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A unique viral reservoir landscape in HIV-1 elite controllers

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Author contributions

Whole genome amplification, HIV-1 sequencing: CJ, XL, KBE, JMC, BR, KC, JEB

Integration site analysis of in-vivo infected cells: CJ, KBE

Integration site analysis of *in-vitro* infected cells: ES, ANE

RNA-Seq: SH, XS

ATAC-Seq: XS

Analysis of HIV-1 RNA transcripts: CJ, XL, KBE, JEB, MO

Bioinformatics analysis: CG

Viral outgrowth assays: JMC, SMYC, LNB, SES, JAV, RFS, JMS

Contribution of PBMC and tissue samples: MJP, RH, MS, JM, PDB, TWC, SGD, BDW

HLA class I typing: MC IPDA: GML, RFS, JMS

Data interpretation, analysis, presentation: CJ, XL, CG, ML, XGY

Preparation and writing of manuscript: CJ, XL, CG, ML, XGY

Critical review and edits to manuscript: CG, XS, KBE, RH, ANE, MC, SGD, RFS, BDW

Research idea and concept and study supervision: ML and XGY

Data availability statement

RNA-Seq and ATAC-Seq data have been deposited in a public repository (NCBI GEO, accession number GSE144334). Due to study participant confidentiality concerns, full-length viral sequencing data cannot be publicly released, but will be made available to investigators upon reasonable request and after signing a coded tissue agreement. The Los Alamos HIV Sequence Database Hypermut 2.0 and the Los Alamos HIV Immunology Database 2.0 are available at www.hiv.lanl.gov. The iMethyl database is available at http:// imethyl.iwate-megabank.org. ROADMAP epigenomic data are available at http://www.roadmapepigenomics.org.

ANE has received fees from ViiV Healthcare Co. within the past year for work unrelated to this project. All other authors declare that conflicts of interest do not exist.

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Untreated HIV-1-infected individuals who durably control HIV-1 replication below detection thresholds of commercial viral load assays (here termed "elite controllers", ECs) may represent the closest possible approximation to a natural cure of HIV-1 infection¹. Previous studies have linked elite HIV-1 control to specific variations of the human HLA class I gene locus², and to the presence of highly-functional cellular immune responses³ with stronger abilities to kill virally-infected cells³, target mutationally-constrained epitopes⁴ and limit viral escape⁵. Although the persistence of small, replication-competent proviral reservoirs has been documented in ECs^{6,7}, the characteristics and possible distinguishing features of reservoir cells in this specific group of individuals remain poorly defined.

To address this question, we applied FLIP-Seq (Full-Length Individual Proviral Sequencing)⁸ to profile the proviral reservoir landscape at single-genome resolution to a large cohort of ECs who maintained undetectable HIV-1 plasma viral loads for a median of 9 (range: 1–24) years based on commercially-available PCR assays. A reference cohort of HIV-1-infected individuals treated with suppressive antiretroviral therapy (ART) for a median of 9 (range: 2–19) years was recruited for comparative purposes (Extended Data Table 1). Collectively, our analysis of a large number of individual HIV-1 proviral genomes (n=1,385 from 64 ECs and n=2,388 from 41 ART-treated individuals) demonstrated that the median number of proviral amplification products (intact and defective) per person was significantly lower in ECs relative to ART-treated individuals (Fig. 1a). Frequencies of nearfull length, genome-intact proviral sequences (IPs) lacking defined lethal sequence defects were also markedly reduced in ECs, although their quantitative spectrum varied considerably (Fig. 1b). Of note, IPs made up a significantly larger proportion of all proviral sequences in ECs at both the cohort level (Fig. 1c) and the per study participant level (Fig. 1d), compared to ART-treated individuals; in four ECs, IPs accounted for 100% of detected proviral species. Intra-individual proviral sequence diversity, determined by pair-wise comparisons of all IPs within a given study participant, was smaller in ECs (Fig. 1e, Extended Data Fig. 1a). Interestingly, within IPs from ECs, optimal cytotoxic T lymphocyte (CTL) epitope sequences restricted by autologous HLA class I isotypes displayed more limited evidence of mutational escape (Fig. 1f, Extended Data Fig. 1c-f). These data suggest that IPs from ECs were seeded early in the disease process and persisted long-term.

For a deeper analysis of the proviral reservoir structure, we initially focused on two ECs in whom no IPs were observed in our initial analysis. In EC1, an individual who had

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maintained drug-free HIV-1 control for a recorded time of 12 years with only one documented viremia of 56 HIV-1 RNA copies/ml in 23 viral load tests spanning this period (Extended Data Fig. 2), escalating the number of analyzed PBMC to the limit of available cells yielded a single IP from a total of 1.02 billion PBMC; 21 defective proviruses, many of which belonged to a sequence-identical cluster, were also observed (Fig. 1g). In EC2, in whom only a single documented episode of 93 HIV-1 RNA copies/ml was noted in 39 viral load tests spanning >24 years of follow-up without ART exposure (Extended Data Fig. 2), we failed to detect even a single IP from more than 1.5 billion PBMC, while 19 defective proviral species, including near full-length sequences with lethal hypermutations, were observed, clearly documenting that this individual had been infected with HIV-1 in the past (Fig. 1g). Members of a sequence-identical cluster of defective proviral sequences with large deletions were noted in samples collected in 2009 and in 2019 from EC2, demonstrating a profound durability of a clonal population of cells harboring this sequence.

Moreover, a subsequent quantitative viral outgrowth assay (qVOA) with 340 million resting CD4⁺ T-cells isolated from approximately 1 billion PBMC (collected in 2019), and an additional qVOA involving 41 million total CD4⁺ T-cells isolated from 158.5 million PBMC (collected in 2009) did not retrieve a single replication-competent viral species. The recently-developed intact proviral DNA assay (IPDA) did not reveal evidence of IPs in 14 million resting CD4⁺ T-cells, while confirming the presence of defective HIV-1 DNA sequences (Extended Data Fig. 1b). In addition, an analysis of 7.72 million gut cells collected by colonoscopy from the rectum (2.08 million CD45⁺ mononuclear cells and 2.30 million CD45⁻ cells) and terminal ileum (1.99 million CD45⁺ mononuclear cells and 1.35 million CD45⁻ cells) by FLIP-Seq did not reveal any intact or defective proviruses in EC2.

To the authors' knowledge, the absence of IPs in such extremely large numbers of analyzed cells has only been documented in the "Berlin Patient" who underwent an allogeneic hematopoietic stem cell transplantation from a donor who was homozygous for CCR5 Δ 32 9 , which resulted in what is widely considered a sterilizing cure of HIV-1 infection. Indeed, in our hands, an analysis of 113 million available PBMC from the "Berlin Patient" (collected in 2017 and 2018) retrieved not a single intact or defective proviral sequence using FLIP-Seq (Fig. 1a–b). Although the logic of scientific discovery 10 will never allow us to confirm that EC2 has achieved a sterilizing cure of HIV-1 infection through natural immune-mediated mechanisms, it is noteworthy that we have failed to falsify this hypothesis, despite analyzing massive amounts of cells with a range of complementary, highly-sensitive detection techniques.

We next performed a phylogenetic analysis of all IPs obtained from 50 ECs and 37 ART-treated individuals. In both groups, we readily observed large clusters of sequences that were completely identical over entire analyzed viral genomes (Fig. 1h), strongly suggesting that they originate from clonally-expanded HIV-1-infected cells that pass on identical copies of IPs during cell divisions. The proportions of these clonally-expanded IPs were significantly higher in ECs compared to ART-treated individuals (Extended Data Fig. 1g–h). A number of these sequences were also retrieved from qVOA, documenting that these IPs are indeed fully replication-competent (Fig. 2–3).

For a detailed analysis of the viral reservoir landscape in ECs, we focused on eleven ECs (EC3-13) in whom large clusters of identical IPs were detected and from whom sufficient numbers of cells were available. In these ECs, we frequently observed oligoclonal, and sometimes almost monoclonal compositions of the entire intact proviral reservoir landscape (Figures 2-3, Extended Data Fig. 3). Notably, such a narrowly-focused viral reservoir configuration consisting of rather few distinct IPs but displaying relatively large expansions of identical IP clones is compatible with very low, if any, levels of ongoing viral replication in these ECs. This viral reservoir structure is atypical relative to the more diverse spectrum of IPs previously described in long-term ART-treated individuals^{8,11}. Instead, the viral reservoir landscape in EC3-13 is more reminiscent of the oligoclonal viral reservoir structure of IPs typically observed in chronic HTLV-1 infection, a retroviral disease characterized by deep proviral latency that limits active viral transcription and replication, such that viral propagation occurs almost exclusively by mitotic spread during clonal proliferation of infected T cells¹². Based on these considerations, we hypothesized that IPs from ECs maintain a state of deep, long-lasting latency, possibly due to chromosomal integration into genomic regions not permissive to active viral transcription.

To investigate chromosomal positions of IPs, we used MIP-Seq (Matched Integration site and Proviral Sequencing)¹³ to analyze integration sites (IS) in conjunction with corresponding proviral sequences. Briefly, proviral DNA was diluted to single genome levels, subjected to phi29-catalyzed whole-genome amplification, and subsequently exposed to FLIP-Seq⁸ and IS analysis using "integration site loop amplification" or ligationmediated PCR¹⁵. These experiments, performed in the eleven ECs (EC3–13), identified a total of 92 IS corresponding to IPs, of which 33 were associated with unique chromosomal locations (Supplementary Table 1). These IS of IPs were preferentially located in chromosomes 7, 17 and 19, and to a lesser extent in chromosomes 16 and 18 (Fig. 4a, Extended Data Fig. 5a). Consistent with previous studies 13 in which a total of 100 pairs of IPs and corresponding IS (n=73 IPs with unique IS) were analyzed in long-term ART-treated individuals, proviral species that displayed complete sequence identity shared the same IS, confirming their clonal origin. Notably, upstream HIV-1 long terminal repeat regions, which are not included in typical FLIP-Seq assays^{8,11} but were specifically amplified in these individuals, also displayed complete sequence identity within analyzed proviral clones (Extended Data Fig. 4).

Interestingly, IS analysis revealed that a significantly larger proportion of IPs from ECs were located in non-genic/pseudogenic regions, relative to IPs from long-term ART-treated individuals analyzed using the same approach 13 (45% vs. 17.8% of distinct IPs, respectively, p=0.0051; 40.2% vs. 13% of all IPs, respectively, p<0.0001), and in comparison to prior studies analyzing HIV-1 IS in ART-treated individuals 14,16 without distinguishing intact from defective proviruses (Fig 4b, Extended Data Fig. 5b). A closer investigation revealed that the non-genic IS of IPs from ECs were frequently positioned in or surrounded by centromeric satellite or microsatellite DNA (EC3–7, Fig. 2a–e), non-coding regions of the human genome that consist of dense heterochromatin "gene deserts" 17 that are typically disfavored for HIV-1 integration 18. Localization of proviral sequences in such centromeric satellite DNA has been associated with deep viral latency in functional viral reactivation studies 19,20 and was exquisitely rare 21 or entirely undetectable in prior investigations

involving ART-treated individuals¹³. In our study, the integration of IPs into centromeric satellite or microsatellite DNA was observed for a total of 8 unique IPs (24% of distinct IPs, 20.7% of all IPs) and occurred at least once in five (EC3–5, EC7–8; Fig. 2a–c, e; 3a) of the 11 ECs analyzed. Additionally, three IS of IPs were located in centromeric non-genic DNA surrounded by satellite DNA (EC3, EC6; Fig. 2a, d). Notably, as many as six different IS of IPs were located in or surrounded by centromeric satellite DNA in EC3 (Fig. 2a). In addition to this highly disproportionate overrepresentation of centromeric satellite DNA among IS of IPs from ECs, EC10 and EC13 harbored integrations of clonal IPs in a large non-genic region in proximity to non-centromeric micro-satellite DNA on chromosome 16 (Fig. 3c, f). Thus, in total, 39.4% of all 33 distinct IPs (32.6% of all 92 IPs) from ECs were located within or in proximity to satellite or microsatellite DNA.

Corresponding to the disproportionate enrichment of non-genic IS in ECs, we noted that the number of genic IS associated with IPs was significantly diminished in ECs, relative to ARTtreated individuals¹³. These genic IS were almost exclusively located in introns of genes that, in comparison to long-term ART-treated individuals, showed weaker transcriptional activity (Extended Data Fig. 7a) and displayed an opposite orientation relative to the harboring host gene in approximately 60% of all sites analyzed (Extended Data Fig. 7b-c). Genes encoding for members of the Zinc-Finger protein (ZNF) family, in particular for Krueppel-associated box domain-containing ZNF (KRAB-ZNF)²², accounted for 33% of all 18 genes harboring distinct IPs in ECs (corresponding to 49% of all 55 genic integration events of IPs), a notable enrichment relative to ART-treated individuals (Fig. 4c, Extended Data Fig. 5c). Of note, clonal IPs were frequently integrated into KRAB-ZNF genes located in defined regions of chromosome 19 (Ref. 23), which are extensively occupied by the heterochromatin proteins CBX1 and SUV39H1²⁴ and display highly distinct chromatin features, with profound enrichment for repressive chromatin marks that extensively cover the bodies of ZNF genes, but selectively spare the corresponding host transcriptional start sites (TSS)²⁴. Interestingly, a prior computational, genome-wide analysis of chromatin states based on a combinatorial evaluation of multiple different chromatin marks in their respective spatial context revealed that repetitive satellite DNA and ZNF genes share a common, highly distinct chromatin state (referred to as "ZNF genes & repeats")²⁵. When combined, IPs located either in satellite DNA or in ZNF genes represented >45% of all 33 independent IPs and >60% of all 92 IPs in ECs, proportions that were significantly increased relative to ARTtreated individuals (Fig. 4d, Extended Data Fig. 5d).

For a formal analysis of proviral IS positioning relative to active transcription units in the host DNA, we performed RNA-Seq-based gene expression profiling in autologous total CD4⁺ T-cells, as well as autologous central-memory (CM) and effector-memory (EM) CD4⁺ T-cell subsets which harbor the majority of all HIV-1 IS²⁶. These experiments demonstrated a significantly increased chromosomal distance between IS of IPs and the most proximal host TSS in ECs, relative to long-term ART-treated individuals¹³ (Fig. 4e). Simultaneously, we calculated the chromosomal distance between IS coordinates of IPs and accessible chromatin, as determined by genome-wide ATAC-Seq data obtained from autologous CD4⁺ T-cells. Although IS in satellite and microsatellite DNA were excluded from this analysis (and from the subsequent analysis using ChIP-Seq, Hi-C-Seq and methylation-Seq data; see below) due to the reduced ability to map next-generation sequencing reads to repetitive

genomic DNA regions²⁷, we noted that IS in cells from ECs were located at significantly increased distances to accessible chromatin, compared to those from ART-treated individuals¹³ (Fig. 4f). These differences were observed when clonal sequences were counted only once (Fig. 4e–f) but were also notable when all clonal sequences were considered individually (Extended Data Fig. 5e–f).

In a subsequent analysis, we calculated the number of DNA reads associated with defined epigenetic histone marks in the proximity to viral IS using ChIP-Seq data from primary memory CD4⁺ T-cells available from the ROADMAP Epigenomics Project²⁵. In comparison to ART-treated individuals ¹³, this analysis revealed a marked enrichment for the repressive histone features H3K9me3 (Chr19, Chr7) and/or a de-enrichment for the activating chromatin feature H3K4me1 (Chr19, Chr17) at IS of IPs from ECs (Fig. 4g); a trend for differential expression of additional activating and inhibitory chromatin modifications in proximity to IS of IPs from ECs and ART-treated individuals was also noted (Extended Data Fig. 6a-d). Furthermore, an alignment of IS coordinates to three-dimensional chromosomal contact data generated by Hi-C-Seq²⁸ demonstrated a significantly-increased proportion of IPs from ECs located in compartment B, containing mostly closed chromatin. This effect was particularly obvious for IS in KRAB-ZNF genes on Chromosome 19 in ECs, which were all located in subcompartment B4 (Fig. 4h, Extended Data Fig. 5g). This very small compartment (accounting for approximately 0.3% of the human genome) is known to harbor dense heterochromatin marks²⁸ and represents a highly atypical chromosomal IS location for HIV-1 in non-controller individuals ¹³. A profoundly-increased frequency of IPs from ECs in compartment B was also noted when Hi-C-Seq data from Jurkat cells²⁹ were used for alignment (Extended Data Fig. 6e-f).

Taking advantage of previously-published genome-wide bisulfite sequencing data in CD4⁺ T-cells³⁰, we observed that the frequency of hypermethylated (>90% methylation) cytosine residues was significantly higher in proximity to IPs from ECs, relative to IS of IPs from long-term ART-treated individuals¹³ (Fig. 4i). These data suggest that chromosomal regions more susceptible to DNA methyltransferases represent preferential sites for the long-term persistence of IPs in ECs, arguably because the integration into hypermethylated genomic DNA might facilitate deep latency of IPs and protect against immune-cell targeting. Given that closely-neighboring cytosine residues are likely to share the same methylation status³¹, these results raise the possibility that HIV-1 promoter methylation, previously shown to induce proviral HIV-1 silencing in *in-vitro* assays³², may contribute to durable transcriptional repression of IPs from ECs. The frequencies of IPs located in laminaassociated domains (LADs), genomic regions that interact with the inner nuclear membrane, contain mostly closed chromatin and represent a rare target for HIV-1 integration³³, were not significantly different between IPs from ECs and ART-treated individuals when clonal sequences were counted only once; however, a significant enrichment of IPs from ECs in LADs was noted when clonal IPs were counted as independent proviruses (Extended Data Fig. 7d-e).

Given that non-coding centromeric satellite DNA is a highly disfavored target site for HIV-1 integration ¹⁸, the disproportionately increased number of IS in satellite DNA described here is a particularly striking feature of ECs. Notably, ECs expressed normal mRNA levels of

LEDGF/p75 and CPSF6 (Extended Data Fig. 7f), host factors that interact directly with HIV-1 proteins to bias HIV-1 IS selection to active transcription units^{34,35}. Although protein levels of these molecules were not assessed, these results suggest that there is no increased susceptibility of centromeric satellite DNA to HIV-1 integration in ECs. To further address this, we infected CD4⁺ T-cells from n=12 ECs from our study cohort and n=9 HIV-1 negative healthy individuals with a GFP-encoding HIV-1 construct, followed by sorting of GFP⁺ and GFP⁻ CD4⁺ T-cells and a subsequent IS analysis. These experiments, retrieving >120,000 independent HIV-1 integration coordinates, demonstrated that IS in satellite DNA accounted for extremely low proportions of all integration events (0.04–0.06% in GFP⁺ and 0.11–0.12% in GFP⁻ CD4⁺ T-cells), irrespective of the analyzed study cohort (Extended Data Fig. 8a–b, Supplementary Table 2). Moreover, there was no evidence for preferential targeting of non-genic chromosomal regions or genes encoding for KRAB-ZNF proteins in *in-vitro* infected CD4⁺ T-cells from ECs (Extended Data Fig. 8b–c).

In conclusion, this work identifies a markedly distinct intact proviral reservoir landscape in PBMC from individuals with durable natural control of HIV-1, characterized by IS features highly suggestive of deep latency. For additional functional validation of this conclusion, we analyzed the frequency of HIV-1 RNA transcripts in ECs and ART-treated individuals; these additional experiments demonstrated that the number of HIV-1 RNA copies, normalized to the corresponding number of IPs, was significantly lower in ECs (Fig. 4j). As such, ECs seem to exemplify attributes of a "block and lock" mechanism³⁶ of viral control, defined by silencing of proviral gene expression through chromosomal integration into repressive chromatin locations³⁷. We propose that the distinct reservoir configuration in ECs is not related to altered IS preferences during acute infection in ECs, but instead represents the result of cell-mediated immune selection forces that preferentially eliminate proviral sequences more permissive to viral transcription, in a process that we suggest referring to as the "autologous shock and kill" mechanism. In contrast, less transcriptionally active proviral sequences with features of deep latency, leading to lower vulnerability to immune recognition, seem to persist long term. In very rare cases, such as in EC1 and EC2, such selection forces may have accomplished near total clearance of all IPs, raising the possibility that a sterilizing cure of HIV-1 infection can, at least in principle, spontaneously occur through natural, immune-mediated mechanisms. Future studies will be necessary to determine whether signs of immune-mediated selection pressure on viral reservoir cells are also visible in IPs from lymphoid tissues, which harbor the majority of viral reservoir cells³⁸.

While our data strongly suggest that deep latency plays a role in maintaining spontaneous, drug-free control of HIV-1 in some ECs, deep viral latency is not completely permanent or irreversible, as reflected by our ability to retrieve replication-competent virus from ECs in *in-vitro* outgrowth assays. However, *in-vitro* viral outgrowth assays with maximum stimuli are unlikely to adequately reflect susceptibility to viral reactivation *in vivo*; indeed, *in-vitro* viral outgrowth may largely be a stochastic process^{11,39}, and may occur independently of molecular pathways fine-tuning *in-vivo* viral outgrowth behavior. Nevertheless, it is likely that deep viral latency in ECs is a dynamic process, and that occasional bursts of viral transcription may occur despite genomic and epigenetic IS features restricting viral gene expression. In fact, a proviral landscape with low permissiveness to viral reactivation stimuli

may expose the immune system to a tailored viral antigen dose that can maintain a highly-functional antiviral T-cell response, a hallmark of antiviral immunity in ECs³, without supporting high-level viral replication promoting cytotoxic T-cell exhaustion. Therefore, a reciprocal equilibrium between a weakly-inducible viral reservoir and an efficient HIV-1-specific CD8+ T-cell response may represent the cornerstone of natural HIV-1 immune control. Given that evidence for selection of IPs with features of deeper latency was also observed in long-term ART-treated individuals, albeit at weaker degrees¹³, it is hoped that future longitudinal evaluations will be informative for designing strategies to induce a long-term drug-free remission of HIV-1 infection in larger populations of individuals.

Methods

Study participants

HIV-1-infected study participants were recruited at the Massachusetts General Hospital (MGH), the Brigham and Women's Hospital (BWH, both in Boston, MA, USA) and at the University of California, San Francisco (UCSF) at the Zuckerberg San Francisco General Hospital (San Francisco, CA, USA). PBMC and tissue samples were obtained according to protocols approved by the respective Institutional Review Boards. Clinical and demographical characteristics of study participants are summarized in Extended Data Table 1.

Droplet digital PCR

PBMC or CD4⁺ T-cells enriched from total PBMC using CD4 T Cell Isolation Kit (Miltenyi Biotec #130-096-533) were subjected to DNA extraction using commercial kits (Qiagen DNeasy #69504). We amplified total HIV-1 DNA using droplet digital PCR (Bio-Rad), using primers and probes described previously⁸ (127 bp 5'LTR-gag amplicon; HXB2 coordinates 684–810). PCR was performed using the following program: 95°C for 10 min, 45 cycles of 94°C for 30s and 60°C for 1 min, 72°C for 1 min. The droplets were subsequently read by the QX200 droplet reader and data were analyzed using QuantaSoft software (Bio-Rad).

Whole genome amplification

Extracted DNA was diluted to single viral genome levels according to ddPCR results, so that 1 provirus was present in approximately 20–30% of wells. Subsequently, DNA in each well was subjected to multiple displacement amplification (MDA) with phi29 polymerase (Qiagen REPLI-g Single Cell Kit #150345), per the manufacturer's protocol. Following this unbiased whole genome amplification⁴⁰, DNA from each well was split and separately subjected to viral sequencing and integration site analysis, as described below. If necessary, a second-round MDA reaction was performed to increase the amount of available DNA.

HIV near full-genome sequencing

DNA resulting from full-genome amplification reactions was subjected to HIV-1 near full-genome amplification using a 1-amplicon and/or non-multiplexed 5-amplicon approach, as described before ¹³. PCR products were visualized by agarose gel electrophoresis (Quantify One and ChemiDoc MP Image Lab, BioRad). All near full-length and/or 5-amplicon

positive amplicons were subjected to Illumina MiSeq sequencing at the MGH DNA Core facility. Resulting short reads were de novo assembled using Ultracycler v1.0 and aligned to HXB2 to identify large deleterious deletions (<8000bp of the amplicon aligned to HXB2), out-of-frame indels, premature/lethal stop codons, internal inversions, or packaging signal deletions (\(\geq 15\) bp insertions and/or deletions relative to HXB2), using an automated in-house pipeline written in Python programming language (https://github.com/BWH-Lichterfeld-Lab/Intactness-Pipeline)⁴¹, consistent with prior studies^{8,42,43}. Presence/absence of APOBEC-3G/3F-associated hypermutations was determined using the Los Alamos National Laboratory (LANL) HIV Sequence Database Hypermut 2.0⁴⁴ program. Viral sequences that lacked all mutations listed above were classified as "genome-intact" sequences. Sequence alignments were performed using MUSCLE⁴⁵. Phylogenetic distances between sequences were examined using maximum likelihood trees in MEGA (www.megasoftware.net) and MAFFT (https://mafft.cbrc.jp/alignment/software), and visualized using Highlighter plots (https://www.hiv.lanl.gov/content/sequence/HIGHLIGHT/highlighter_top.html). Viral sequences were considered clonal if they had completely identical consensus sequences; single nucleotide variations in primer binding sites were not considered for clonality analysis. Clades of intact HIV-1 proviral sequences were determined using the LANL HIV Sequence Database Recombinant Identification Program⁴⁶. Within intact HIV-1 clade B sequences, the proportions of optimal CTL epitopes (restricted by autologous HLA class I alleles) matching the clade B consensus sequence and CTL escape variants restricted by selected HLA class I alleles and supertypes described in the LANL HIV Immunology Database (www.hiv.lanl.gov) were determined.

Integration site analysis

Integration sites associated with each viral sequence were obtained using integration site loop amplification (ISLA), using a protocol previously described by Wagner et al¹⁴, or by ligation-mediated PCR¹⁵ (Lenti-XTM Integration Site Analysis Kit (Takara Bio #631263)); DNA produced by whole-genome amplification was used as template. For selected clonal sequences, viral-host junction regions were also amplified using primers annealing upstream of the integration site in host DNA and downstream of the integration site in viral DNA. Resulting PCR products were subjected to next-generation sequencing using Illumina MiSeq. MiSeq paired-end FASTQ files were demultiplexed; small reads (142 bp) were then aligned simultaneously to human reference genome GRCh38 and HIV-1 reference genome HXB2 using bwa-mem⁴⁷. Biocomputational identification of integration sites was performed according to previously-described procedures ^{14,48}: Briefly, chimeric reads containing both human and HIV-1 sequences were evaluated for mapping quality based on (i) HIV-1 coordinates mapping to the terminal nucleotides of the viral genome, (ii) absolute counts of chimeric reads, (iii) depth of sequencing coverage in the host genome adjacent to the viral integration site. The final list of integration sites and corresponding chromosomal annotations was obtained using Ensembl (v86, www.ensembl.org), the UCSC Genome Browser (www.genome.ucsc.edu) and GENCODE (v29, www.gencodegenes.org). Repetitive genomic sequences harboring HIV-1 integration sites were identified using RepeatMasker (www.repeatmasker.org).

Cell sorting and flow cytometry

PBMC were stained with monoclonal antibodies to CD4 (1:50, clone RPA-T4, Biolegend #300518), CD3 (1:50, clone OKT3, Biolegend #317332), CD45RO (1:40, clone UCHL1, Biolegend #304236) and CCR7 (1:40, clone G043H7, Biolegend #353216). Afterwards, cells were washed and CD45RO+ CCR7+ (central-memory) and CD45RO+ CCR7- (effector-memory) and CD3+ CD4+ (total) CD4+ T-cells were sorted in a specifically designated biosafety cabinet (Baker Hood), using a FACS Aria cell sorter (BD Biosciences) at 70 pounds per square inch. Cell sorting was performed by the Ragon Institute Imaging Core Facility at MGH and resulted in isolation of lymphocytes with the defined phenotypic characteristics of >95% purity. Data were analyzed using FlowJo software (Treestar).

RNA-Seq

Total RNA was extracted from sorted CD4⁺ T-cell populations using a PicoPure RNA Isolation Kit (Applied Biosystems #KIT0204). RNA-Seq libraries were generated as previously described⁴⁹. Briefly, whole transcriptome amplification (WTA) and tagmentation-based library preparation was performed using SMART-seq2, followed by sequencing on a NextSeq 500 Instrument (Illumina). The quantification of transcript abundance was conducted using RSEM software (v1.2.22) supported by STAR aligner software (STAR 2.5.1b) and aligned to the hg38 human genome. Transcripts per million (TPM) values were then normalized among all samples using the upper quantile normalization method.

ATAC-Seq

A previously-described protocol with some modifications^{50,51} was used. Briefly, 20,000 sorted cells were centrifuged at 1500 rpm for 10 min at 4°C in a pre-cooled fixed-angle centrifuge. All supernatant was removed and a modified transposase mixture (including 25 μl of 2x TD buffer, 1.5 μl of TDE1, 0.5 μl of 1% digitonin, 16.5 μl of PBS, 6.5 μl of nuclease-free water) was added to the cells and incubated in a heat block at 37°C for 30 min. Transposed DNA was purified using a ChIP DNA Clean & Concentrator Kit (Zymo Research #D5205) and eluted DNA fragments were used to amplify libraries. The libraries were quantified using an Agilent Bioanalyzer 2100 and the Q-QubitTM dsDNA High Sensitivity Assay Kit. All Fast-ATAC libraries were sequenced using paired-end, singleindex sequencing on a NextSeq 500/550 instrument with v2.5 Kits (75 Cycles). The quality of reads was assessed using FastQC (https://www.bioinformatics.babraham.ac.uk). Low quality DNA end fragments and sequencing adapters were trimmed using Trimmomatic (http://www.usadellab.org). Sequencing reads were then aligned to the human reference genome hg38 using a short-read aligner (Bowtie2, http://bowtie-bio.sourceforge.net/bowtie2/ index.shtml) with the non-default parameters "X2000", "non-mixed" and "non-discordant". Reads from mitochondrial DNA were removed using Samtools (http://www.htslib.org). Peak calls were made using MACS2 with the callpeak command (https://pypi.python.org/pypi/ MACS2), with a threshold for peak calling set to FDR-adjusted p<0.05.

Viral outgrowth assays

CD4⁺ T-cells were isolated from PBMC using EasySep™ Human CD4 Positive Selection Kit II (STEMCELL Technologies #17852). Cells were plated in limiting dilutions based on the intact provirus reservoir size determined through FLIP-Seq. Irradiated feeder PBMC were added at 1×10^5 cells/well. Cells were activated with 1 µg/mL PHA for four days, which was subsequently washed away and 10,000 MOLT-4 CCR5+ cells (NIH AIDS Reagent Program #4984) were added to propagate infection. On the thirteenth and twentieth days, culture supernatants from each well were individually incubated with 10,000 TZM-bl cells (NIH AIDS Reagent Program #8129) to drive Tat-dependent luciferase production. On the fifteenth and twenty-second days, TZM-bl cells were lysed, and luciferase activity was measured using Britelite Plus (PerkinElmer #6066761). Luciferase positive wells were defined as having signal levels >3 fold higher than negative controls. Cells from positive wells were then harvested and plated into lower compartments of Transwell tissue culture inserts (Costar® 6.5 mm Transwell®, 0.4 µm Pore Polyester Membrane Inserts, STEMCELL #38024), while 1×10^6 MOLT-4 cells were placed in upper compartments. After five additional days of culture, MOLT-4 cells from the upper wells were harvested, subjected to FLIP-Seq. Large scale quantitative viral outgrowth measurements on EC2 were performed by a similar standard method⁵² with a p24 ELISA assay used to detect outgrowth.

IPDA

The intact proviral DNA assay (IPDA) uses digital droplet PCR to quantitate proviruses lacking overt fatal defects, especially large deletions and hypermutation, and was performed as previously described⁵³.

In-vitro infection assays

CD4⁺ T-cells were stimulated in RPMI medium supplemented with 10% fetal calf serum, recombinant IL-2 (50 U/ml), and an anti-CD3/CD8 bispecific antibody (0.5 μ g/ul, NIH AIDS Reagent Program #12277). Cells were infected on day 5 with a GFP-encoding NL4–3 construct with a BAL-derived R5-tropic envelope⁵⁴ at a multiplicity of infection (MOI) of 0.1 for 4 h at 37°C. After 2 washes, cells were resuspended in medium and plated at 5 × 10⁵ cells/well in a 24-well plate. On day 5, GFP⁺ and GFP⁻ CD4⁺ T-cells were sorted. Cells were processed to DNA extraction and integration site analysis using ligation-mediated PCR according to a previously-described protocol⁴⁸.

Analysis of cell-associated HIV-1 RNA

Total cell-associated RNA and DNA was extracted in parallel from the same PBMC sample, using the GenElute RNA/DNA/Protein Purification Plus Kit (Sigma #RDP300) according to the manufacturer's protocol. RNA was reverse transcribed into cDNA using a polyadenylation-RT reaction⁵⁵ to efficiently detect HIV-1 RNA transcripts, followed by ddPCR-based amplification with primers and probes spanning the HIV-1 TAR region, as described before⁵⁵. Simultaneously, cell-associated DNA was subjected to ddPCR-based amplification of the RPP30 gene to determine cell counts in PBMC samples, using probes and primers described previously⁵⁶. HIV-1 RNA copies per million PBMC were normalized

to the corresponding number of intact proviruses per million PBMC (determined by FLIP-Seq).

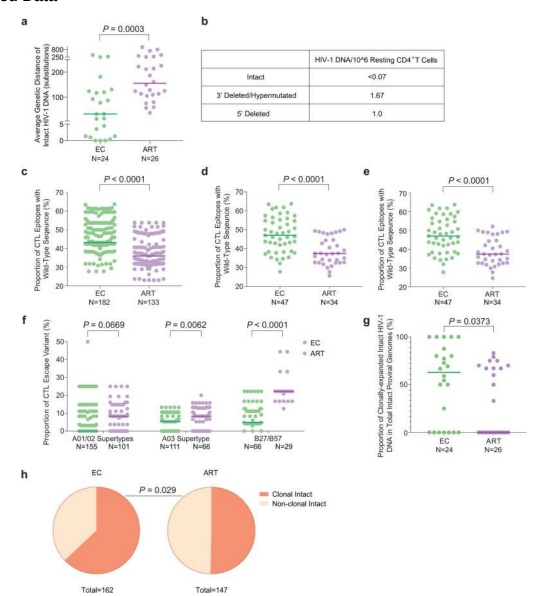
Statistics

Data are presented as pie charts, bar charts, scatter plots with individual values or heatmaps. Differences were tested for statistical significance using Mann-Whitney U test (two-tailed), Fisher's exact test (two-tailed), or Chi-squared test (two-tailed), as appropriate. p-values of <0.05 were considered significant, false discovery rate (FDR) correction was performed using the Benjamini-Hochberg method⁵⁷. Analyses were performed using Prism (GraphPad Software, Inc.), SPICE⁵⁸ and R (R Foundation for Statistical Computing⁵⁹).

Study approval

Study participants gave written informed consent to participate in accordance with the Declaration of Helsinki. The study was approved by the institutional review boards of MGH, BWH and UCSF.

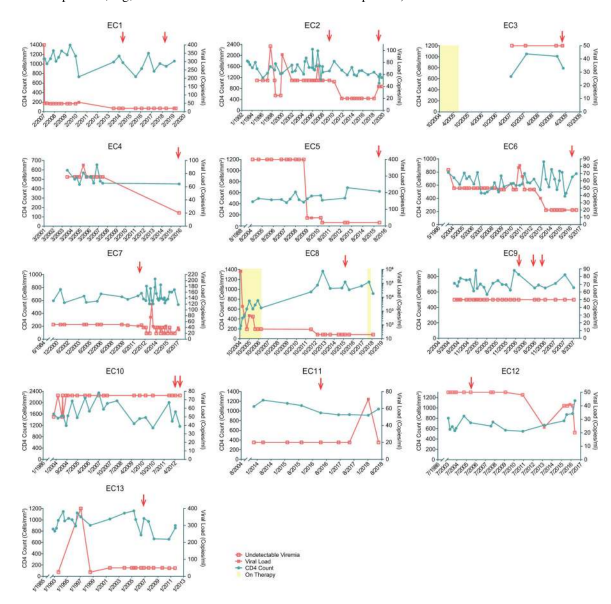
Extended Data



Extended Data Figure 1: Viral sequence analysis of intact HIV-1 proviruses from ECs.

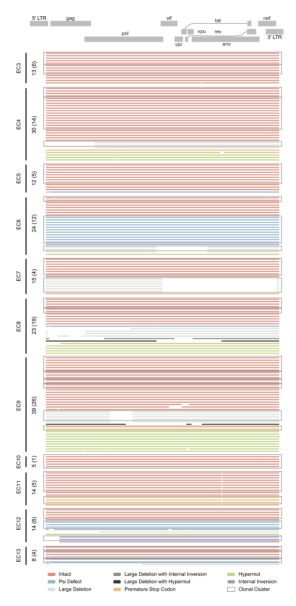
(a): Genetic distance (expressed as average number of base pair substitutions) among all intact near full-length proviral sequences obtained from each study participant. Clonal sequences were considered as individual sequences; participants with at least two intact proviruses are included (n=175 intact proviral sequences from 24 ECs and n=147 intact proviral sequences from 26 ART-treated individuals). (b): Frequencies of proviral species (copies per million resting CD4+ T-cells) detected by IPDA from EC2. (c): Proportion of optimal CTL epitopes (restricted by autologous HLA class I isotypes) with wild-type sequences. Each dot represents one intact proviral sequence. N=182 and N=133 HIV-1 clade B intact sequences from 47 ECs and 34 ART-treated individuals are included, respectively. Optimal CTL epitopes matching the clade B consensus sequences were considered as wild-type sequences. Clonal sequences were considered as individual sequences. (d-e): Average

frequencies of autologous HLA-restricted optimal CTL epitopes with wild-type sequences calculated from intact proviruses in each study participant. Clonally-expanded sequences were counted either once (d) or individually (e). Each dot represents one study participant. (f): Proportion of CTL escape variants (restricted by HLA-A01/A02 supertypes, HLA-A03 supertype, or HLA-B*27/B*57). Each dot represents one intact proviral sequence. Clonal sequences were counted individually. (g-h): Proportion of clonal intact proviruses among all intact proviruses within each study participant (g) or within all intact proviruses from ECs and ART-treated individuals(h). Study participants in whom at least two intact proviruses were detected are included in (g) and (h). (Two-tailed Mann Whitney U tests were used for panels a, c-g; two-sided Fisher's exact test was used for panel h).



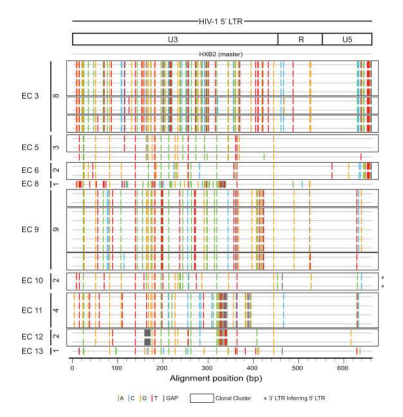
Extended Data Figure 2: Longitudinal evolution of CD4⁺ T-cell counts and HIV-1 viral loads in EC1-EC13.

The recorded diagnosis date of HIV-1 infection for each study participant is shown as the first date on x-axis. PBMC sampling time points are indicated by red arrows.

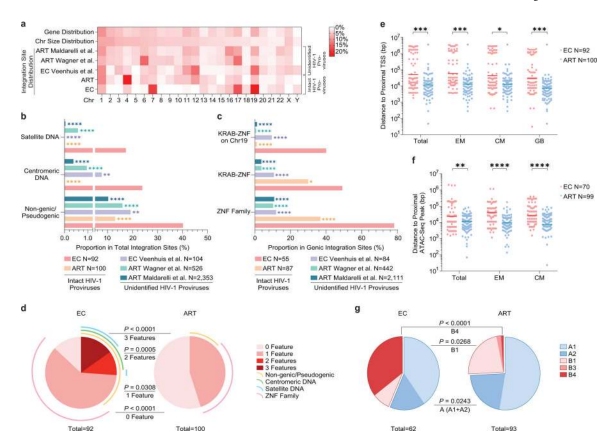


Extended Data Figure 3: Diagrams reflecting the structural composition of proviral reservoirs in ECs.

Virograms reflect the genetic coverage of individual sequences of proviral genomes analyzed in EC3-EC13. Numbers of total near full-length proviral sequences obtained from each individual are shown on the vertical axis; numbers of independent sequences are indicated in brackets. Open boxes indicate clonal clusters.



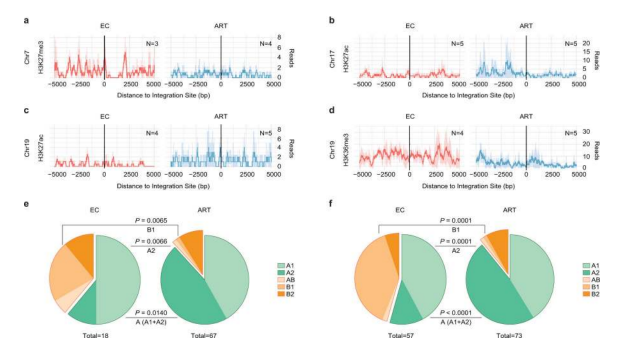
Extended Data Figure 4: Highlighter plot reflecting variations in HIV-1 DNA sequences in 5' LTR regions from intact proviruses isolated from indicated ECs, relative to HXB2. Numbers of 5' LTR sequences of intact proviruses obtained from each individual are shown on the vertical axis. Open boxes indicate clonal clusters.



Extended Data Figure 5: Chromosomal integration site features of intact proviruses from ECs after counting clonal sequences individually.

(a): Heatmap indicating the relative proportion of proviral integration sites of intact proviruses in each chromosome in ECs, relative to corresponding data from long-term ART treated individuals¹³. Proviral integration site data from prior publications are shown for comparative purposes (Veenhuis et al. ⁷, Maldarelli et al ¹⁶, Wagner et al. ¹⁴); integration sites from intact and defective proviruses were not distinguished in these studies. Contributions of each chromosome to total number of genes (first row) and to total size of human genome (second row) are included as references. (b-c): Proportion of near full-length intact proviruses located in indicated genomic regions. Data from near full-length intact proviral sequences in long-term ART-treated individuals (ART) are shown for reference purpose¹³; chromosomal integration sites from unselected (intact and defective) proviral sequences in ECs (Veenhuis et al.⁷) and in ART-treated individuals (Maldarelli et al.¹⁶, Wagner et al.¹⁴) are also shown for comparison. (d): SPICE diagrams⁵⁸ demonstrating proportion of intact proviruses with indicated chromosomal integration site features in ECs and ART-treated individuals. (e-f): Chromosomal distance between integration sites of intact proviruses and the most proximal transcriptional start sites (TSS, determined by RNA-Seq) (e) or to the most proximal ATAC-Seq peak (f) in autologous total, central-memory and effector-memory CD4⁺ T-cells and in GB. Horizontal lines reflect the geometric mean. (g): Proportions of proviral sequences located in structural compartments A and B, as determined based on Hi-C-Seq data published by Rao et al²⁸. Chromosomal integration regions not covered in the study by Rao et al. were excluded from analysis. (f-g): Sequences in genomic regions included in the blacklist for functional genomics analysis identified by the ENCODE and

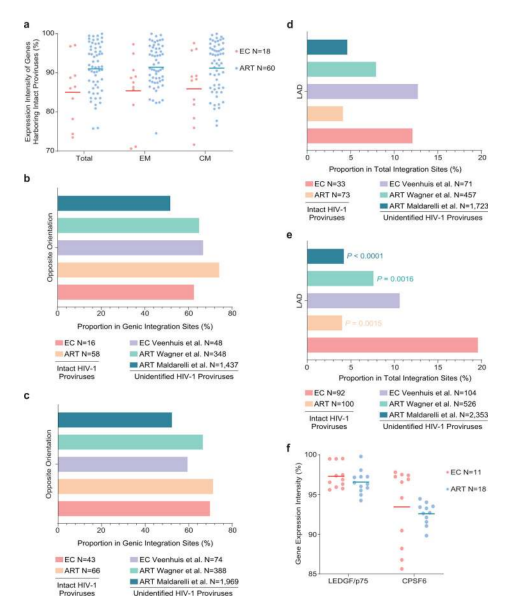
modENCODE consortia²⁷ were excluded due to absence of reliable ATAC-Seq and Hi-C-Seq reads in such repetitive regions. (a-g): All members of clonal clusters were included as individual sequences. (****p<0.001, ***p<0.001, **p<0.01, *p<0.05, FDR-adjusted two-sided Fisher's exact tests were used for panels b and c; two-sided Fisher's exact tests were used for panel d and g, FDR-adjusted two-tailed Mann Whitney U tests were used for panels e and f; all comparisons were made between ECs and reference groups).



Extended Data Figure 6: Epigenetic features of chromosomal integration sites of intact proviruses from ECs.

(a-d): Numbers of DNA sequencing reads associated with activating (H3K27ac) or repressive (H3K27me3) histone protein modifications in proximity to integration sites from ECs and long-term ART-treated individuals; median and confidence intervals (defined by one standard deviation) of ChIP-Seq data from primary memory CD4+ T-cells included in the ROADMAP repository²⁵ are shown. Negative distances indicate genomic regions upstream of the HIV 5' LTR host-viral junction, while positive distances indicate regions downstream of the 3' LTR viral-host junction. DNA sequencing reads associated with H3K36me3, an activating chromatin mark that is atypically enriched in KRAB-ZNF genes on Chromosome 19, are also shown²⁸. (e-f): Proportions of intact proviral sequences located in structural compartments A and B (and associated sub-compartments) by counting clonal sequences once (e) or by counting clonal sequences individually (f), as determined based on alignment of chromosomal integration sites of intact proviruses to Hi-C-Seq data from Jurkat cells²⁹. Chromosomal integration regions not covered in the Jurkat cell study were excluded from the analysis. Compartment B4 was not assessed in the source data²⁹ for this analysis. Two-sided Fisher's exact tests were used for statistical comparisons, nominal pvalues are reported. (a-f): Sequences in genomic regions included in the blacklist for functional genomics analysis identified by the ENCODE and modENCODE consortia²⁷

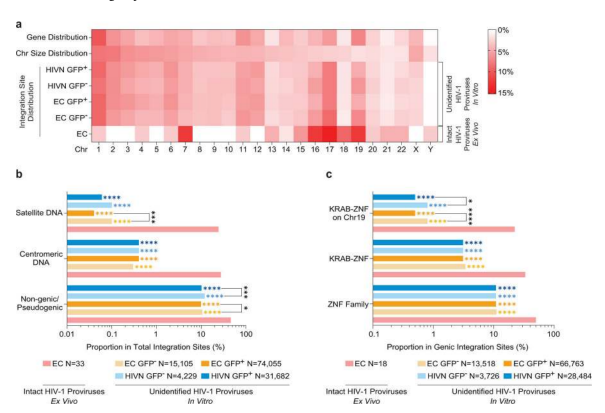
were excluded due to absence of reliable ChIP-Seq and Hi-C-Seq reads in such repetitive regions.



Extended Data Figure 7: Accessory chromosomal integration site features of intact proviral sequences from ECs.

(a): Expression of host genes harboring intact proviral sequences in ECs and long-term ART-treated individuals, as determined by autologous RNA-Seq data in total, central-memory (CM) and effector-memory (EM) CD4⁺ T-cells. Gene expression percentiles are indicated. (b-c): Orientation of intact proviruses relative to host genes in ECs and long-term ART-treated individuals. All data for genic integration sites with exclusive orientation towards host genes are included. Integration site data from prior studies involving ECs (Veenhuis et al.⁷) and ART-treated individuals (Maldarelli et al.¹⁶, Wagner et al.¹⁴) are shown for comparative purposes. (d-e): Proportion of intact proviruses from ECs and long-term ART-treated individuals in lamina-associated domains (LAD), determined using Lamin B1-DNA

adenine methyltransferase Identification (DamID) by Robson et al.⁶⁰ for resting Jurkat cells. Integration site data from prior studies involving ECs (Veenhuis et al.⁷) and ART-treated individuals (Maldarelli et al.¹⁶, Wagner et al.¹⁴) are shown for comparative purposes. (b, d): Clonal proviruses were counted once; (c, e): clonal proviruses were counted as individual sequences (FDR-adjusted two-sided Fisher's exact tests). (f): Expression of LEDGF/p75 and CPSF6 mRNA in autologous total CD4+ T-cells from ECs and long-term ART-treated individuals, as determined by RNA-Seq. Gene expression percentiles are indicated. (a, f): Horizontal lines reflect the geometric mean. All comparisons were made between ECs and reference groups.



Extended Data Figure 8: Chromosomal integration site features of *in-vitro* infected CD4⁺ T-cells from ECs and HIV-1 negative study participants.

(a): Heatmap indicating the relative proportion of proviral integration sites in sorted GFP +/GFP- *in-vitro* infected CD4+ T-cells (determined by LM-PCR⁴⁸) from ECs and HIV-1 negative study participants (HIVNs), relative to proviral integration sites of intact proviruses in each chromosome in ECs; integration sites from intact and defective proviruses were not distinguished in *in-vitro* infection studies. Data from GFP+ (n=74,055) and GFP- (n=15,105) CD4+ T-cell populations from ECs and from GFP+ (n=31,682) and GFP- (n=4,229) CD4+ T-cell populations from HIVNs were included. Contributions of each chromosome to total number of genes (first row) and to total size of human genome (second row) are included as references. (b-c): Proportion of proviral integration sites located in indicated genomic regions (b) or defined genes (c). Data from near full-length intact proviral sequences in ECs are indicated for reference. (****p<0.0001, ***p<0.001, *p<0.05, FDR-adjusted two-sided Fisher's exact tests or two-tailed Chi-square tests were used as

appropriate; p-values indicating comparisons made between ECs and each *in-vitro* infection group are shown in corresponding colors).

Extended Data Table 1:

Demographical and clinical characteristics of all study participants.

	Elite Controllers (EC)	ART-treated Participants (ART)
Number of participants	64	41
Age in years*	57 (31 – 75)	55 (34 – 73)
Female (%)	18.75%	21.95%
CD4 counts*	$908^{\dagger}(450 - 2,282)$	726 (316 – 1,649)
Viral loads	Under limit of detection	Under limit of detection
Number of viral load tests*	18 (3 – 91)	32.5 (4 – 73)
HLA-B*27/B*57 (%)	27.34% [‡]	8.75%
Time since diagnosis (year)*	17 (1 – 34)	17 (5 – 35)
Recorded duration of undetectable viremia (year) *	9 (1 – 24)	9 (2 – 19)

[,] median with range;

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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 $^{^{\}dagger}$, P= 0.0006, tested using two-tailed Mann-Whitney U test;

 $^{^{\}ddagger}$, P = 0.0012, tested using two-sided Fisher's exact test.

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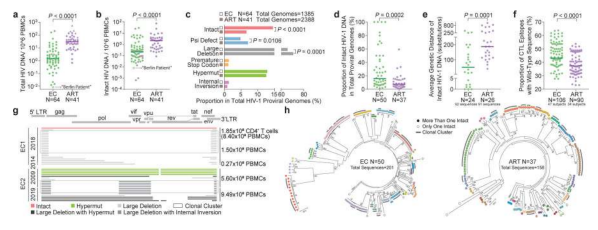


Figure 1: Proviral reservoir landscape in HIV-1 ECs.

(a-b): Relative frequencies of total (a) and near full-length intact (b) HIV-1 DNA sequences in ECs and ART-treated individuals (ART). Grey symbols: Limit of detection (expressed as 1 copy/total number of analyzed cells without target identification). Circles: Proviral sequences obtained from unfractionated PBMC; triangles: proviral sequences retrieved from isolated CD4⁺ T-cells and normalized to PBMC. (c): Proportions of proviral sequences that are genome-intact or display defined structural defects among all proviral genomes. (d): Proportion of IPs among all proviral genomes from each study participant. Only individuals with at least one detected IP are shown. (e): Average genetic distance between distinct IPs obtained from each study participant. Participants with at least two detectable IPs are included. (f): Proportion of optimal CTL epitopes (restricted by autologous HLA class I isotypes) with wild-type clade B consensus sequences. Each dot represents one IP. Clonal sequences are counted once. (g): Diagrams reflecting all proviral HIV-1 sequences isolated from EC1 and EC2. Left vertical axis: Dates of sample collection; right vertical axis: Numbers of cells analyzed. (h): Circular maximum-likelihood phylogenetic trees for all IPs from ECs and ART-treated individuals. Dots with the same colors indicate IPs detected in the same individuals. Clonal sequences, defined by complete sequence identity, are indicated by grey arches. Bootstrap analysis with 1000 replicates was performed to assign confidence to tree nodes; bootstrap support values >70% are shown in the trees. Two-tailed Mann Whitney U tests were used for panels a-b, d-f; False Discovery Rate (FDR)-adjusted twotailed Fisher's exact tests were used for panel c.

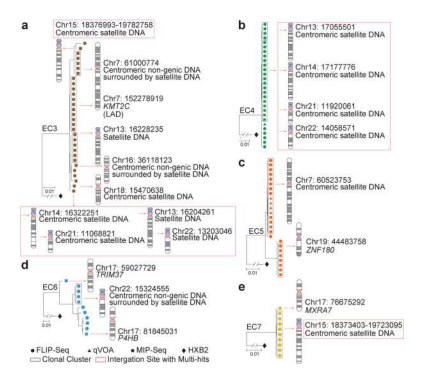


Figure 2: Increased frequency of IPs integrated in centromeric satellite DNA in ECs. (a-e): Data indicate linear maximum-likelihood phylogenetic trees for IPs from five ECs. Coordinates and relative positioning of IS are depicted; genes harboring IS are italicized. Clonal IPs, defined by identical proviral sequences and identical corresponding IS, are highlighted in curved black boxes. Red boxes reflect multi-hit IS that cannot be definitively mapped to one particular genomic location due to positioning in repetitive centromeric satellite DNA present in multiple regions of the human genome. LAD, lamina associated domain.

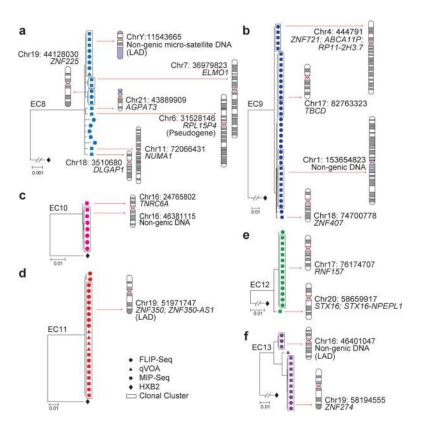


Figure 3: Preferential location of IPs from ECs in genes encoding for KRAB-ZNF proteins. (a-f): Linear maximum-likelihood phylogenetic trees demonstrate IPs from indicated study participants. Coordinates and relative positioning of IS are depicted. Other pertinent information is as defined in the legend to Figure 2.

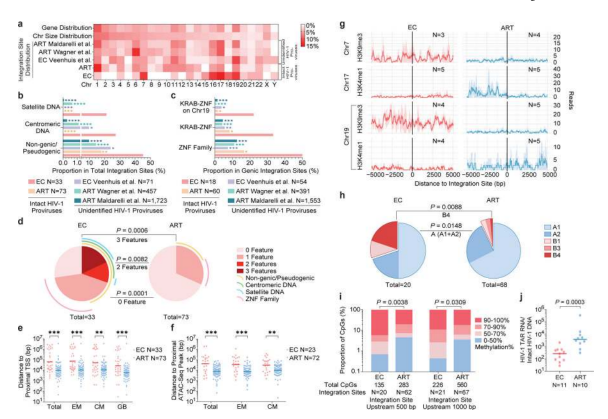


Figure 4: Distinct genomic and epigenetic features of IS of IPs from ECs.

(a): Relative proportion of proviral IS of IPs in each chromosome. Contributions of each chromosome to total number of genes (first row) and to total size of human genome (second row) are included as references. (b-c): Proportion of IPs located in indicated genomic regions. (a-c): Data from IPs in ART-treated individuals ¹³ (ART) and from unselected (intact and defective) proviral sequences in ECs⁷ and in ART-treated individuals ^{14,16} are shown as references. (d): SPICE diagrams demonstrating proportions of IPs with indicated IS features in ECs and ART-treated individuals. (e-f): Chromosomal distance between IS of IPs and the most proximal TSS in autologous total, EM or CM CD4+ T-cells or from Genome Browser (GB) (e), or to the most proximal ATAC-Seq peaks (f) in autologous total, EM and CM CD4⁺ T-cells. Horizontal lines reflect the geometric mean. (g): Numbers of DNA sequencing reads associated with activating (H3K4me1) or repressive (H3K9me3) histone protein modifications in proximity to IS from ECs and long-term ART-treated individuals¹³; median and confidence intervals (one standard deviation) of ChIP-Seq data from primary memory CD4⁺ T-cells included in the ROADMAP repository²⁵ are shown. (h): Proportions of IPs located in structural compartment A and B (and associated sub-compartments), as determined by Hi-C-Seq data²⁸. IS in regions not covered in ref.²⁸ were excluded. (i): Numbers of cytosine residues with indicated levels of methylation (derived from CD4⁺ Tcells in the iMethyl database³⁰) in proximity (500 or 1000 bp upstream of the 5' LTR hostviral junction) to IS from ECs and ART-treated individuals. (j): Frequencies of HIV-1 RNA transcripts in PBMC from ECs and ART-treated individuals, normalized to the corresponding number of IPs determined by FLIP-Seq. (a-i): Clonal sequences were only counted once. (f-i): Sequences in genomic regions included in the ENCODE blacklist²⁷ were

excluded. ****p<0.0001, ***p<0.001, **p<0.01, *p<0.05; (b/c/d/h): two-sided Fisher's exact tests; (e/f/j): two-sided Mann Whitney U tests; (i): two-tailed Chi-square test; (b/c/e/f/i): FDR-adjusted p-values; (d/h/j): nominal p-values. All comparisons were made between ECs and reference groups.