

Annotation

AnnotSV and knotAnnotSV are run with default parameters, on either filtered or verified predictions. A custom annotation track with known mobile element insertions and their frequencies has been added and is automatically included. When invoked, the annotation step executes both AnnotSV which searches for elements overlapping the predicted insertions, and KnotAnnotSV which takes the annotations predicted by AnnotSV and renders them into an HTML annotation report with clickable links.

Whole-genome sequencing

Whole-genome sequencing (WGS) data were obtained from frozen human *postmortem* tissue taken from primary motor cortex of 162 individuals from Project MinE described in detail in (lacoangeli et al., 2019; Project MinE ALS Sequencing Consortium, 2018). Samples were sequenced on the Illumina Hiseq X platform, using PCR-free library preparation, 150 bp paired-end reads, with 30x coverage depth. The full list of IDs of subjects from Project MinE included in this study is in Table S4.

Lab validation

Nested PCR was used to validate one novel, high frequency HERV-K detected by RetroSnake located approximately at Chr6:153429801. The protocol was applied to one Project MinE sample predicted to contain the HERV-K insertion and two samples in which it was predicted to be absent. 50ng genomic





DNA was used for all PCR reactions with Q5 High-fidelity 2X master-mix (NEB) in the presence of 10pmol of primers for 30 amplification cycles. The first PCR step used flanking external chromosome 6 primers which were designed using PrimerBlast (forward: CTCACTTCTCCCCCTTGTG, reverse: TAACCAAATGT GCGGCTGCT). The PCR product was purified using the QIAquick protocol (Qiagen). This was followed by a second PCR reaction using the forward flanking primer and an internal primer derived from the HERV-K LTR5_Hs sequence (TTTCAGAGAGCACGGGGTTG). This protocol should therefore amplify the 5′ junction between genomic DNA (chromosome 6) and the predicted HERV-K LTR. The final PCR product was TOPO cloned in the pCR™Blunt II-TOPO™ vector (Invitrogen) for 30 min at 22°C, and the TOPO cloning product was introduced into DH10B *E. coli* by chemical transformation. Transformed bacteria were selected in presence of kanamycin and plasmid DNA was isolated from single colonies by miniprep (QIAprep Spin Miniprep, QIAGEN). The insert size in the plasmids was verified by EcoRI digest, and plasmids with the expected insert size (~350bp) were sequenced by Sanger sequencing (GENEWIZ). A chromosome 6 contig (NCBI nucleotide assession AL356963) and the HERV-K LTR sequence (Dfam DF0000558.4) were aligned to the sequencing read using MacVector 17 (MacVector). The sequencing read is representative of two independent bacterial clones from the patient's PCR reaction.