A quantitative approach for measuring the reservoir of latent HIV-1 proviruses

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A stable latent reservoir for HIV-1 in resting CD4+ T cells is the principal barrier to a cure1-2. Curative strategies that target the reservoir are being tested and require accurate, scalable reservoir assays. The reservoir was defined with quantitative viral outgrowth assays for cells that release infectious virus after one round of T cell activation1. However, these quantitative outgrowth assays and newer assays for cells that produce viral RNA after activation3 may underestimate the reservoir size because one round of activation does not induce all proviruses2. Many studies rely on simple assays based on polymerase chain reaction to detect proviral DNA regardless of transcriptional status, but the clinical relevance of these assays is unclear, as the vast majority of proviruses are defective7-9. Here we describe a more accurate method of measuring the HIV-1 reservoir that separately quantifies intact and defective proviruses. We show that the dynamics of cells that carry intact and defective proviruses are different in vitro and in vivo. These findings have implications for targeting the intact proviruses that are a barrier to curing HIV infection.

Evaluation of cure strategies requires assays that detect infected cells and distinguish intact proviruses from the vast excess of defective proviruses. We define ‘intact proviruses’ as those that lack overt fatal defects such as large deletions and hypermutation7-8, and recognize that some proviruses defined in this way may have minor defects that affect fitness. We analysed 431 near full genome HIV-1 sequences obtained by single genome analysis9 from 28 HIV-1-infected adults. The near full genome sequencing (nFGS) methods used identify defects throughout the genome except the 5′ long terminal repeat (LTR). Consistent with previous reports8,9, only 2.4% of proviruses were intact (Fig. 1a). The remaining 97.6% had fatal defects including deletions, encompassing on average 49.6% of the genome, and/or G→A hypermutation, which altered start codons and/or introduced stop codons in most open-reading frames (ORFs)8. Most defective proviruses had defects in most HIV-1 genes (Fig. 1b, Supplementary Table 1). Ninety-seven per cent of defective proviruses had defects that affected the transcriptional activator Tat (Fig. 1b, Supplementary Table 1) and might not be efficiently transcribed after latency reversal. Thus, cure interventions dependent on viral gene expression4,5 may affect cells with intact and defective proviruses differently. Hence, separate quantification is essential.

Standard PCR assays use short subgenomic amplicons in conserved regions (Fig. 1c, d) and do not distinguish intact and defective proviruses. Deletions occur throughout the genome (Fig. 1e), affecting not only the fraction of proviruses detected (Fig. 1f) but also the fraction of detected proviruses that are intact (Fig. 1g). Most standard PCR assays, less than 10% of detected sequences are intact (Fig. 1g). Thus, efficacy of a cure intervention causing a selective one log reduction in intact proviruses may be inappropriate (Fig. 1h).

Interrogating individual proviruses simultaneously at several positions could differentiate intact from defective proviruses. Analysis of nFGS data revealed that strategically placed amplicons in the packaging signal (Ψ) and env regions could jointly identify more than 90% of deleted proviruses as defective (Fig. 2a). Hypermutated proviruses must also be identified (Fig. 1a). Seventy-three per cent of these have mutations in the GG→AG context (Extended Data Fig. 1a). Most also have GA→AA mutations. Only 27% of hypermutated proviruses had only GA→AA mutations, and most of these also had deletions (Extended Data Fig. 1a). Therefore, we focused on GG→AG hypermutation. We identified a conserved region in the Rev-response element (RRE) with adjacent consensus sites (TGGG) for the responsible enzyme, APOBEC3G10 (Fig. 2b–e). Of the sequences with GG→AG hypermutation, 97% had one or more mutations in this region (Fig. 2c, e), with 13 distinct patterns (Fig. 2f). Using mutant plasmids carrying each pattern, we developed allelic discrimination probes (Extended Data Fig. 1b) that correctly identify 95% of hypermutated sequences as defective (Fig. 2f).

These analyses allowed design of a droplet digital PCR (ddPCR) method that distinguishes most deleted and/or hypermutated proviruses from intact proviruses using two amplicons and hypermutation discrimination probes (Fig. 3a, b). Genomic DNA (gDNA) is isolated using an optimized method to minimize DNA shearing between targeted regions (see below) and partitioned into nanodroplets such that individual droplets rarely contain more than one provirus (P = 0.00416). Proviruses within droplets are analysed simultaneously at the Ψ and env regions via multiplex PCR, with the env PCR also discriminating hypermutated proviruses. Intact proviruses give amplification at both regions (Fig. 3a, b). Intact proviruses per 106 cells are calculated using separate amplification of a cellular gene (RPP30) after correction for DNA shearing.

The ability of this intact proviral DNA assay (IPDA) to distinguish intact and defective proviruses was verified using plasmid controls that represent proviruses with different defects. These templates gave positive droplets in the expected quadrants: quadrant 1 (Q1) for 3′ deletion and/or hypermutation and Q4 for 5′ deletion (Extended Data Fig. 2). Cultured clonal populations of infected patient cells also give single quadrant patterns (see below). By contrast, uncultured polyclonal patient CD4+ T cell populations gave droplets in all four quadrants, allowing separate quantification of intact and defective proviruses (Fig. 3c).

Validation of the IPDA is described in Extended Data Figs. 3 and 4. DNA shearing is essential for droplet formation but artificially reduces

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double-positive (Q2) droplets while increasing Q1 and Q4 droplets. To control for shearing, two regions of a cellular gene (RPP30) with the same spacing as the $\Psi$ and env amplicons are amplified in a separate multiplex ddPCR on each sample. This gives a DNA shearing index (DSI) to correct raw HIV-1 ddPCR results (see Methods). To validate this approach, DNA from JLat cells, which contain a single provirus, was subjected to different amounts of shearing and analysed by ddPCR for HIV-1 and RPP30. At all levels of shearing, DSI was similar for HIV-1 and RPP30. Thus, RPP30 shearing can be used to correct for HIV-1 shearing (Fig. 3d). At all levels of shearing, corrected results were close to the expected values of one copy per cell for HIV-1 (Fig. 3e) and four copies per cell for RPP30 (JLat cells are tetraploid for RPP30). With correction for shearing, the IPDA gives highly accurate discrimination between molecularly defined standards representing intact and defective proviruses (Extended Data Fig. 5). In acute in vitro infections, high molecular mass DNA from productively infected CD4$^+$ T cells showed a substantial fraction of intact proviruses as assessed by IPDA and confirmed by nFGS (Extended Data Fig. 6). Thus, the assay can reliably quantify intact proviruses in most DNA samples after correction for shearing.

Using samples of only $5 \times 10^6$ cells, we measured intact and defective proviruses in 62 infected individuals who had suppression of viraemia on antiretroviral therapy (ART) ($n = 57$) or ‘elite’ control of HIV-1 without ART ($n = 5$) (Supplementary Table 3). The median DSI was 0.32 (interquartile range $\pm 0.07$; Extended Data Fig. 7), well within the range for which accurate correction is possible. Intact proviruses were rare (approximately 100 per $10^6$ resting CD4$^+$ T cells) and greatly outnumbered by defective proviruses (Fig. 3f). These results provide the first demonstration, to our knowledge, using a method not requiring long-distance PCR that the proviral landscape is dominated by defective proviruses. We found strong correlations between IPDA and quantitative viral outgrowth assay (QVOA) measurements on the same
samples ($r = 0.48$, $P = 0.003$, $n = 35$, Fig. 3g), even though the fraction of intact proviruses induced by a single round of T cell activation in the QVOA is small and variable between individuals (Fig. 3h).

Using nFGS data, we compared the IPDA with the widely used gag PCR with respect to desired assay characteristics. IPDA detects a larger fraction of proviruses (Fig. 3i) and separately enumerates intact and two class of defective proviruses. More importantly, the IPDA excludes 97% of proviruses with defects detectable by nFGS (Fig. 3i). The fraction of defective proviruses excluded by standard PCR assays is much lower and almost zero for Alt PCR assays. It is only 30% for gag PCR. Most proviruses (approximately 70%) classified as intact by IPDA lack defects detectable by nFGS. By contrast, this value is less than 10% for gag PCR (Fig. 3i). Thus, the IPDA provides a scalable alternative that is much more selective for intact proviruses. Most defects missed by IPDA are small deletions not overlapping IPDA amplicons. In addition, although nFGS data used to design the assay covers 97.6% of the non-redundant HIV-1 sequence, small deletions in the remaining portion of the genome may be present in some proviruses. The assay was designed for use in treated patients and does not distinguish integrated proviruses from unintegrated linear or circular forms. These are rare in patients on long-term ART. Further sequence data and improved ddPCR technology may allow even greater selectivity for intact proviruses. Importantly, selective intervention-induced reductions in intact proviruses would be apparent with the current IPDA but not with standard assays (Fig. 3i).

The latent reservoir measured by QVOA undergoes slow decay ($t_{1/2} = 44$ months)$^{12,13}$. For most patients, intact proviruses declined with a $t_{1/2}$ of approximately 44 months. Some patients showed even slower decay ($t_{1/2} = 100–300$ months), and for 3 out of 14 patients, there was no decay (Fig. 4b, c). Thus, the IPDA can measure changes consistent with known reservoir dynamics and possibly define subpopulations with slower decay.

The dynamics of defective proviruses in the same patients showed greater variability, with increases over time in some patients (Fig. 4a, c, Extended Data Fig. 8). Although the mean decay slopes did not differ, the standard deviations of decay slopes were much greater for defective proviruses (Extended Data Fig. 9), complicating interpretation of assays dominated by defective forms. Increases in infected cell frequency can reflect the proliferation of infected cells$^{14,19}$. Although the IPDA cannot demonstrate the presence of expanded clones, it can detect increases in infected cell frequency due to clonal expansion. We asked whether the differential dynamics of cells with intact and defective proviruses could be due to uneven distribution in subsets of memory CD4$^+$ T cells$^{20}$ with different proliferative potential. Intact and defective proviruses were found in expected ratios in central, transitional and effector memory subsets (Fig. 4d). We then asked whether intact and defective proviruses imposed different constraints on proliferation. We used the IPDA to track individual infected cells from treated patients after in vitro stimulation with anti-CD3/28 (Fig. 4e). Microcultures seeded with around one infected cell per well were subjected to four rounds of stimulation in the presence of antiretroviral drugs. Analysis of more than 1,700 microcultures showed that positive droplets were detected in only one out of three possible quadrants for most positive wells, as expected for cultures initiated with a single infected cell and cultured with antiretroviral drugs (Fig. 4e). Most (97.6%) proviruses were defective (Fig. 4f). Importantly, the IPDA also counts proviruses in each well, allowing the direct quantification of infected cell proliferation. Some cells carrying defective proviruses showed enormous clonal expansion (1,000-fold), whereas cells with intact proviruses were rarely detected and showed little proliferation (Fig. 4f). Despite fewer wells with intact
proviruses, differences in the mean number of intact and defective proviruses per well remained highly significant ($P = 0.0029$ to $P < 0.0001$ by unpaired $t$-test with Welch’s correction for intact proviruses versus proviruses with hypermutation and/or 3′ deletions). These differences are minimal estimates because the fraction of positive microcultures with intact proviruses was lower than expected based on IPDA analysis of the starting population (3.3% versus 7.5%), indicating that some cells with intact proviruses had hypermutation and/or 3′ deletions. These differences were confirmed by unpaired $t$-test for paired non-parametric values. e. Use of DSI to correct raw ddPCR output for RPP30 and HIV. Mean and s.d. of copies per cell of RPP30 (blue) and HIV (orange) are shown before (circles) and after (triangles) correction for shearing. f. IPDA results on CD4+ T cells from infected individuals ($n = 62$) with plasma HIV-1 RNA below the limit of detection. Data are geometric mean ± s.e.m. See Supplementary Table 3 for patient characteristics. Polymorphisms precluding amplification with either primer/probe set were not observed in this cohort and would require triage primer/probe sets incorporating rare polymorphisms. g. Correlation between infected cell frequencies measured by QVOA and IPDA on the same samples of CD4+ T cells from treated patients ($n = 36$). IUPM, infectious units per million cells. h. IPDA/QVOA ratios for samples from Fig. 3g. Horizontal bars indicate geometric mean and 95% confidence interval. i. Bioinformatic comparison of standard gag PCR and IPDA with respect to the percentage of proviruses amplified, percentage of defective proviruses excluded, percentage of amplified proviruses that are intact, and percentage loss in assay signal after a selective tenfold reduction in intact proviruses. Fn, fraction. All fractions expressed as percentages. See Fig. 1f–h for details.

Analysis of proviral integration sites of expanded clones demonstrated that the cultures were clonal and that integration into genes with cancer association was not required for proliferation (Fig. 4f, Supplementary Table 4). Instead, proliferative potential is related to proviral defects. nFGS confirmed the clonal nature of the proviruses in each well and demonstrated directly that the IPDA correctly identifies the presence and nature of defects in proviruses (Fig. 4g). Of 12 clonally expanded defective proviruses sequenced, 9 were defective in all HIV-1 ORFs, and none was fully competent for the expression of Tat or of Vif, Vpr, Vpu, Env or Nef. Low or absent expression of these genes would allow stimulated cells to escape viral cytopathic effects that could limit proliferation.

The failure of cells carrying intact proviruses to expand after repetitive in vitro T cell receptor stimulation provides insights into the mechanisms that drive in vivo clonal expansion and into cure strategies that involve T cell activation. The reservoir of intact proviruses undergoes slow decay in vivo, as shown here and previously. Thus, observed clonal expansions must be more than offset by the death of infected cells. Strong T cell receptor stimulation may induce viral gene expression and cell death or impaired proliferation as shown in our in vitro experiments. Clonal expansion in vivo may be driven by stimuli including homeostatic cytokines that allow proliferation without virus production.
Our results show that the small subset of proviruses with the potential to cause viral rebound show different dynamics than the vast excess of defective proviruses captured in standard PCR assays, emphasizing the importance of direct measurement of intact proviruses. The availability of a scalable assay for intact proviruses should accelerate cure research.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, statements of data availability and associated accession codes are available at https://doi.org/10.1038/s41586-019-0898-8.

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Competing interests Aspects of IPDA are subject of a patent application PCT/ US16/28822 filed by Johns Hopkins University. K.M.B. and R.F.S. are inventors on this application. Acclevir Diagnostics holds an exclusive license for this patent application. G.M.L. is an employee of and shareholder in Acclevir Diagnostics. R.F.S. holds no equity interest in Acclevir Diagnostics. R.F.S. is a consultant on cure-related HIV research for Merck and Abbvie.

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METHODS

Study participants. Characteristics of study participants are given in Supplementary Table 3. The Johns Hopkins Institutional Review Board and the UCSF Committee on Human Research approved this study. All participants provided written consent before enrolment. Except where indicated, participants were HIV-1-infected adults on suppressive ART with undetectable plasma HIV-1 RNA levels (<50 copies per ml) for more than 6 months. Chronic phase (CP)-treated subjects are defined as subjects starting ART more than 180 days from the estimated date of infection. Acute phase-treated subjects started ART less than 100 days after the estimated date of infection. For longitudinal analysis, additional peripheral blood mononuclear cell (PBMC) samples were obtained from 10 HIV-1-infected men followed in the Baltimore-Washington DC centre of the Multicentre AIDS Cohort Study (MACS) who had undetectable plasma HIV RNA (less than 20 copies per ml by Roche Taqman assay) at all semiannual study visits for 5 years or more, with no blips or missed visits. PBMC cryopreserved at visits at least 5 years apart, and viably stored as per MACS protocols, were studied. Characteristics of these 10 men are given in Supplementary Table 3 (CP31–CP49).

CD4+ T cell isolation. PBMCs were isolated by density centrifugation using Ficoll-Paque PLUS (GE Healthcare Life Sciences) per the manufacturer’s instructions. Untouched total CD4+ T cells were then enriched from PBMCs using negative immunomagnetic selection using the EasySep Human CD4+ T-Cell Enrichment Kit (StemCell Technologies). In some experiments, resting CD4+ T cells (CD4+, CD69+, CD25+ and HLA-DR−) were isolated using a second negative selection step (CD25−Biotin; Anti-Biotin MicroBeads; CD69 MicroBead Kit II; Anti–HLA-DR MicroBeads, all from Miltenyi Biotec). Resting CD4+ T cell purity was consistently greater than 95% as assessed using flow cytometry.

Bioinformatic analysis. We constructed an alignment of 431 published full-length proviral sequences obtained from 28 HIV-1-infected adults by single genome analysis. Of these individuals, 19 started suppressive ART during chronic infection, and 9 started suppressive ART during acute infection. Twenty-four individuals had prolonged suppression of viraemia (>6 months) when studied, and five were viraemic when studied (one individual was studied at two time points, before and after ART). Clinical characteristics of these individuals are given in the relevant publications. The proportion of different types of defects in HIV-1 proviruses varies with the stage of disease at which ART is started and the level of viraemia. For most analyses, we used sequences from patients who initiated ART during chronic infection and who had sustained suppression of viraemia to below 50 copies HIV-1 RNA per ml of plasma for >6 months. These sequences (n = 211) are thus representative of the most common group of infected individuals who would be eligible for cure interventions. The proviral landscape in patients starting ART during acute infection is also dominated by defective sequences, with a higher fraction showing hypermutation. In the analysis of deletions, common length polymorphisms were not included. Terminal deletions preclude integration and thus a large number of these deletions in the 5′ LTR are not available because the nFGS methods do not capture this region. Hypermutation occurred with or without deletions. For analyses of specific types of defect, we used all sequences in the database with that defect. For example, for analysis of hypermutation, we used all database sequences with hypermutation (n = 100) and additional hypermutated sequences obtained by single genome env sequencing. Hypermutation in the GG or GA context was confirmed using the Hypermut algorithm. Positions of primers for ddPCR analysis were evaluated using a sliding window analysis of two hypothetical 100-base-pair amplicons, scoring for lack of significant overlap (>5%) with mapped deletions in the database sequences with deletions. Optimal discrimination between intact and deleted sequences is obtained with a 5′ amplicon in the Ψ region and a 3′ amplicon in env. Ψ is the site of frequent small deletions and is included in larger 5′ deletions. Sequence conservation was evaluated using separate alignments of US clade B sequences from the Los Alamos HIV sequence database (https://www.hiv.lanl.gov/).

IPDA. A variety of experimental conditions were used in the development of the IPDA. The following is an optimized recommended procedure. In general, the IPDA is performed on DNA from 5 × 10⁶ CD4+ T cells. Genomic DNA is extracted using the QIAamp DNA Mini Kit (Qiagen) with precautions to avoid excess DNA fragmentation. DNA concentrations are determined using the Qubit3.0 and Qubit dsDNA BR Assay Kit (ThermoFisher Scientific). Quantification of intact, 5′ deleted, and 3′ deleted and/or hypermutated proviruses is carried out using primer/probe combinations optimized for subtype B HIV-1 (Accelver Diagnostics). The primer/probe mix consists of oligonucleotides for two independent hydropolymerase probe reactions that interrogate conserved regions of the HIV-1 genome to discriminate intact from defective proviruses (Supplementary Table 5). HIV-1 reaction A targets the packaging signal (Ψ) that is a frequent site of small deletions and is included in many large deletions in the proviral genome. The Ψ reaction A is positioned at HXB2 coordinates 692–797. This reaction uses forward and reverse primers, as well as a 5′ FAM-labelled probes: a 5′ VIC-labelled probe specific for wild-type proviral sequences, and a 5′ unlabelled probe specific for APOBEC-3G hypermutated proviral sequences (Supplementary Fig. 2). Successful amplification of HIV-1 reaction B produces a VIC fluorescence in droplets containing a wild-type form of RRE, detectable in channel 1 of the droplet reader, whereas droplets containing a hypermutated form of RRE are not fluorescent.

Droplets containing HIV-1 proviruses are scored as follows. Droplets positive for FAM fluorescence only, which arises from Ψ amplification, score as containing 3′ defective proviruses, with the defect attributable to either APOBEC-3G-mediated hypermethylation or 3′ deletion. Droplets positive for VIC fluorescence only, which arises from wild-type RRE amplification, score as containing 5′ defective proviruses, with the defect attributable to 5′ deletion. Droplets positive for both FAM and VIC fluorescence score as containing intact proviruses (Fig. 3a, b). Double-negative droplets contain no proviruses or rare proviruses (approximately 3.8%) with defects affecting both ampiclons. An important aspect of the quantification of intact proviruses is correction for DNA shearing between amplicons, which artificially reduces Q2 droplets while increasing Q1 and Q4 droplets (Fig. 3b). This can be done accurately through ddPCR analysis of a host gene, which also provides a measure of input cell number. In principle, any host gene can be used, with two ddPCR amplicons spaced at the same distance as the HIV-1 amplicons described above. As demonstrated in Fig. 3d, e, this approach allows for accurate correction for DNA shearing. Simultaneous quantification of DNA copy number and input genome equivalents is performed using another aliquot of the same DNA sample. For the studies described here, oligonucleotides for two independent hydrolysis probe reactions that interrogate the human RPP30 gene (chromosome 10: 90,880,081 on GRCh38) were used (Accelver Diagnostics). In this primer/mix, the total distance between duplex reactions and individual amplicon sizes are equivalent to those of the HIV-1 proviral discrimination reactions described above. Optimal reaction melting temperatures are equivalent for all reactions. Comprehensive oligonucleotide analysis predicted no potential off-target nucleotide binding or amplification (BLASTn) and no self-dimerization, hetero-dimerization, or primer hairpin formation. RPP30 reaction A uses a 5′ FAM-labelled hydrolysis probe, and RPP30 reaction B uses a 5′ HEX-labelled hydrolysis probe. Droplets positive for both 6-FAM and HEX fluorescence score as containing an unsheared RPP30 gene fragment, whereas droplets positive for only a single fluorescence score as containing part of a sheared RPP30 gene fragment. The ratio of dual fluorescent to single fluorescent droplets is used to calculate a DSI, and this index is applied to both genome copy number input reactions and HIV-1 proviral discrimination reactions to correct for experimentally observed DNA shearing.

The Bio-Rad QX200 AutoDG Digital Droplet PCR system using the appropriate manufacturer supplied consumables and the ddPCR Supermix for probes (no dUTPS) (Bio-Rad Laboratories). For HIV-1 proviral discrimination reactions, 700 ng of genomic DNA was analysed in each well. For DNA shearing and copy number reference reactions, 7 ng of genomic DNA was analysed in each reaction well. Multiple replicate wells were performed for each reaction type to ensure consistent quantification, and replicate wells were merged during analysis to increase IPDA dynamic range. The thermal cycling program used for all reactions, with a 2 °C ramp rate, is given in Supplementary Table 6. In general, parallel processing and analysis of uninfected donor CD4+ T cells performed using another aliquot of the same DNA sample. For the studies described here, oligonucleotides were used to calculate a DSI, and this index is applied to both genome copy number input reactions and HIV-1 proviral discrimination reactions to correct for experimentally observed DNA shearing.

QVOA. The QVOAs were performed as previously described, MOLT-4/CCR5 cells and CD4+ T cells were added on day 2 of the culture and the culture supernatants were examined for the p24 viral capsid protein by ELISA (PerkinElmer) after 14 and 21 days. For the studies described here, QVOAs were performed as previously described, MOLT-4/CCR5 cells and CD4+ T cells were added on day 2 of the culture and the culture supernatants were examined for the p24 viral capsid protein by ELISA (PerkinElmer) after 14 and 21 days. Results were expressed as infectious units per million cells (IUPM) CD4+ T cells calculated using maximal likelihood as described by previously (UIMPStats). T cell subset analysis. Resting CD4+ T cells from HIV-1-infected donors on suppressive ART were isolated as described above. To sort resting memory subsets, we incubated cells with FcεR block (BD Pharmingen) for 10 min before staining with a FITC-labelled antibody to CD3 (Biologen; clone HT13A), phycoerythrin (PE)-labelled antibody to CD4 (Biologen; clone RPA-T4), allophycocyanin (APC)-labelled antibody to CD45RO (Biologen; clone UCHL1), BV421-labelled antibody to CD27 (Biologen; clone O323) and PE-labelled antibody to CCR7 (Biologen; clone G043H7). Dead cells were excluded using propidium iodide.

Cell lines. The JLaT0 full-length clone (clone 6.3) from E. Verdin was obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH. Cells were authenticated through short tandem repeat analysis and tested negative for mycoplasma.

VQA. The VQAAs were performed as previously described, MOLT-4/CCR5 cells and CD4+ T cells were added on day 2 of the culture and the culture supernatants were examined for the p24 viral capsid protein by ELISA (PerkinElmer) after 14 and 21 days. Results were expressed as infectious units per million cells (IUPM) CD4+ T cells calculated using maximal likelihood as described by previously (UIMPStats). T cell subset analysis. Resting CD4+ T cells from HIV-1-infected donors on suppressive ART were isolated as described above. To sort resting memory subsets, we incubated cells with FcεR block (BD Pharmingen) for 10 min before staining with a FITC-labelled antibody to CD3 (Biologen; clone HT13A), phycoerythrin (PE)-labelled antibody to CD4 (Biologen; clone RPA-T4), allophycocyanin (APC)-labelled antibody to CD45RO (Biologen; clone UCHL1), BV421-labelled antibody to CD27 (Biologen; clone O323) and PE-labelled antibody to CCR7 (Biologen; clone G043H7). Dead cells were excluded using propidium iodide.
PE mouse IgG2α, BV421 mouse IgG1κ, and APC IgG2ακ isotype antibodies were used in fluorescence-minus-one controls to set sorting gates. Memory cells were distinguished from naïve cells by the presence or absence of CD45RO staining, respectively. Central memory cells were distinguished from effector and transitional memory subsets by the presence of CCR7 as described previously32. CCR7- cells were subdivided into effector memory (TEM) (CD45RO+CCR7-), or transitional memory (TM) (CD45RO+CCR7-CD27+) cells as described previously20. Central, effector and transitional resting memory subsets were sorted using a Beckman Coulter MoFlo XDT Cell sorter.

Clonal microcultures. Purified resting CD4+ T cells from HIV-1-infected donors on suppressive ART were analysed for proviral DNA copies using gag quantitative PCR as previously described33. The resulting values were corrected for gag- proviruses, and cells were plated at approximately one infected cell per well (2,000–4,000 total resting CD4+ T cells) in 96-well plates. The cells were then stimulated with anti-CD3/CD28 Dynabeads (25 μl per million cells; Thermo Fisher Scientific) in RPMI containing 10% fetal bovine serum and 100 U ml-1 IL-2 (Novartis) for 7 days in the presence of tenofovir disoproxil fumarate (10 μM) and emtricitabine (10 μM). Half of the medium in each well was removed and replaced with fresh medium, anti-CD3/CD28 Dynabeads and antiretroviral drugs weekly for 2–3 weeks. DNA isolation was performed on cells from each well using a Quick-DNA 96 Kit (Zymo Research Corporation). One-quarter of the extracted DNA was analysed by the IPDA to determine the type of provirus and proviral copy number in each well. The remaining DNA was used for integration site analysis and full genome sequencing.

Integration site analysis. Integration site analysis was performed using previously described linker ligation method26,34,35. Sites for which both the 5’ and 3’ junctions were captured are shown. Cancer associations were determined as previously described35.

Full genome sequencing. Full genome sequencing was performed at the single molecule level as previously described8.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability
The IPDA was developed through an analysis of published near full genome HIV-1 sequences (refs8,9, GenBank accession numbers KX505390–KX505744 and KU677989–KU678196, respectively). All other data are available from the corresponding author upon reasonable request.

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Extended Data Fig. 1 | Analysis of hypermutation. a, Most hypermutated proviruses show both GG→AG and GA→AA patterns of hypermutation. Analysis based on hypermutated full-genome sequences in the database (n = 100). Sequences were analysed for GG→AG and GA→AA patterns using all available sequences for each clone and the Los Alamos Hypermut algorithm\(^\text{17}\). b, Hypermutation discrimination using two probes in the RRE of the env gene. The intact probe hybridizes with a region containing two adjacent APOBEC3G consensus sites (red underline) in intact proviruses. It is labelled with a fluorophore (VIC) and a quencher (Q). Also present in the reaction is a hypermutated probe that lacks the fluorophore and does not bind (arrow) to intact proviral sequences owing to G→A mutations at both APOBEC3G consensus sites. Dashed boxes indicate the nucleotide positions of sequence differences between the intact and hypermutated probes. The hypermutated probe preferentially binds to the same region in hypermutated proviruses. It lacks a fluorophore and prevents binding of the fluorophore-labelled intact probe (arrow). Therefore, no fluorescent signal is generated for 95% of hypermutated proviruses (Fig. 2f).
**Extended Data Fig. 2** | Plasmid controls show the specificity of the IPDA. 

**a**, Maps of proviral plasmid control templates. Plasmids E44E1, 39G2 and 4F11 have deletions in the indicated regions (white). Plasmid 2G10 is a heavily hypermutated patient-derived sequence with G→A mutations in the probe-binding region of the env amplicon (enlarged region). Plasmid 19B3 has GA point mutations in this region. These G→A mutations (red) occur at two APOBEC3G consensus sites (TGGG, underlined) in this region. These plasmids have been previously described12,34.

**b**–**e**, IPDA on plasmids representing the indicated defective proviruses showing positive droplets only in the expected quadrants.
Extended Data Fig. 3 | IPDA accuracy, reproducibility and limit of quantification. 

**a**, Correlation between expected and IPDA-measured frequencies of intact proviruses per 10^6 cells. Genomic DNA from uninfected donor CD4^+ T cells was spiked with JLat6.3 DNA cell equivalents and subjected to a serial fourfold dilution. This material was then analysed by the IPDA, and the IPDA-measured frequencies of intact proviruses per million cells were compared to the expected frequencies (998, 249.5, 62.4, 15.6 and 3.9 intact proviruses per 10^6 CD4^+ T cells). This experiment was performed independently three times. The agreement between the expected and IPDA-measured frequencies of intact proviruses was determined using Pearson correlation.

**b**, Reproducibility of the IPDA across independent assay runs. Reproducibility was assessed by determining the coefficient of variation (CV) across three independent assay measurements of genomic DNA from uninfected donor CD4^+ T cells spiked with JLat6.3 cell equivalents and subject to serial fourfold dilution, as described in **a**.

### Table

| Expected intact proviruses per 10^6 CD4^+ T cells | Intact Proviruses per 10^6 input CD4^+ T-cells by IPDA |
|-------------------------------------------------|------------------------------------------------------|
|        | Run 1 | Run 2 | Run 3 | Average | StDev | % CV |
| 998.0  | 836.4 | 953.8 | 925.1 | 905.1   | 61.2  | 7    |
| 249.5  | 200.7 | 190.2 | 206.0 | 199.0   | 8.0   | 4    |
| 62.4   | 64.0  | 72.3  | 57.5  | 64.6    | 7.4   | 12   |
| 15.6   | 17.6  | 19.9  | 16.0  | 17.8    | 1.9   | 11   |
| 3.9    | 2.7   | 3.8   | 3.8   | 3.4     | 0.6   | 18   |

**Replicate 1**

Pearson r = 0.9998

R^2 = 0.9996

P < 0.0001

**Replicate 2**

Pearson r = 0.9982

R^2 = 0.9964

P < 0.0001

**Replicate 3**

Pearson r = 0.9996

R^2 = 0.9999

P < 0.0001
Extended Data Fig. 4 | IPDA reproducibility. a–c. Frequencies of cells containing proviruses with 3′ deletions and/or hypermutation (a), 5′ deletions (b), or no defects (intact; c) in CD4+ T cells from 28 treated patients. Each data point represents a replicate IPDA determination from a single sample from the indicated patient. The mean and s.e.m. of the replicates are plotted. The variability between patients is much greater than the variation between replicates from a single patient. Technical replicates are shown to indicate low intrinsic variability of the IPDA.
Extended Data Fig. 5 | Plasmid controls confirm specificity of the IPDA. a, Map of the plasmid pNL4-3 carrying an intact HIV-1 provirus. Positions of the Ψ (blue) and env (green) IPDA amplicons and of a distinct set of plasmid shearing control (PSC, magenta boxes) amplicons are indicated. Spacing between PSC amplicons is equal to spacing between Ψ and env amplicons. Dotted lines indicate positions of deletions in plasmids carrying previously described defective proviruses E44E11 and 4F12 with 5′ and 3′ deletions, respectively. b, IPDA analysis of the indicated ZrAl-cut plasmids representing intact, 5′-deleted and 3′-deleted proviruses. c, Summary of droplet counts for the experiment shown in b. E44E11 and 4F12 give positive droplets only in quadrants 4 and 1 (Q4 and Q1), respectively. For pNL4-3, more than 95% of droplets are in Q2, with the remainder attributable to shearing between the Ψ and env amplicons. d, Analysis of shearing. For IPDA analysis of patient samples, shearing was measured using amplicons in the RPP30 gene (Fig. 3a, d, e).

For plasmid control experiments, shearing of ZrAl-cut plasmids was analysed using two sets of amplicons, the Ψ and env IPDA amplicons and the equally spaced PSC amplicons shown in a. ddPCR analysis was done on fresh (D0) maxipreps of pNL4-3 linearized with ZrAl at a concentration mimicking patient samples. To assess the effects of higher levels of DNA fragmentation, IPDA analysis was also done on pNL4-3 DNA that had been incubated at 4 °C for 5 days (D5), and on pNL4-3 DNA cut with both ZrAl and EcoRI (two cuts). The mean and range of duplicate determinations of the DSI is shown for each set of amplicons. The DSI was the same for the IPDA and PSC amplicons at three different levels of shearing. The DSI was used to correct the IPDA droplet counts in e. Negative values were set to 0. e, Uncorrected and DSI-corrected IPDA analysis of the intact proviral construct pNL4-3 at different levels of fragmentation. After correction, positive droplets were almost exclusively in Q2 even at higher levels of fragmentation.
Extended Data Fig. 6 | Sequence analysis of Q2 proviruses. a, Sorting of productively infected CD4+ T cells. Cell preparations with a high fraction of intact proviruses were obtained by infecting CD4+ T lymphoblasts with a replication-competent HIV-1 carrying GFP in the nef ORF (R7-GFP37). After 48 h, GFP+ cells were collected by sorting. Genomic DNA was isolated, subjected to pulse field electrophoresis to remove unintegrated intermediates, and analysed by IPDA. b, IPDA analysis of high molecular mass DNA from sorted cells. Droplets in Q1 and Q4 largely reflect the shearing of intact proviruses (DSI = 0.46) during DNA isolation and purification. c, Frequency of intact proviruses in GFP− and GFP+ cells before and after correction for shearing. After correction for shearing, the frequency of intact proviruses in sorted GFP+ cells is close to the expected value of 1. d, Map of the HIV-1 genome in GFP-expressing HIV-1 vector R7-GFP used in a. GFP is inserted in the nef ORF. Positions of outer primers in the LTR and GFP used in single genome amplifications are indicated. e, Sequence analysis of nine independent single genomes. Arrows indicate positions of the ψ and env IPDA amplicons. Orange lines indicate intact sequence without deletions or hypermutation and identical to R7-GFP except for single base mutations (black lines).
Extended Data Fig. 7 | DSI for patient samples. The DSI was determined by ddPCR using two amplicons in a cellular gene (RPP30) spaced at exactly the same distance as the \( \Psi \) and env amplicons. It is the fraction of templates in which DNA shearing has occurred between the amplicons. Horizontal bars indicate median and interquartile range; data from \( n = 62 \) patient samples.
Extended Data Fig. 8 | In vivo decay rates of cells with intact and defective proviruses. The frequency of cells carrying intact proviruses, proviruses with 3′ deletion and/or hypermutation (3′ del/hyper), and proviruses with 5′ deletions (5′ del) was measured in resting CD4+ T cells from patients on long-term suppressive ART. Data are plotted in terms of decay rate assuming exponential decay. Half-life values for the same decay curves are shown in Fig. 4c. Negative decay rate indicates proliferation.
Extended Data Fig. 9 | Variability in decay slopes. Shown are the mean and s.d. of the decay slopes for intact and defective proviruses in infected individuals on ART sampled longitudinally (n = 14). Analysis based on decay data in Fig. 4a.
Reporting Summary

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Statistical parameters

When statistical analyses are reported, confirm that the following items are present in the relevant location (e.g. figure legend, table legend, main text, or Methods section).

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- Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
- A full description of the statistics including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
- Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated
- Clearly defined error bars
- State explicitly what error bars represent (e.g. SD, SE, CI)

Our web collection on statistics for biologists may be useful.

Software and code

Policy information about availability of computer code

Data collection
No software was used in data collection.

Data analysis
Sequence data were analyzed with alignments generated manually with Bioedit (http://www.mbio.ncsu.edu/bioedit/bioedit.html).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

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All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The IPDA was developed through an analysis of published near full genome HIV sequences. These sequences are described in Bruner et al. (reference 13 in the revised manuscript, Genebank accession numbers KX505390-KX505744) and Imamichi et al. (reference 14 in the revised manuscript, Genbank accession numbers KU677989-KU678196. There are no restrictions on the availability of these data.
Field-specific reporting

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Sequence analysis was performed on a database of all available full genome HIV sequences meeting study criteria (n=431). |
|-------------|---------------------------------------------------------------------------------------------------------------|
| Data exclusions | No sequences were excluded but specific analyses were carried out on sequences from the relevant patient populations (i.e. viremic or aviremic, acute or chronic) depending on the research question as described in the text. |
| Replication | Assay reproducibility is described in Extended Data Figures 3 and 4. |
| Randomization | This was not a clinical trial. |
| Blinding | This was not a clinical trial. |

Reporting for specific materials, systems and methods

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | Unique biological materials |
| ☑   | Antibodies |
| ☑   | Eukaryotic cell lines |
| ☑   | Palaeontology |
| ☑   | Animals and other organisms |
| ☑   | Human research participants |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | ChIP-seq |
| ☑   | Flow cytometry |
| ☑   | MRI-based neuroimaging |

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s) The J-Lat clone (#6.3) was obtained from the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH.

Authentication The cells were shown the contain the expected 1 copy of HIV / cell as described in the text.

Mycoplasma contamination The cells were negative for mycoplasma.

Commonly misidentified lines (See ICLAC register) No misidentified cells were used.

Human research participants

Policy information about studies involving human research participants

Population characteristics Blood samples were obtained from HIV-1 infected adults on suppressive ART with undetectable plasma HIV-1 RNA levels (< 50 copies per mL) for >6 months.

Recruitment Patients were recruited through care providers at the Moore Clinic (JH) or the Scope Study (UCSF). All participants provided written informed consent.