Quantification of Unintegrated HIV-1 DNA at the Single Cell Level In Vivo

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Abstract

In the nucleus of HIV-1 infected cells, unintegrated HIV-1 DNA molecules exist in the form of one and two LTR circles and linear molecules with degraded extremities. In tissue culture they are invariably more numerous than the provirus, the relative proportion of integrated to unintegrated forms varies widely from ~1:1 to 1:10 and even over 1:100. In vivo, this ratio is unknown. To determine it, single nuclei from two infected patients with a known provirus copy number were microdissected, HIV DNA was amplified by nested PCR, cloned and individual clones sequenced. Given the extraordinary sequence complexity, we made the assumption that the total number of distinct sequences approximated to real number of amplifiable HIV-1 DNA templates in the nucleus. We found that the number of unintegrated DNA molecules increased linearly with the proviral copy number there being on average 86 unintegrated molecules per provirus.

Introduction

The phenomenal intrapatient variation of human immunodeficiency virus type 1 (HIV-1) genome needs no introduction [1,2,3,4,5,6,7]. The absence of proofreading mechanisms associated with the reverse transcriptase, the high recombination rate and high turnover are the main factors [8,9,10]. Whenever multiple infections occur, recombination invariably follows. Recombination is present at all levels of HIV genetics [11,12]. Within an infected individual recombinant genomes show up in network analyses of HIV sequences [13,14,15,16]. In an animal model, macaques inoculated simultaneously with SIVmac239 vpr and Δvpx or Δvpx and with SIVmac239Δeux, the emergence of wild-type virus was detected in blood in as little as 2 weeks post-inoculation [17,18]. Finally, some strains in widespread circulation are clearly composites of at least 2–3 other clades [19,20,21,22].

Recently, it was shown that in infected patients ~85% of infected CD4+ T cells in blood contain only one copy of HIV-1 DNA [23]. This would suggest a limited potential for recombination in virus produced by these cells. However, HIV replicates mainly in secondary lymphoid organs. The spleen is a secondary lymphoid organ replete with white pulps and germinal centres and it is in such structures that HIV recombination will occur. Indeed, the majority of HIV-1 infected cells in vivo harbour multiple proviruses and additional unintegrated DNA molecules [4,5], the average proviral copy number per infected cell being 3–4 with a range of 1–8 [3]. Greater than 75% of infected cells harbour two or more proviruses [5]. The ratio of unintegrated/provirus DNA molecules was not addressed [5]. It is well known that proviruses are accompanied by unintegrated DNA forms, either as covalently closed circles with one or two LTRs or linear molecules with ends degraded by exonucleases [24,25,26,27]. In tissue culture experiments the relative proportion of integrated to unintegrated forms varies widely from ~1:1 to 1:10 and even over 1:100 [28,29].

How can this ratio be addressed for single cells in vivo? Fluorescence in situ hybridization (FISH) can quantify the number of proviruses per cell [5]. In this work, Jung et al., reported extensive genetic variability within the hypervariable V1V2 region of viruses from two patients. Indeed, the vast majority of sequences were unique. This is perhaps not too surprising as the hypervariable V1V2 region of env is one of the most variable regions of the HIV-1 genome and thus offers the greatest resolution. This choice also meant there was no interference from the HIV-1Δenv probe used for FISH [5]. Given the phenomenal variation for these two patients, it is probably a reasonable assumption to equate the number of unique sequences with total number of distinct molecules within the nucleus. Accordingly, if sequencing was performed on the nuclei of single cells, it is possible to estimate the number of unintegrated DNA copies per nucleus.

Materials and Methods

Patients/Ethics Approval

The two patients, B and R, have been already described [1,4,5,30]. Briefly, patient B was at stage clinical B1 and had a blood CD4 count of 583/µl and a plasma viremia titre of 3,900 RNA copies/ml. Patient R was at stage C2, while the blood CD4 count and viremia were 317/µl and 126,000 RNA copies/ml respectively. Splenic tissues from patients B and R came from
Hôpital Saint-Louis (Paris, France), [1,4,5,30]. Ethics approval for the two patients have been obtained from Hôpital Saint-Louis (Paris, France). Informed consent were obtained from the patients B and R.

Preparation of Cells for Fluorescence In situ Hybridization

Frozen spleen cells from HIV-infected patients were thawed and stimulated with 2.5 μg/ml phytohaemagglutinin (PHA) (Difko, Detroit, MI, USA) in the presence of 10 μM azidothymidine (AZT) (Sigma, Taufkirchen, Germany) to prevent the virus spread in the cell culture [1,4,5,30]. After two days of culture in RPMI 1640 (Lonza, Verviers, Belgium) with 10% fetal calf serum (Invitrogen, Karlsruhe, Germany), 1% penicillin/streptomycin (Biochrom AG, Berlin, Germany) and IL-2 (100 U/ml; Chimron Behring, Marburg, Germany), the cell nuclei were prepared via incubation with 0.1 M KCl for 30 min at 37°C, fixed with methanol/acetic acid 3:1 at 4°C and stored at 20°C. The HIV FISH was performed according to our previous HIV protocol [3]. Nuclei with defined HIV provirus numbers were microdissected using a PALM Robot16 MicroBeam, according to the manufacturer’s instructions (PALM Bernried, Germany), and transferred to PCR tubes with the laser pressure catapulting technique.

PCR Amplification, Cloning and Sequencing

A fragment of the HIV envelope genome (V1V2) was amplified employing a semi-nested procedure. In order to increase sensitivity and specificity, hot start PCR was performed. First round primers were LV15 5’-gccacatctggtagttgcc and LV13 5’-CTTCTAGATGAGGAAAGCTTCC while primers SK122 and SK123 5’-CTAAAGCCATGTTGTTAATTCCG and 5’-TGCTCTAAAAGATATCTTGAGCA were used for the second round. The first and second round of PCR concerned standard amplification, the reaction parameters were 95°C for 5 min, followed by 35 cycles (95°C for 30 s, 55°C for 30 s, and 72°C for 30 s) and finally for 10 min at 72°C. The buffer conditions were 2.5 mM MgCl₂, 50 mM KCl, 10 mM Tris-HCl (pH 8.3), 200 μM of each dNTP, 100 μM of each primer, and 2.5 units of Taq DNA polymerase (Cetus) in a final volume of 50 μl. The equivalent of 0.2 l of the first round reaction is used as template in the second round. Amplification products were purified from agarose (Qiagen II kit, Qiagen, France) and directly cloned into the pCR2.1-TOPO vector included in the TOPO TA Cloning Kit (Invitrogen Corp., San Diego, CA). After transformation of E. coli TOP10F’ Blue cells, up to 4000 clones were picked and sequenced by GATC Biotech.

Determination of PCR-mediated Recombination Frequency

To appreciate the PCR-mediated recombination frequency, ten picograms each of DNA corresponding to V1V2 mini-prep clones 01 and 16 were mixed and subjected to 35 cycles of PCR with primers L1 and L2 and 1/10 of the first PCR was used to amplify for 35 cycles with primers SK122 and SK123 under the same conditions as those described above [2,5,31]. The DNA was purified and cloned into TOPO TA cloning site. About 500 colonies were screened with 4 different 32P-labeled oligonucleotides (Probes A1 5’ AACACAAATATCTAGGCAA and A2 5’ TGATCTCTCTATGTTATCG for clone 01 and B1 5’ GTCATCTTTATATCAGGC and B2 5’ TATAGGAAATGACTACTA for clone 16, data not shown). Plaques giving positive hybridization signals with both couple of primers A1+B2 or/and B1+A2 were considered PCR-mediated recombinants. Recombination was confirmed by sequencing and a recombination frequency of 2/500 or ~4×10⁻³ established.

Results

Quantification of Unintegrated HIV-1 DNA

For sequencing, how many molecules should be sequenced to detect the real number of distinct viral DNA in a sample? We considered P, the probability of finding a new sequence, and N as the absolute number of distinct sequences, while x and y are the number of total and unique sequences experimentally determined respectively. The relationship between P and N is obtained from the equation, P = (N−y)/N (Figure 1). For an x/y ratio ~2 approximately 80% of unique sequences (y) can be identified. HIV-positive interphase nuclei from two patients (B and R) were laser microdissected and transferred to PCR tubes and the V1V2 env region amplified. In order to increase sensitivity and specificity, hot start PCR was performed. Four splenocytes from patient R harbouring a single provirus constituted the starting point (Figure 2). Small numbers of sequences were sufficient for the number of unique sequences (y) to plateau. The values of y ranged from 12–14 to 39, which translates into values of N from 13–16 and 42 respectively (Figures 2). Given the underlying assumption that all genomes are genetically unique, the unintegrated/provirus ratios are ≥13:1 to 42:1.

A similar analysis was performed on the nuclei of six splenocytes with 2 or 6 proviruses from patient B (Figures 3A and 4A). The number of unique sequences identified ranged from 131 to 172 for cells harbouring 2 proviruses, and between 359 and 677 for those with 6 proviruses (Figure 4A). Values of N ranged from 150–927, while the unintegrated/dNA/provirus ratios (Z) were of the order of 75–155 (Figure 4A). Approximately 927 unique HIV genomes were predicted for nucleus B06 (Figures 3A and 4A). A selection of V1V2 protein sequences from B06 is shown in Figure 3B, a number of which are arguably recombinants. Most probably if more sequences from other cells were sequenced it would be possible to identify more recombinants. Indeed, given the

\[ \frac{y}{x} = \frac{N}{1 - e^{x/N}} \]

\[ x = -\ln(p)/(1 - p) \]

\[ P = (N - y)/N \]

Figure 1. Relationship between the probability P of scoring for a new sequence as a function of the ratio of genetically distinct (y) and total (x) sequences.

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frequency of HIV recombination and the 70% fraction of multiply infected splenocytes for patient B, probably all sequences were recombined in a relatively recent time frame. Recombination was confirmed by sequencing and a recombination frequency of 2/500 or \(4 \times 10^{-3}\) established. In parallel, Taq polymerase mutation frequency was shown to be \(f \approx 10^{-3}\), suggesting that the high mutation frequency observed was not associated to Taq polymerase.

**G to A Hypermutation at a Single Cell Level**

The B6 nucleus also harboured two G->A hypermutated sequences which were interesting. Such genomes are associated with editing by host cell cytidine deaminases APOBEC3F and 3G in the context of a \(Δvif\) genotype. The enzymes show distinct dinucleotide contexts associated with editing, notably \(T\_p\_C\) for APOBEC3F and \(C\_p\_C\) for APOBEC3G, where the edited base is underlined [32,33,34,35,36,37]. The two sequences showed remarkable selectivity for dinucleotide editing context. For clone 06 \(T\_p\_C\) was very strongly preferred over \(C\_p\_C\), while for clone 79 the opposite was apparent (Figure 3C). These striking differences suggest that occasionally only A3F or A3G are packaged, which is feasible given the estimations of low levels packaged into virions [38]. Presumably hypermutated genomes with less startling biases probably reflect co-packaging of both APOBEC3 proteins.

**Discussion**

For these two patients, proviruses are accompanied by a very large number of unintegrated forms, varying from \(Z = 13–155\) molecules per provirus (Figure 4A). As only 10 nuclei were analysed (4 from patient R +6 from B), this number is by no means a maximum. Interestingly the relationship between N and pv is linear (Figure 4B). While the sample size of this present study is low and thus has limited statistical power, the results obtained for the patients are similar. Given these large values for Z, it is logical that many splenocytes harbour unintegrated DNA alone. As the present study focussed on cells with FISH-positive proviruses such cells were not scored. While the values for Z are large compared to those derived from tissue culture, a recent report in a very different setting, peripheral blood mononuclear cells from HIV-infected Elite suppressors identified an unintegrated/integrated DNA ratios of between 10/1 to up to 10,000/1 [39].
The extraordinary genetic variation among HIV DNA molecules found in different nuclei indicates that the source of virus infection was very complex. Within an established HIV-infected patient there are three sources of genetically complex virus. These are circulating virus, virus on follicular dendritic cells (FDC) and virus associated with dendritic cells (DC) presented to CD4^+^ T lymphocytes. As the infection frequency of splenocytes was 1%, this rules out circulating virus, as the major source. The virus on FDC surfaces is essentially in the form of immune complexes and is mainly presented to B cells [40]. By contrast DCs present very efficiently HIV to CD4^+^ T lymphocytes [41]. While the present data cannot distinguish between the FDC and DC sources, the cellular immunology of DCs pleads in favour of the latter.

Using cell free virus to infect cells ex vivo, treatment of the target cell by proteasome inhibitors enhances viral growth ~3 fold indicating that more viruses infect and enter a cell than make it to the nucleus [42]. Assuming a comparable phenomenon in vivo the number of incoming virions would be of the order of 40–2800. If other catabolic pathways were operative, the number of virions would necessarily be even greater.

Infection of a single cell by many virions could well be a general phenomenon in virology. However, in order to quantify the number of incoming virions some trait is needed to distinguish them from the replication template for progeny virus. For retroviruses the incoming genome is RNA while the template for transcription is DNA. For RNA viruses, such as poliovirus, there is no such trait distinguishing parent and daughter genomes. Of course recombinants are a tell tale sign. Following vaccination with the three attenuated polioviruses, recombinants among them have been described, as have recombinants between vaccinating strains and wild type poliovirus and other enteroviruses [43,44]. Although negative stranded viruses such as influenza are known to recombine rarely, recombinants can be identified [45]. Along with the above data such examples indicate that multi-infection is probably commonplace, an inevitable consequence of the capacity of a cell to produce hundreds to thousands of virions in a very small space.

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**Figure 3. Phenomenal sequence diversity within single cell B06.** A) Quantification of unintegrated HIV DNA in a single splenocyte from patient B with 6 proviruses. B) Env V1V2 amino acid sequences from patient B6 are shown and compared to an arbitrarily selected sequence as reference. Only differences are shown. The sequences are coded by capital letters to the right with obvious recombinant molecules indicated. Dots denote synonymous substitutions, while arrows correspond to recombination sites. C) Dinucleotide context associated with two singular APOBEC3 edited genomes. The ordinate represents the substitution frequency as a function of the 5’ nucleotide.

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Author Contributions
Conceived and designed the experiments: RS AM. Performed the experiments: RS. Analyzed the data: RS AM. Contributed reagents/materials/analysis tools: RS AM. Wrote the paper: RS AM.

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Figure 4. Quantitation of HIV DNA molecules in single nuclei. A) Tabulation of the number of detected and projected unintegrated forms for nuclei from splenocytes of patients R and B. pv, proviral copy number detected by FISH; y, # of unique sequences identified; x, # of sequences determined; P, probability of finding a new unique sequence; N, absolute # of unique sequences; Z, unintegrated DNA/provirus ratio. B) Linear correlation between unintegrated and proviral copy numbers.
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