HIV-1 viral blips are associated with repeated and increasingly high levels of cell-associated HIV-1 RNA transcriptional activity

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Objective: Some HIV+ patients, virally suppressed on ART, show occasional ‘blips’ of detectable HIV-1 plasma RNA. We used a new highly sensitive assay of cell-associated HIV-1 RNA to measure transcriptional activity in PBMCs and production of infectious virus from the viral reservoir, in patients with and without ‘blips’.

Design/methods: RNA and DNA extracted from cells in 6 ml of peripheral blood, from suppressed patients with one to two ‘blip’ episodes over the past 2 years of ART (n = 55), or no ‘blips’ (n = 52), were assayed for HIV-1 RNA transcripts and proviral DNA targeting the highly conserved ‘R’ region of the LTR. Follow-up samples were also collected. Purified CD4+ T cells were cultured with anti-CD3/CD28/CD2 T-cell activator to amplify transcription and measure replication competent virus.

Results: HIV-1 RNA transcripts ranged from 1.3 to 5415 copies/10^6 white blood cells. ‘Blip’ patients had significantly higher levels vs. without blips (median 192 vs. 49; P = 0.0007), which correlated with: higher levels of inducible transcripts after activation in vitro, sustained higher HIV-1 transcription levels in follow-up samples along with increasing HIV-1 DNA in some, and production of replication-competent HIV-1.

Conclusion: Viral ‘blips’ are significant reflecting higher transcriptional activity from the reservoir and contribute to the reservoir over time. This sensitive assay can be used in monitoring the size and activity of the HIV-1 reservoir and will be useful in HIV-1 cure strategies. Copyright © 2021 The Author(s). Published by Wolters Kluwer Health, Inc.

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Introduction

Most HIV-1-infected individuals on antiretroviral therapy (ART) successfully reduce HIV-1 RNA plasma viral load (pVL) to less than 20 copies/ml, the lower limit of detection for standard diagnostic RT-PCR assays [1,2]. Nonetheless, occasional viral blips (elevated pVL between 20 and 200 copies/ml) occur [3–6]. These blips could be related to assay variation, given the extremely sensitive nature of the pVL diagnostic test [7–11] or a brief cytokine/antigen-driven increase in HIV-1 replication from the HIV-1 reservoir in the context of an unrelated illness [12], or residual activation from the reservoir [13–16]. Importantly, no new resistance mutations have been seen before, during, or shortly after blips [17].

The clinical significance of viral blips is unclear. Some studies have shown an association with progression of disease [3,13–16,18], whereas most do not [4–6,19]. Recent evidence suggests that they impede the ART-related decline in the HIV-1 latent reservoir [20]. A better understanding of the precise nature of blips would help address these uncertainties.

The research-based highly sensitive single copy RNA assay of HIV-1 has quantified very low levels of residual pVL on ART [18,21–26]. However, this methodology is technically challenging, because of the very low concentration of HIV virions in suppressed patients in plasma, on ART. Other assays have been developed to quantify the intact proviral HIV-1 DNA reservoir [27–31], including the Intact Proviral DNA Assay (IPDA) [27] and the quadruplex qPCR (Q4PCR) assay [28] but are impractical for routine clinical use. The latter two have been designed to increase the probability of quantifying only intact HIV-1 DNA using two or four highly conserved regions (reviewed in [29]). However, infected cells containing replication-competent provirus are very rare, usually containing only one copy of HIV proviral DNA per infected cell.

Instead, cell-associated RNA assays can detect many copies of spliced and unspliced RNA [30,31]. Furthermore, measurements of cell-associated HIV-1 RNA in PBMC cultures activated in vitro with either anti-CD3/CD28 [32,33] or PMA/ionomycin [34], have found HIV-1 is able to newly infect reporter CD4+ T-cell lines, thereby corresponding to replication-competent HIV-1 in circulating cells. Indeed, levels of cell-associated HIV RNA may be a surrogate for reservoir activity: higher levels prior to ART discontinuation correlate with quicker pVL rebound during treatment interruptions [35,36].

We used a recently described, extremely sensitive assay to detect intracellular HIV-1 RNA and total HIV-1 DNA [37], targeting the highly conserved ‘R’ region of the LTR. This ‘Double-R assay’ is able to detect all HIV-1 mRNA as well as total integrated HIV-1 DNA. Herein we used this sensitive assay to better understand HIV-1 transcriptional activity and DNA in PBMCs from blip-experienced and blip-negative patients. We addressed the significance of the ex-vivo HIV-1 transcript activity in peripheral blood and compared it with transcript activity of in-vitro activated purified CD4+ T cells.

Methods

Study design, DNA and RNA extraction
This study was approved by the St Vincent’s Hospital Human Research Ethics Committee (HREC LNR/16/SVH/327) and used standard-of-care samples sent to the NSW State Reference laboratory for HIV, St Vincent’s Hospital between 2017 and 2020 for monitoring pVL and CD4+ T-cell counts. Demographics and HIV-1 disease characteristics available from St Vincent’s Hospital patient database, are shown in Table S1, http://links.lww.com/QAD/C214. Also, an elite controller study was approved by South Western Sydney Local Health District Human Research Ethics Committee (HREC 2020/ETH00235). We defined viral blips as elevated routine patient monitoring pVL between 20 and 200 copies/ml. The median interval between analysis time points for this observational study was 6 months [interquartile range (IQR) = 5–8]. White blood cells (WBCs) were prepared from 6 ml of fresh anticoagulated whole blood in ACD (acid citrate dextrose) tubes, as previously described [37]. DNA and RNA were extracted using the Maxwell RSC automated extraction platform (Promega, Madison, Wisconsin, USA), with the Maxwell RSC Buffy Coat DNA kit (Promega) and Maxwell RSC Simply RNA Tissue kit (Promega), respectively, according to the manufacturer’s protocol. WBCs were counted after red blood cell lysis using TC20 Automated Cell Counter (Bio-Rad, Hercules, California, USA), and these counts were used to normalize the HIV-1 DNA and RNA results as copy numbers/10⁶ cells.

The double-R assay based on πCode End-Point PCR assay
We used a recently reported assay to detect intracellular HIV-1 transcription and total HIV-1 DNA [37], targeting the highly conserved ‘R’ region in both the 5’-LTR and 3’-LTR regions (Fig. S1, http://links.lww.com/QAD/C213). The Double-R assay is based on the precision image pi-code (πCode) MicroDiscs detection platform [37–39], and is at least 27 times more sensitive than the current gold standard Real-Time PCR assay [37]. HIV-1 copy number per patient sample, in duplicate, was determined from a standard curve generated with HIV-1 plasmid controls, with 8 points of 0.73–1600 HIV-1 copies/μl. The HIV-1 copy number was normalized per one million of WBCs and used as the standardized unit.
Analysis of intact sequences in proteinase/reverse transcriptase and integrase region
To confirm the presence of intact HIV-1 sequences, protease/reverse transcriptase (PR/RT) and integrase regions were amplified with the in-house Drug Resistance assay, at the NSW State Reference laboratory for HIV, St Vincent's Hospital [37,40]. HIV-1 subtype was identified using the HIV Drug Resistance Database, Stanford University [41]. Identification of insertions, deletions, stop codons and APOBEC3G-related mutations in PR/RT and integrase regions was conducted using the same Stanford HIV Drug Resistance Database.

Isolation of CD4^+ T cells by negative selection and culture with ex-vivo T-cell activation
CD4^+ T cells were isolated from 3 to 4 ml of fresh ACD-anticoagulated whole blood using the RosetteSep CD4 Enrichment Cocktail (StemCell Technologies, Vancouver BC, Canada), following the manufacturer's instructions. Briefly, 50 μl of RosetteSep reagent was mixed per milliliter of whole blood, incubated at room temperature (RT) for 20 min, and then layered on Ficoll-Hypaque for density centrifugation. The layer of CD4^+ -enriched cells was washed twice with PBS and resuspended in 2 ml of RPMI containing 20%FCS and 20 IU/ml of IL-2 (Roche, Basel, Switzerland; 10799068001). Cells were counted using TruCount tubes (BD Biosciences, San Jose, California, USA) and simultaneously analyzed for purity, by flow analysis on a 5-laser Fortessa as previously described [42]. Typically, 1.5–2 × 10^6 cells at more than 96% purity of CD4^+ CD4^- cells were obtained and incubated in two wells of a 48-well plate (Corning, Corning, New York, USA). One well was left with IL-2 only, and to the second well was added 5 μl of anti-CD3/CD28/CD2 (StemCell Technologies). Cells were cultured for 3 days in 5% CO2 at 37°C, before RNA extraction, using the Maxwell RSC SimplyRNA Tissue kit. HIV-1 RNA transcript numbers in these cultures were normalized to copy numbers/input 10^6 CD4^+ T cells.

Reverse transcriptase assay of supernatants from activated CD4^+ T and SupT1 cells
Reverse transcriptase activity in culture supernatants was determined as previously described [43,44]. Reverse transcriptase activity was used to measure the amount of HIV-1 in 10 μl of culture supernatant from activated CD4^+ T cells, collected after days 6 and 9 of culture (see Fig. S7 for further details, http://links.lww.com/QAD/C213).

Statistical analysis
Standard curves from known concentrations of HIV-1 plasmid copy numbers were generated with GraphPad Prism v7 (GraphPad Software, San Diego, California, USA). HIV-1 transcription levels in blip versus without blip groups were analyzed by nonparametric Mann–Whitney test. Pearson correlation was used for analysis of HIV-1 transcription and HIV-1 DNA levels. HIV-1 transcription levels in IL-2-only cultures versus IL-2 with the anti-CD3/CD28/CD2 activator cultures were analyzed by the parametric paired t test.

Results
Intracellular HIV-1 transcript activity and total HIV-1 DNA levels in peripheral blood
The Double-R assay (Fig. S1a and b, http://links.lww.com/QAD/C213) detected cell-associated HIV-1 transcriptional activity from both patient groups (see Table S1 for patient group details, http://links.lww.com/QAD/C214). Individual patient HIV-1 transcriptional activity, expressed as copies/10^6 WBCs, had a range greater than 3 log_{10}, from 1.3 to 5415 copies/10^6 WBCs (Fig. 1a and b, Fig. S2a and b, http://links.lww.com/QAD/C213). The median HIV-1 RNA transcription levels in the ‘blip’ group was significantly higher than the ‘without blip’ group (Fig. 1a: 192 vs. 49 copies/10^6 cells; P = 0.0007 and Fig. S2a, http://links.lww.com/QAD/C213). The median total HIV-1 DNA levels in the blip group was also significantly higher than the without blip group (Fig. 1b: 88 vs. 23 copies/10^6 cells; P < 0.0001 and Fig. S2b, http://links.lww.com/QAD/C213).

We then assessed the correlation between HIV-1 RNA transcription levels and HIV-1 DNA levels in each of the two groups, and identified a strong correlation between intracellular HIV-1 RNA transcripts and HIV-1 DNA for each patient group (Fig. 1c and d; P < 0.0001 and Fig. S2c and d, http://links.lww.com/QAD/C213).

Importantly, Fig. 1c also shows that, at any given HIV-1 DNA level, the number of transcripts varied by up to 3 log_{10} between patients.

Follow-up sample analysis in HIV-1 transcript activity and total HIV-1 DNA levels
We further investigated whether the initial high levels of HIV-1 transcription and DNA were also seen in follow-up samples over a median of 17 months (IQR = 11–20). Ten blip and twelve without blip patients had multiple samples from different time points assessed (Fig. 2a and b and Fig. S3, http://links.lww.com/QAD/C213, 4, http://links.lww.com/QAD/C213).

Overall, the significant elevation of HIV-1 transcripts in blip patients compared to without blip patients was maintained (Fig. 2); initially with median 209 vs. 38 respectively; P = 0.011) and at last follow-up with median 305 vs. 67, respectively; P = 0.020. HIV-1 total DNA was significantly higher in blip patients initially (median 50 vs. 17; P = 0.020) and at last follow-up (median 118 vs. 15; P = 0.0011) (see Fig. S3, http://links.lww.com/QAD/C213 and Fig. S4 for details, http://links.lww.com/QAD/C213)
Isolation and in-vitro activation of purified CD4⁺ T cells

As HIV-1 infection is known to persist in peripheral blood memory CD4⁺ T cells [32,45–48], we investigated the effect on HIV transcripts of using strong polyclonal stimulation of purified CD4⁺ T cells. In 15 consecutive patients, constituting nine blip patients and six without blip patients, CD4⁺ T cells were isolated from 3 to 4 ml of whole blood samples, using negative selection, to achieve greater than 96% purity (Fig. S5a, http://links.lww.com/QAD/C213). The purified CD4⁺ T cells were set up as 3-day cultures containing IL-2 in the presence of a highly effective T-cell activator combination of anti-CD3, anti-CD28 and anti-CD2 monoclonal antibodies (anti-CD3/CD28/CD2 activator). A parallel culture containing only IL-2 was also studied for each patient sample.

HIV-1 transcripts were detected in CD4⁺ T cells ex vivo, prior to culture, as described above, for these 15 patients (day 0, left in Fig. 3a). The purified CD4⁺ T cells cultured with IL-2 had approximately similar transcript levels to the ex-vivo results from the same patients (middle column versus first column in Fig. 3a). With the anti-CD3/CD28/CD2 T-cell activator and IL-2, the transcription levels were greatly increased in all but two patients (right column in Fig. 3a).

Overall, transcriptional activity in the CD4⁺ T cells ex vivo, prior to culture, highly correlated with...
transcriptional activity in the purified CD4$^+$ T cells cultured in vitro in the presence of IL2 and anti-CD3/CD28/CD2 activator ($***P<0.001$, Fig. 3b).

**Presence of replication-competent HIV-1 in latent reservoir**

We extended the analysis of the intracellular mRNA obtained from the activated CD4$^+$ T cells, using HIV-1 Proteinase/Reverse Transcriptase and integrase region sequences (Fig. 3a, Table S2, http://links.lww.com/QAD/C215). The sequence data revealed that the HIV-1 promoter was able to transcribe long sequences of ‘intact HIV-1 mRNA’ in most of the activated cultures: 13 out of 15 samples in HIV-1 PR/RT region and 12 out of 15 samples in HIV-1 IN region. We could not identify any insertions, deletions, stop codons, or APOBEC3G-related mutations from a total $\approx$3500 bp long sequence analyses (Table S2, http://links.lww.com/QAD/C215 and Table S3, http://links.lww.com/QAD/C216). The anti-CD3/CD28/CD2 activator combination was able to induce intact, nonmutated HIV-1 RNA sequences from its promoter in all but three patients (Fig. 3a).

Two samples (ID953, 956) out of 15 for the HIV-1 PR/RT region, and three (ID952, 953, 956) out of 15 samples for the HIV-1 integrase region, failed to amplify by PCR for HIV-1 sequencing (Table S3, http://links.lww.com/QAD/C216). Out of these three samples, two showed a less than two-fold increase of HIV-1 transcriptional activity after anti-CD3/CD28/CD2 activation, compared with the IL-2-only culture (Fig. 3a and Table S2, http://links.lww.com/QAD/C215). Importantly, the two patients with the lowest levels of intracellular HIV-1 transcriptional activity in CD4$^+$ T cells at day 0 showed the lowest levels of in-vitro reactivation from latent infection of HIV-1 within CD4$^+$ T cells, and failed to show intact HIV-1 RNA sequences in the PR/RT and integrase regions (patients ID953 and ID956 in Fig. 3a).

Therefore, transcriptional activity detected by the double-R assay in CD4$^+$ T cells ex vivo, prior to culture, correlated with the production of replication-competent HIV-1 in cultures with maximal activation in vitro.

Consistent with intact sequences, viral outgrowth occurred in the anti-CD3/CD28/CD2-activated sequences from these samples, indicated with ‘F’ within the dot. The patient ID952 was labelled as ‘F’ because of failure to identify intact HIV-1 sequences in IN region; however, we identified the presence of intact HIV-1 sequences in both PR and RT regions (Table S2, http://links.lww.com/QAD/C215). (b) Correlation of HIV-1 RNA transcription in CD4$^+$ T cells after in vitro 3-day culture in the presence of IL2 and T-cell activator, versus HIV-1 RNA transcription per 10^6 CD4$^+$ T cells in peripheral blood, ex vivo, prior to culture. Blip patients are shown by red symbols and without blip-experienced patients are shown by blue symbols.
cultures. HIV-1 was detected in the culture supernatants of the activated CD4+ T cells on days 6–10 (Fig. S7a, http://links.lww.com/QAD/C213), by reverse transcriptase assay. Furthermore, inoculation from the activated CD4+ T-cell cultures into secondary cultures of the CD4+ SupT1 cell line demonstrated productive infection, by reverse transcriptase assay (Fig. S7b, http://links.lww.com/QAD/C213). These results were also confirmed in analyses done on an elite controller showing extremely low HIV-1 transcripts by the double R assay and validated by purified CD4+ T-cell culture activity (Table S4, http://links.lww.com/QAD/C217; Table S5, http://links.lww.com/QAD/C218; Table S6, http://links.lww.com/QAD/C219 and Fig. S8 for further details, http://links.lww.com/QAD/C213).

Discussion

In this study, patients with past viral blips had significantly increased cell-associated HIV-1 transcriptional activity temporally remote from the blip episodes, compared to without blip patients as assessed by the Double-R assay. This significant increase was maintained in follow-up samples, and in some, there was an increase in HIV-1 DNA in PBMCs reflecting an increase in that reservoir. Further, the transcriptional activity in ex-vivo CD4+ T cells had significant correlation with transcriptional activity in in-vitro activated purified CD4+ T cells, harbouring HIV-1 intact HIV-1 sequences and production of replication-competent HIV-1.

Our results are in accord with recent studies and also extend these studies. Several groups have shown that blips may be associated with high-baseline pVL or progression of disease [3,13–16]. A large Swiss longitudinal HIV Cohort Study with 1057 individuals with analysis of total HIV-1 DNA in PBMCs showed clear evidence that blips impede the decline in the HIV-1 latent reservoir [20]. The current study extends these data by showing that regardless of the timing of blip episodes, high-level HIV transcripts are sustained in blip patients. Even some without blip patients at follow-up after prolonged suppression of pVL (>31 months) had borderline levels of HIV-1 transcripts (between 49 and 192 RNA copies/WBCs: definition of borderline in Fig. S3, http://links.lww.com/QAD/C213).

The single copy RNA assay has shown that most clinically suppressed patients have persistent low-level viremia but it has only been used in research and clinical trial contexts [18,21,23–26] and has the disadvantage of a limited range from 0 to 20 copies. The Double-R assay has an improved dynamic range of at least 3 log10 copies making it more suitable for routine clinical monitoring.

Indeed, the Double-R assay has significant advantages over other methodologies in terms of detecting any residual HIV-1 transcripts to reflect HIV-1 promotor activity. The Double-R assay detects HIV-1 promotor activity at both ends of 5’-LTR and 3’-LTR of all transcripts including unspliced and spliced intracellular HIV-1 RNA. It requires only 4–6 ml of whole blood and is not labour-intensive or resource-intensive, in contrast to the previous research-based viral outgrowth or limiting dilution assays. The Double-R assay targets the ‘R’ region, just downstream of the HIV-1 promoter ‘U3’ region in the LTR; it is highly conserved and essential region for HIV-1 integration [50,51]. The U3 region is also essential for HIV-1 transcription but can be modulated by promoter-targeted short hairpin RNA to induce HIV-1 transcriptional gene silencing akin to a latent HIV-1 infection [52–56]. It is likely that when the promoter region in the HIV-1 LTR DNA mutates, HIV-1 would not be able to efficiently produce HIV-1 transcripts.

Our target ‘R’ region sequence to detect HIV-1 transcripts is quite different from previously described targets in unspliced and spliced HIV-1 RNA assays [32,57–59]. Previous studies, that clearly showed a correlation between relatively higher cell-associated HIV RNA levels in PBMC, and more rapid rebound during treatment interruptions, were mainly based on unspliced transcripts but in close to half of patients during ART, these were at or below their assay’s limit of detection [35,36].

Several HIV-1 DNA-based analyses of the memory CD4+ T-cell reservoir [60] using next-generation sequencing (NGS) have reported that the majority (>90%) of the integrated HIV-1 genome is replication-incompetent [27,48,61], with newer approaches now used to identify intact proviral genomes [62–70], particularly in Effector Memory CD4+ T cells [62]. However, the disadvantage of these approaches is that intact virus is present close to, or below, the lower limit of detection [62]. In contrast, our HIV-1 RNA transcript analysis following very efficient activation of CD4+ T cells, with greatly increased intracellular mRNA because of the additional CD2 co-stimulation that increases signalling proximal to the T-cell receptor, separate from CD28 co-stimulation via PI3K/Akt [71,72], had the distinct advantage of being able to detect activatable and intact HIV-1 sequences at very low levels.

Analysis of transcriptional activity by the Double-R assay has some limitations and the current study was not designed to quantify exactly the proportion of replication-competent virus relative to the HIV-1 transcriptional activity in CD4+ T cells. The amount of transcriptional activity detected by the Double-R assay would almost certainly be an overestimate compared with the amount of intact HIV-1 transcripts in the CD4+ T cells in peripheral blood, which will be the subject of future studies.

Our results have several significant consequences, which are hypothesis-generating. First, the variability in
transcriptional activity may relate to particular HIV-1 characteristics or possibly differential effects of antiretroviral drugs. Further research addressing these possibilities is required. Second, measurement of transcriptional activity in peripheral blood CD4⁺ T cells can act as a surrogate marker for the HIV-1 reservoir. Whilst we did not measure the reservoir in other tissues or cell types, a recent study examining many separate tissue reservoirs in autopsy samples suggests that CD4⁺ T cells in the peripheral blood are nevertheless responsible for dissemination of HIV-1 from reservoir sites of tissue replication [73]. Third, the reservoir is dynamic rather than stable, with a bi-directional relationship between blips and reservoir size. Blips may both contribute to, and reflect, activity and size of the reservoir. This residual transcription may eventually represent a new target of antiretroviral therapy.

In conclusion, viral blips are significant. High-sensitivity reservoirs of memory CD4⁺ T cells will enable new scalable opportunities to more closely monitor treated patients experiencing blips, and also study the efficacy of treatments aimed at achieving HIV eradication.

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Conflicts of interest

K.S. receives research funds from Denka Co. Ltd. K.S. is the original inventor under WO2018/045425 (PCT/ AU2017/050974) patent, titled ‘Methods of detecting Lentivirus’ of HIV-1 detection targeting ‘R’ region.

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