HIV-1 Integration Landscape during Latent and Active Infection

Graphical Abstract

Highlights

- Integration sequencing identifies clonally expanded and single HIV-1 integrations in human subjects
- Large clonal families of HIV-1+ cells are likely not part of the latent reservoir
- HIV-1 integrates near or into a 30 bp INT-motif found in Alu repeats

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In Brief

HIV-1-infected CD4+ T cells that undergo clonal expansion are able to proliferate because their proviruses are defective. Conversely, the replication-competent reservoir is likely found in the subset of CD4+ T cells that carry unique integrations and remain quiescent.

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HIV-1 Integration Landscape during Latent and Active Infection

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SUMMARY

The barrier to curing HIV-1 is thought to reside primarily in CD4+ T cells containing silent proviruses. To characterize these latently infected cells, we studied the integration profile of HIV-1 in viremic progressors, individuals receiving antiretroviral therapy, and viremic controllers. Clonally expanded T cells represented the majority of all integrations and increased during therapy. However, none of the 75 expanded T cell clones assayed contained intact virus. In contrast, the cells bearing single integration events decreased in frequency over time on therapy, and the surviving cells were enriched for HIV-1 integration in silent regions of the genome. Finally, there was a strong preference for integration into, or in close proximity to, Alu repeats, which were also enriched in local hotspots for integration. The data indicate that dividing clonally expanded T cells contain defective proviruses and that the replication-competent reservoir is primarily found in CD4+ T cells that remain relatively quiescent.

INTRODUCTION

Despite effective therapy, HIV-1 can persist in a latent state as an integrated provirus in resting memory CD4+ T cells (Chun et al., 1997; Finzi et al., 1997; Wong et al., 1997). The latent reservoir is established very early during infection, (Chun et al., 1998), and because of its long half-life of 44 months (Finzi et al., 1999), it is the major barrier to curing HIV-1 infection (Siliciano and Greene, 2011).

The HIV-1 latent reservoir has been difficult to define, in part because reactivation of latent viruses is difficult to induce and to measure. Viral outgrowth assays underestimate the size of the reservoir, while direct measurements of integrated HIV-1 DNA overestimate the reservoir because a large fraction of the integrated viruses are defective (Ho et al., 2013). Although the latent reservoir remains to be completely defined, establishing the reservoir requires intact retroviral integration into the genome and subsequent transcriptional silencing (Siliciano and Greene, 2011). Whether or not the genomic location of the integration impacts on latency is debated (Jordan et al., 2003; Jordan et al., 2001; Sherrill-Mix et al., 2013). However, HIV integration into the genome is known to favor the introns of expressed genes (Han et al., 2004), some of which, like BACH2 and MKL2, carry multiple independent HIV-1 integrations in different individuals and are considered hotspots for integration (Ikeda et al., 2007; Maldarelli et al., 2014; Wagner et al., 2014). However, there is currently no precise understanding of the nature of these hotspots or why they are targeted by HIV-1.

Viremia rebounds from the latent reservoir after interruption of long-term treatment with combination anti-retroviral therapy (cART). When it does, it appears to involve an increasing proportion of monotypic HIV-1 sequences, suggesting the proliferation of latently infected cells (Wagner et al., 2013). Based on this observation and the finding that a subset of cells bearing integrated HIV-1 undergoes clonal expansion in patients receiving suppressive anti-retroviral therapy, it has been proposed that the clonally expanded cells play a critical role in maintaining the reservoir (Maldarelli et al., 2014; Wagner et al., 2014).

To obtain additional insights into the regions of the genome that are favored by HIV-1 for integration and the role of clonal expansion in maintaining the reservoir, we developed a single cell method to identify a large number of HIV-1 integration sites from treated and untreated individuals, including “viremic controllers” who spontaneously maintain viral loads of <2,000 RNA copies/ml and “typical progressors” who display viral loads >2,000 RNA copies/ml.
RESULTS

Integration Library Construction

Twenty-four integration libraries were constructed from CD4+ T cells from 13 individuals: three provided longitudinal samples before and after (0.1–7.2 years) initiation of therapy; four were untreated; two were treated; and four were viremic controllers (Table S1). Patients were grouped into three categories based on viral loads and therapy: (1) viremic progressors were untreated individuals with viral loads higher than 2,000 viral RNA copies/ml of plasma; (2) progressors were treated individuals whose initial viral loads were higher than 2,000 viral RNA copies/ml before therapy; (3) controllers were individuals who maintain low viral loads spontaneously in the absence of therapy (<2,000 viral RNA copies/ml). The frequency of latently infected, resting CD4+ T cells in our patients was similar to that reported by others as measured by quantitative viral outgrowth assay (Table S1 and Laird et al., 2013).

Libraries were produced from genomic DNA by a modification of the translocation-capture sequencing method that we refer to in this paper as integration sequencing (Figure 1A) (Janovitz et al., 2013; Klein et al., 2011). Virus integration sites were recovered by semi-nested ligation-mediated PCR from fragmented DNA using primers specific to the HIV-1 3' LTR (Table S2). PCR products were subjected to high-throughput paired-end sequencing, and reads were aligned to the human genome. Since sonication is random, it produces unique linker ligation points that identify the specific integration events in each infected CD4+ T cell, which allows both single-cell resolution and identification of expanded clones of cells with identical integrations (Berry et al., 2012 and Figure 1A). Thus, integration sequencing can enumerate both the number of integration sites and the number of infected cells.

A total of 6,719 unique virus integration sites were determined (Table S3): 873 unique integrations in viremic controllers; 987 in viremic untreated progressors; and 4,859 integrations in treated progressors (Figure 1B).

Integrations Are Enriched in Introns of Highly Expressed Genes

We analyzed the genomic location of the integration sites obtained from viremic controllers and untreated and treated progressors and compared our results to published data obtained from HIV-1-infected individuals (Han et al., 2004; Ho et al., 2013; Ikeda et al., 2007; Schröder et al., 2002). In agreement with the work of others, the majority of integration sites in each group are genic (Figure 1C and Figure S1A). Moreover,
integrations are found more frequently in the introns of highly expressed genes, and there is a slight bias for viral orientation that leads to convergent transcription (Figures 1D, 1E, and 1F and Figures S1B–S1E) (Mitchell et al., 2004; Schröder et al., 2002). Thus, the general features of integrations defined by integration sequencing are similar to those obtained by others.

Although the differences between groups were small in magnitude, they were significant in that treated progressors had a smaller proportion of integrations in genic regions (p < 0.0001 and p < 0.0001, respectively) and in highly expressed genes (p < 0.0001 and p < 0.0001, respectively) when compared to viremic controllers and untreated progressors (Figures 1C and 1E). Conversely, the proportion of viral integrations in genes expressed at lower levels was increased in treated progressors compared to viremic controllers and untreated progressors (p = 0.002 and p < 0.0001, respectively). Viremic controllers and treated progressors were not significantly different from each other in terms of the level of expression of the genes at the sites of integration (Figure 1E). Thus, therapy is associated with a relative decrease in the number of cells with viral integrations in highly expressed genes.

Identification of Clonally Expanded Cells Containing Integrated HIV-1

Since we shear DNA ends randomly to produce our libraries and by paired-end sequencing can determine the precise site of both the integration and sheared end, we infer that identical integrations with unique sheared ends arise from clones of expanded cells (Figure 1A). Integrations can therefore be classified as clonally expanded (i.e., identical integrations with distinct sheared ends, deriving from the clonal expansion of an original unique, single integration event) or single integrations (i.e., unique integration site with a single sheared end).

Clonally expanded viral integrations were present in all individuals irrespective of therapy or viremia (Table S3). However, the proportion of clonally expanded viral integrations is significantly lower in viremic controllers (30%) and viremic progressors (27%) than in treated progressors (40%) (p < 0.0001 and p < 0.0001, Figure 2A and Figure S2A). Although the size of individual clones varied from 2 to 295 cells (Figure S2B), the relative increase in clonally expanded integrations during therapy consistently
translated into an increase in the number of infected cells that derive from expanded clones (Figure 2B). The percentage of cells containing clonally expanded HIV-1 integrations was similar in untreated progressors (78%) and controllers (79%), but it was significantly increased in treated progressors (90%) (p < 0.0001, Figure 2B and Figure S2C). Thus, therapy is associated with an increase in the frequency of clonal HIV-1 integrations and infected clonally expanded cells.

To determine whether the position of viral integration in the genome correlates with clonal expansion, we compared the location of genomic clonally expanded to single integrations. Both types of integrations favored genes and their introns (Figures 2C and 2D and Figures S2D and S2E). However, the proportion of clonally expanded integrations in intergenic regions was greater than that of single integrations (Figure 2C, p < 0.0001). Moreover, of the integrations in genes, single integrations were more likely to be found in highly expressed genes than clonal integrations (Figure 2E, p < 0.0001, and Figure S2F). Thus, cells harboring viral integrations in intergenic regions and genes that are expressed at lower levels are more likely to be clonally expanded.

Large Expanded Clones Are Found in Memory Cells
Central memory cells are thought to be the major source of the HIV-1 reservoir (Chomont et al., 2009). To investigate the nature of the cells that comprise the expanded clones, we performed virus integration sequencing on genomic DNA from sorted central, transitional, and effector memory CD4+ T cells (Figure 2F). In both individuals studied, all three subsets of CD4+ T cells contained expanded clones (Figures 2G and 2H and Figures S2G and S2H). Thus, central, transitional, and effector memory T cells, all of which have undergone antigen-stimulated cell division, harbor the expanded clones of HIV-1 integrants.

Clonally Expanded Integrations Increase after Therapy
The proportion of clonally expanded viral integrations is increased in treated progressors (Figure 2A and Wagner et al., 2014). To further examine the effect of therapy on clonal expansion, we analyzed longitudinal samples from three typical progressors before and during therapy (Table S1). We found an increase in the number of clonally expanded integrations throughout the treatment period of up to 7.2 years in two of the three patients (Figure 3A, p = 0.017, and Figure S3A) as well as an increase in the number of cells that contained clonally expanded viral integrations (Figure S3B). Correspondingly, there

Figure 3. Clonally Expanded Viral Integrations Increase and Single Integrations Decrease during Therapy
Graphs show data from patients 1 (blue), 2 (red), and 3 (green) from longitudinal time points (Table S1). Time was normalized from 0 to 1 (727 days pre-therapy to 2617 days post-therapy). Dotted line at t = 0.21 marks therapy initiation. Trendline was determined by linear regression model. Solid lines indicate significant change in proportion of events; dashed lines indicate insignificant change in proportion of events.

(A) Proportion of clonally expanded viral integrations (Int).
(B) Proportion of single viral integrations.
(C) Proportion of genic clonally expanded viral integrations.
(D) Proportion of genic single viral integrations.
(E) Proportion of intergenic clonally expanded viral integrations.
(F) Proportion of intergenic single viral integrations.
See also Figure S3.
was also an overall decrease over time in single integrations (p = 0.017), with a half-life of 127 months assuming a non-linear regression model for one-phase decay (Figure 3B). Thus, our data suggest that the numbers of single integrations decay very slowly over time, while clonally expanded integrations increase with time on cART.

The increase in the number of clonal integrations during cART did not favor genic or intergenic regions (p = 0.65), indicating that this effect is independent of the location of the integration in the genome (Figures 3C and 3E and Figure S3C). In contrast, single integrations decrease significantly in genic regions (Figure 3D, p = 0.036, and Figure S3D) and increase proportionally in intergenic regions (Figure 3F, p = 0.036). Thus, the fate of cells harboring single viral integrations in cART treated progressors differs from clonal integration. Moreover, the fate of single integrations is dependent on their location in the genome, whereas the clonal integrations are not. These results suggest that cells bearing genic single integrations are selected against during therapy and that clonal expansion is not.

Clonally Expanded Integrations in the Same Genes in Multiple Patients

In the three progressors who provided longitudinal samples, ~5% of the clonal integrations persisted through successive time points without selection for genic or intergenic regions compared to all clonal integrations (Figures 4A and 4B). Furthermore, of the genic integrations that persisted, there was also no selection for or against those in highly expressed genes (Figure 4C). Thus, the persistent clonal integrations are indistinguishable from the larger pool of clonally expanded viral integrations in terms of their position in the genome.

To determine whether specific genes or groups of genes are permissive for clonal expansion, we looked for overlap in genic integration sites between samples (Figures 4D–4F). Despite a higher number of single integrations, there was much greater overlap of the genes that harbor clonally expanded integrations between individuals irrespective of treatment or level of viremic control (p < 0.0001) (Figures 4D–4F). On average, there is 13% and 3% overlap between genes harboring clonally expanded and single viral integrations, respectively. The genes containing clonally expanded viral integrations in multiple patients are expressed at lower levels than genes containing overlapping single viral integrations (Figure 4G). Taken together, these results suggest that cells that carry integrations in highly expressed genes tolerate clonal expansion less well than cells with integrations in genes with lower levels of expression.

Since clonal integrations have been associated with genes involved in malignant transformation (Wagner et al., 2014), we examined our entire dataset for enrichment of integrations in cancer-associated genes (n = 743 cancer-associated genes [Vogelstein et al., 2013; Zhao et al., 2013]). Although there was an overall enrichment for integrations in cancer genes (329/4,410 = 7.5%) compared to all genes in the human genome (743/25,660 = 2.8%) (p < 0.0001), this preference does not seem to be significant because it is similar to the overall preference for integration into highly expressed genes (Figure S4). Furthermore, we observed no overrepresentation of single, clonal, or persistent integrations in cancer genes (Figure 4H).

Importantly, a significant decrease in integrations in cancer-related genes was observed in longitudinal samples (Figure 4I), suggesting that these are selected against with therapy.

Expanded Clones Contain Defective Viruses

Our method of integration sequencing captures the end of the 3' LTR and identifies the genomic site of viral integration. To determine whether the viruses found in expanded clones are intact, we used nested integration site-specific PCR primers that were anchored in the host genome to amplify the 5' LTRs of 75 expanded clones from eight individuals (Figure 5A and Table S2). The clones selected for PCR verification varied in size from 5 to 200 out of 0.3–2 × 10^6 CD4 T cells. Of the 75 sequences obtained, 24 showed fragmented 5' LTRs flanked by the correct genomic site, and an additional 44 of the proviruses did not have a recoverable 5' end (Figure 5B). The remaining eight viruses with intact 5' LTRs were amplified in limiting dilution conditions using integration site-specific primers and HIV-1 primers (Figure 5C).

Three of the eight viruses could not be amplified; four had large deletions in Env, one had a frameshift mutation in pol, and one had undergone APOBEC3G-mediated hypermutation to produce a premature stop codon in env (Figure 5D and Data S1). Thus, we were unable to find a single intact integrated provirus among 75 expanded clones.

Hotspots for Virus Integration

Overlap between integrations in the genes of different patients suggests the existence of hotspots for HIV-1 integration. A number of individual genes have been identified as preferential sites for HIV-1 integration, including BACH2, MKL2, DMNT1, MDC1, and STAT5B (Keda et al., 2007; Maldarelli et al., 2014; Wagner et al., 2014). To identify hotspots for HIV-1 integration genome-wide, we subjected our dataset to hot_scan analysis (Silva et al., 2014), which defines hotspots by identifying regions of local enrichment using scan statistics. This analysis identified 55, 85, and 247 hotspots for controllers, viremic, and treated progressors, respectively (Figure 6A). For example, the intron between exons 5 and 6 in MKL2 is a hotspot for integration in patient 11, contains an expanded clonal family in patient 10, and was also identified as a site of enrichment for integration by others (Maldarelli et al., 2014) (Figure 6B).

To validate our in silico analysis and to further characterize the MKL2 hotspot, we sequenced the gag gene from proviruses integrated into MKL2 by amplification with nested genomic primers specific for MKL2 and HIV-1 gag. Sequences obtained from patient 10, who showed only one expanded clone, are very closely related to each other, which is consistent with a single clonally expanded integration (Figure 6C). In contrast, sequences obtained from patient 11 are far more diverse, suggesting that there were several different viral integrations in the MKL2 hotspot (Figure 6C). We conclude that the hotspots defined by hot_scan represent multiple distinct integration events in close proximity.

Viremic progressors had the highest proportion of integration events in hotspots, indicating that, in the case of high-level viremia, there are specific genomic locations that favor integration (Figure 6D). Although the majority of all integrations fall outside of hotspots (Figure 6D), hotspots resemble other integrations in that
they are preferentially found within genes with a preponderance of these in introns (Figures 6E and 6F). In all cases, hotspots are enriched in highly expressed genes, and consistent with the overall decrease in viral integrations in highly expressed genes during therapy, the proportion of hotspots in these genes also decreases (Figures 6G and 1E). Thus, the general characteristics of hotspots are similar to features of all integrations.

To determine whether there is a relationship between hotspots and clonally expanded viral integrations, we enumerated single and clonally expanded integrations in hotspots (Figure 6H). Only a small fraction (11%–18%) of all single integrations were found in hotspots with untreated viremic progressors showing the highest level (Figure 6H). In contrast, there was a much higher proportion of clonal integrations in hotspots (30%–46%), with the lowest proportion in treated progressors (Figure 6H). This observation is consistent with the finding that there is a greater degree of overlap in genes that harbor clonally expanded rather than single integrations (Figures 4D–4F) and that clonally expanded integrations are more likely to be hotspots than single integrations (Figure 6H, p < 0.0001).
Viral Integration Enriched in Sites Containing a DNA Sequence Motif

To determine whether there are specific genomic features associated with sites of viral integration and hotspots, we examined 100 base pairs (bp) centered on all integration sites for the presence of a consensus sequence (Bailey and Elkan, 1994). We found 7% of all integrations within 100 bp of a highly conserved 30 bp motif (INT-motif) (Figure 7A). The majority of the integrations identified in this analysis were single integration events, with the ratio of single-to-clonal integrations being significantly different from the expected (Figure 7B, p < 0.0001). When HIV-1 integrates directly into the INT-motif, the 5' end of the motif is recurrently found 20 bp from the site of viral integration (Figure 7C). The INT-motif is asymmetrically distributed in Alu repeats, and its position coincides with a peak of viral integration (Figure 7D). Furthermore, there is a significant overall enrichment of integrations inside Alu repeats (Figure 7E) and in close proximity to Alu repeats, irrespective of whether the integration is inside genes or in intergenic regions (Figure 7F). Thus, a preference for Alu is independent of a preference for integration in genes.

Previous studies have shown a preference for integration into Alu repeats, potentially because Alu repeats are enriched in gene-rich regions (Schröder et al., 2002). To further examine the relationship between Alu repeats and transcription of genes, we determined the distance between Alu repeats and the center of all genes. There was no positive correlation between the position of Alu and the level of transcription (Figure 7G). To determine whether the distance between integration and Alu repeats correlates with transcription, we measured the distance between the sites of integration and Alu repeats in all genes (Figure 7H). There was no significant
The difference between integration distance to Alu repeats in highly expressed, silent, or trace level expressed genes. Therefore, the rate of transcription does not impact integration distance to Alu repeats, and integration at these sites must be independent of transcription.

Finally, the number of Alu repeats in a hotspot is directly correlated with the number of integration events in that hotspot (Figure 7I, $p = 0.86$). We conclude that HIV-1 has a preference for integration in close proximity to sites in the genome that are enriched in Alu repeats and that this preference is independent of the level of transcription.

**DISCUSSION**

CD4+ T cells that are actively infected with HIV-1 are rapidly eliminated during anti-retroviral therapy, but this form of treatment is relatively ineffective in selecting against latently infected CD4+ T cells, which have an estimated half-life of 44 months. Abolishing the latent reservoir is the current hurdle to finding a cure for HIV-1 infection. Although we have learned a great deal about the location of the latent compartment and its persistence during therapy, it has been difficult to uncover whether there are specific genomic features associated with latency (Siliciano and...
Figure 7. Consensus Motif for Viral Integration
(A) 30 bp sequence consensus motif (INT motif). 100 bp around all viral integration sites were analyzed for a consensus sequence by MEME (Bailey and Elkan, 1994). 444 integration sites were identified with the INT motif. E value: 6.4 \times 10^{-427}. The dotted line shows the preferred site of integration (see also C). (B) Number of single (S) and clonally expanded (CE) that were identified to contain INT motif within 100 bp of the integration site. \( p < 0.0001 \), using two-proportion z test. (C) Conserved integration site within INT motif. Histogram maps the start site (5' end) of INT motif with respect to the integration site (dotted line). Peak shows that the majority of integration sites occur 20 bp from the 5' end of the motif start site. Shaded region represents the location of the INT motif relative to the majority of the integration sites. (D) Location of integration preference and INT-motif inside Alu repeats is overlapping. (Left) Location of integration site Alu repeats were plotted relative to the midpoint of the repeat. (Right) The location of the start site of INT motifs within Alu repeats. (E) Integrations are enriched inside Alu repeats. Total integrations identified inside Alu repeats were enumerated (red diamond) and compared to the expected value as defined by Monte Carlo simulation. The boxplot displays the variation of the number of random integrations identified inside Alu repeats by each iteration of the simulation. (F) Integrations are near Alu repeats in genes and intergenic regions. Average distance to the nearest Alu repeat for all integrations inside genes or intergenic regions was calculated (red diamond) and compared to the expected distance as defined by Monte Carlo simulation. The boxplot displays the variation of the distance of random integrations from Alu repeats in genes or intergenic regions by each iteration of the simulation. (G) Distance to Alu repeats from the center of highly, medium, low, trace, or silently expressed genes. (H) Distance to Alu repeats in highly, medium, low, trace, or silently expressed genes. (I) Positive correlation between Alu repeats and integrations inside hotspots. Graph shows number of Alu repeats (x axis) versus integrations in hotspots (y axis). Hotspots not containing Alu repeats were removed from this analysis. The scatter plot shows the linear relationship between the number of INT motifs and integrations inside hotspots (Pearson’s correlation, \( p = 0.86 \)).
experiments in cell lines showing that level of HIV-1 transcription is dependent, in part, on the status of surrounding chromatin (Jordan et al., 2003; Jordan et al., 2001; Sherrill-Mix et al., 2013). HIV-1 integration has been studied in multiple cell types, but large libraries of integrations sites in primary infected T cell have only recently become available (Maldarelli et al., 2014; Wagner et al., 2014). Integration sites obtained from in-vitro-infected cell lines and primary T cells are distinct (Brady et al., 2009; Sherrill-Mix et al., 2013). Nevertheless, common features of HIV-1 integration have been defined, including the observation that integration favors Alu repeats (Schröder et al., 2002). This association was thought to be dependent on the presence of these repeats in the introns of gene-rich regions and not on a particular sequence feature (Schröder et al., 2002). However, we observed that integration preference into highly transcribed genes and into Alu repeats seem to be independently important, and furthermore, integrations are enriched near Alu repeats both in genic and intergenic regions. One possible explanation for preference for Alu seems to be the presence of an INT motif. TG-(N)5-7-CA sequence has been associated with sites of HIV-1 integration, but an integration consensus has not been defined (Brady et al., 2009; Holman and Coffin, 2005; Lewinski et al., 2006; Serrao et al., 2014; Wang et al., 2007; Wu et al., 2005). We found a 30 bp INT motif within 100 bp of 7% of all integrations, the large majority of which are single events. As expected, the HIV-1 INT motif contains a signature TG-(N)5-7-CA and can form a hairpin structure, anchored on 5’ NTG-3’, 5’ CAN-3’. This motif is frequently found at the 3’ end of Alu, where it coincides with a peak of viral integration events, and viruses integrated directly in this motif show a dramatic specificity for insertion site. The asymmetric peak and specificity of the integration site are remarkable. Nevertheless, we are likely underestimating the frequency of integrations within Alu because we can only map unique reads.

The observation that HIV-1 prefers to integrate in the neighborhood of Alu repeats is consistent with the finding that different individuals have been reported to have multiple integrations in selected genes (Ikeda et al., 2007; Maldarelli et al., 2014; Schröder et al., 2002; Wagner et al., 2014). Our experiments define a group of overlapping hotspots for integration that share many of the features of all HIV-1 integrations, including preference for introns of highly expressed genes and high density of Alu repeats. Viremic progressors showed the highest levels of hotspot integration, possibly because persistent integration leads to over-representation of these favored sites. Alternatively, these integrations might be positively selected by mechanisms that remain to be determined.

Individuals receiving cART show increasing numbers of cells with identical viral genomes by SGA, suggesting clonal expansion of a subset of cells bearing integrated proviruses (Buzon et al., 2014; Chomont et al., 2009; Wagner et al., 2013). Two independent groups have recently documented the long-term persistence of expanded clones of cells during therapy with cART (Maldarelli et al., 2014; Wagner et al., 2014). Our analysis confirms and extends these observations by showing that, when considered as a group, expanded clones are less likely to occur when the provirus is in a genic region, and clones that are associated with genes tend to be in genes that are expressed at lower levels than single integrations. Thus, proviruses inserted into active regions of the genome, which would be more likely to support viral re-activation during T cell proliferation, are generally selected against during clonal expansion.

Why certain integration sites are permissive for clonal expansion is not known, but finding that expanded clones with integrations occur in cancer-related genes led to the suggestion that integration into genes that regulate cell division promotes proliferation (Wagner et al., 2014). While we also found a higher proportion of integrations in cancer-related genes as compared to random, this bias was not different from that observed for other highly expressed genes favored by HIV-1. Further, we do not see any differential bias for integration in cancer-related genes in clonally expanded cells compared to single integrations and an overall decrease in the number of integrations in cancer related genes during the course of therapy. Since the number and size of clones increase with time on therapy, the data indicate that integration into cancer genes is unlikely to be a general contributor to the proliferation of infected cells.

Our data show that cART selects for expanded clones and that viremic controllers resemble treated progressors in showing a higher proportion of expanded clones than untreated viremics. cART selects for clonal integrations irrespective of the location in the genome. This is in contrast to single integrations, which are selected against by therapy. cART specifically favors the survival of single integrations in intergenic regions and is biased against genic regions, with an overall half-life for single integrations of 127 months. The half-life of single integrations is not too dissimilar from the current estimate for the latent reservoir, which is believed to decay with a half-life of 44 months on cART (Finzi et al., 1999).

The major outstanding question after the discovery of clonally expanded cells with integrated HIV-1 is whether the virus from these cells contributes to the latent reservoir (Maldarelli et al., 2014; Wagner et al., 2014). Several different independent lines of evidence argue against this idea. First, although the latent reservoir is thought to be contained primarily in resting central memory CD4+ T cells (Siliciano and Greene, 2011), we find that clonally expanded viral integrations are found in all three memory T cell compartments. Second, whereas the reservoir appears to decay with time on cART, we find that clonally expanded integrations increase with time and do so irrespective of whether they are found in genes or intergenic regions. In contrast, single integrations in more active parts of the genome, which are more likely to support HIV-1 reactivation, are selected against with time on ART. Finally, all 75 of the clonally expanded proviruses tested were defective, which is in agreement with two examples in the literature (Imamichi et al., 2014; Josefsson et al., 2013). Thus, we conclude that intact virus is not enriched in infected expanded cells. However, we cannot rule out the possibility that a rare clone of cells contains an active virus. Nevertheless, the 90% of all cells bearing integrated proviruses that account for expanded clones of infected cells in cART-treated progressors appear unlikely to be the major source of the rebounding latent reservoir. Instead, the replication-competent reservoir is likely to be contained in the remaining 10% of cells that harbor single integrations that decline with a long half-life on cART (Figure S5).
In conclusion, the data indicate that HIV-1-infected T cells that undergo clonal expansion are able to do so because their proviruses are defective and that the replication-competent reservoir is found in the subset of CD4+ T cells that remain quiescent.

**EXPERIMENTAL PROCEDURES**

**CD4+ T Cell Isolation for Integration Library Construction**

Human samples were collected after signed informed consent in accordance with Institutional Review Board (IRB)-reviewed protocols by all participating institutions. Patients 1, 2, and 3 were selected from the Seattle HIV longitudinal cohort studies at Fred Hutchinson Cancer Research Center. Patients 4, 8, and 9 were recruited from the University of Cologne, and samples were obtained at Rockefeller University (MN5, 0628). Patients 5, 6, and 7 were selected from the Rockefeller University HIV-1 antibody therapy clinical trial. Patients 10, 11, 12, and 13 were selected from a group of elite controllers that were followed at the Ragon Institute in Boston.

CD4+ T cells were isolated from whole PBMC using anti-CD4 microbeads (Miltenyi Biotec). The percentage of live cells was determined by flow cytometry based on forward and side scatter. Purity of CD4+ T cells was determined by labeling isolated cells with anti-human CD3, CD4, CD8, CD19, and HLA-DR and gating on CD3, CD4 double-positive cells. Isolated cells were used for library construction only if purity was >75%. CD4+ T cell subsets were isolated by FACS sorting on a BD Aria II by labeling cells with anti-human CD3, CD4, CD8, CD66b, CD335, HLA-DR, CCR7, CD27, and CD45RA. Analysis of CD4+ T cell subsets was done by pooling cellular DNA isolated from multiple sorts of the same sample.

**Quantitative Viral Outgrowth Assay**

Viral outgrowth was performed as previously described. ([Laird et al., 2013](#))

**Integration Library**

The method for integration library construction was adapted from TC-Seq (Klein et al., 2011).

**DNA Preparation**

DNA from 0.2–2 million CD4+ cells from HIV-1-infected patients was isolated and prepared as previously described (Klein et al., 2011). Fragments were ligated to 200 pmol of annealed linkers (Table S2). Virus sequences were eliminated by digestion with BglII (NEB), and fragments were purified.

**Integration Site Amplification**

Semi-nested ligation-mediated PCR was performed on DNA. All PCRs were performed using Phusion Polymerase (Thermo). DNA was divided into 700 ng aliquots and subjected to single-primer PCR with biotinylated LTR1 (1 × (98C-1 min) 12 × (8C-15 s, 62C-30 s, 72C-30 s) 1 × (72C-5 min)) (Table S2). Each reaction was spiked with pLinker and subjected to additional cycles of PCR (1 × (98C-1 min) 25 × (98C-15 s, 62C-30 s, 72C-30 s) 1 × (72C-5 min)). Products of 300–1,000 bp were isolated by agarose gel electrophoresis and magnetic streptavidin bead purification. Semi-nested PCR was performed on the magnetic beads first with a single primer LTR2 (same cycling conditions as above) followed by spiking in pLinker and additional cycles (Table S2). Products of 300–1,000 bp were isolated by gel electrophoresis.

**Paired-End Library Preparation**

Linkers were digested by Ascl such that a six-nucleotide barcode (CGCGCG) was left on the DNA fragments, indicating linker-dependent amplification. Fragments were blunted by End-It DNA Repair Kit (Epicenter), purified with AmPure beads (Agencourt), and ligated to NextFlex paired-end adapters. Adaptor-ligated fragments were enriched by 35 cycles of PCR with NextFlex primers [1 × (98C-1 min) 35 × (98C-15 s, 66C-30 s, 72C-30 s) 1 × (72C-5 min)], and fragments between 300–1,000 bp were isolated by gel electrophoresis. Two or three libraries were mixed in equimolar ratios and sequenced by either 150 bp paired-end sequencing on Illumina MiSeq or 150 bp or 100 bp paired-end sequencing on an Illumina 2500 HiSeq. Data are accessible via NCBI SRA using the accession number: SRP048222.

**Computational Analysis**

**Read Alignment**

Paired-end reads were mapped to the HIV-1 sequence (designated as a bait) using BLAT ([Kent, 2002](#)) with default settings. Reads that were mapped to the bait without mismatches were checked for the linker barcode in the paired-end read and were mapped to the human genome reference GRCh37/hg19 with Bowtie ([Langmead et al., 2009](#)). Only uniquely mapped reads (allowing for up to two mismatches) were used as defined in the best alignment stratum (command line options: -v2 -all -best-strata -m1). Identical reads generated by PCR amplification were merged.

**Integration Determination**

Once the paired-end reads were properly mapped in the bait and human genome (see above), we determined the integration breakpoint by aligning the remaining nucleotide sequence containing the 3’ terminus of the HIV-1 LTR to the human genome using BLAT (default settings). Only uniquely mapped reads up to 1 kb away from its partner were kept. Adjacent (within 50 nucleotides) putative integrations sites were merged. Finally, the 5’ end of the paired-end reads were used to deduce the integration and shear position sites in the human genome.

**Hotspot Detection**

To detect preferred sites of HIV-1 integration genome wide, we subjected our dataset to hot_scan software analysis ([Silva et al., 2014](#)), which defines hot-spots by scan statistics. Hotspots obtained by hot_scan were defined using different window widths (100, 200, 500, 1,000, 2,000, 5,000, 10,000, 20,000, 50,000, and 100,000 bp).

**Motif Analysis**

To determine a consensus motif, 100 bp flanking each integration site was analyzed for the presence of 30 bp consensus sequence using MEME software ([Bailey and Elkan, 1994](#)).

**Monte Carlo Simulation for Virus Integration and Hotspots**

Monte Carlo simulation was conducted by shuffling the genomic locations of all virus integration sites 10,000 times using bedtools shuffle utility ([Quinlan and Hall, 2010](#)). Then, we compared the observed number with the median number of integrations in the randomized list. We assessed enrichments by p value by counting the frequency of observed events being equal to or higher than the number of randomized events divided by n = 10,000.

**Statistical Analysis**

Proportion test is the standard test for the difference between proportions, also known as a two-proportion z test. We used R’s implementation of this via the prop.test() function.

**Integration Library Verification**

To verify our integration sequencing strategy, we constructed two libraries from DNA isolated from un-infected individuals. We recovered 13 sequences that mapped to integration sites. We subtracted these “integration sites” from all libraries before proceeding.

To test the saturation of our method, two separate integration libraries were constructed from identical samples for three patients. We found that both libraries contained the same expanded clonal families, but the majority of single virus integrations were unique to each sample of cells used for library construction. Single viral integrations found in both libraries were less than 1% of observed viral integrations.

**PCR Verification**

Genomic DNA isolated as described was serially diluted and subjected to nested-PCR using genomic specific primers and primers LTR1 and LTR2 (Table S2) using HotStart Taq Polymerase (QIAGEN) [1 × (98C-14 min) 40 × (98C-30 s, 55C-30 s, 72C-30 s) 1 × (72C-5 min)]. Products were isolated by gel electrophoresis and sequenced directly. Analysis of clones in this manner identified that we underestimate the size of clones by four to five times (data not shown).

**CD4+ T Cell Subset Sorting**

To isolate CD4+ subsets, we labeled PBMCs with antibodies to CD45RA, CD4, CD8, CD66b, CD335, HLA-DR, CD3, and CD27. We separated T cell subsets by FACS Aria (BD Biosciences) to very high purity (>98%).
Virus Sequencing

5’ LTRs from large clones were amplified with nested genomic primers and LTR2Rev (Table S2) using Platinum High Fidelity Taq (Invitrogen) [1 x (98C-14 min) 40 x (98C-30 s, 55C-30 s, 68C-1 min) 1 x (68C-5 min)]. Products were isolated by gel electrophoresis and sequenced directly.

Full-Length Virus

Full-length genomic DNA from infected patients was isolated as described and serially diluted. Each well was filled to a final volume of 50 μl with PCR reaction mixture (Platinum Taq MasterMix, Invitrogen) and primers to amplify virus from a specific integration site in the genome (Table S2 and Ho et al., 2013) using touchdown cycling to increase specificity. Then, 2μl aliquots from the first PCR were subjected to nested genomic PCR and 1% gel electrophoresis. The positive wells were gel purified, and fragments were sequenced directly.

ACCESSION NUMBERS

Data are accessible via NCBI SRA using the accession number SRP045822.

SUPPLEMENTAL INFORMATION

Supplemental Information includes five figures, three tables, and one data file and can be found with this article at http://dx.doi.org/10.1016/j.cell.2015.01.020.

AUTHOR CONTRIBUTIONS

L.B.C. planned and performed experiments, analyzed the data, and wrote the manuscript. I.T.S. performed all bioinformatics analysis. T.Y.O. performed culture gene analysis. R.A.R. performed bioinformatics and statistical analysis. E.H.P., G.H.L., and B.H.H. performed SGA and phylogenetic analysis. J.L.C., M.J.M., C.L., F.K., M.C., and B.D.W. selected and provided clinical samples. J.D.S. and R.F.S. performed QVOA and provided extensive discussion. M.J. planned and performed experiments, analyzed the data, and wrote the manuscript. M.C.N. planned experiments, analyzed the data, and wrote the manuscript.

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Figure S1. Characterization of Integrations in Controllers, Viremic, and Treated Progressors, Related to Figure 1

(A–D) Each dot represents data from an individual integration library.

(A) Proportion of integrations (Int) that are in genes in controllers (C), viremic (V) or treated progressors (T).
(B) Proportion of integrations that are in introns.
(C) Proportion of integrations that are in highly expressed genes.
(D) Proportion of integrations that are integrated in the same transcriptional orientation as host gene.
(E) Enrichment of integrations inside highly expressed genes. Number of HIV integrations in highly, medium, low, trace or silently expressed genes was enumerated (red diamond) and compared to expected number as defined by Monte Carlo simulation. The boxplot displays the variation of the number of random integrations in genes in each iteration of the simulation.
Figure S2. Comparison of Clonally Expanded and Single Integrations, Related to Figure 2

(A) Proportion of integrations (Int) that are clonally expanded in controllers (C), viremic (V) or treated progressors (T); each dot represents data from an individual integration library.

(B) Graph shows the size of proliferating clones of infected cells. Plotted is the number of shears (X-axis) by the number of integrations (Y-axis).

(C-F) Each dot represents data from an individual integration library.

(C) Proportion of infected cells deriving from clonal expansion.

(D) Proportion of clonally expanded (CE) and single (S) genic integrations.

(E) Proportion of clonally expanded and single integrations in introns.

(F) Proportion of clonally expanded and single integrations in highly expressed genes.

(G) Proportion of viral integrations (Int) that are clonally expanded, as identified by the same integration site with multiple shears in sorted CD4⁺ T cell subsets from patient 11.

(H) Proportion of infected cells deriving from clonal expansion in sorted CD4⁺ T cell subsets from patient 11.
Figure S3. Effects of Therapy on Clonally Expanded and Single Integrations, Related to Figure 3

Graphs show data from patients 1 (blue), 2 (red) and 3 (green) from longitudinal time points (Table S1). Time was normalized from 0 to 1 (727 days pre therapy to 2617 days post therapy). Dotted line at t = 0.21 marks therapy initiation. Trendline was determined by linear regression model. Solid lines indicate significant change in proportion of events; dashed lines indicate insignificant change in proportion of events.

(A) Proportion of clonally expanded viral integrations (Int).
(B) Proportion of clonally expanded cells.
(C) Proportion of genic clonally expanded viral integrations.
(D) Proportion of genic single viral integrations.
(E) Proportion of clonally expanded (top) or single (bottom) integrations in genes with high, medium or low expression.
Figure S4. Expression of Cancer-Related Genes with HIV-1 Integrations, Related to Figure 4
Proportion of cancer related (left) or total (right) integrations in genes with high, medium or low expression.
Figure S5. Persistence of HIV-1 in Infected Individuals
A model showing the hypothesized mechanism for HIV-1 persistence. Defective HIV-1 integrated proviruses can persist as memory CD4 T cells, and upon stimulation with antigen, these cells proliferate. Intact HIV-1 proviruses also persist as memory CD4 T cells. However, upon antigenic stimulation cells containing these viruses reactivate and begin producing virus and die by apoptosis.