The Use of Ultra-Sensitive Molecular Assays in HIV Cure-Related Research

Catherine Kibirige
Johns Hopkins Bloomberg School of Public Health, Baltimore, MD, USA

Abstract

Ultra-sensitive laboratory assays based on the Polymerase Chain Reaction (PCR) are playing an increasingly important role in HIV cure-related research. This article reviews the different assays available and how they have evolved. There is a great need for their standardization and for the establishment of reference reagents and testing algorithms to evaluate potential HIV cure-related treatments.

Keywords: HIV; Eradication; Nucleic acid tests; PCR; Standardization; Proviral DNA; 2-LTR

Introduction

The HIV capsid surrounds two copies of genomic ribonucleic acid (RNA). Replication proceeds with reverse transcription of genomic RNA into a deoxyribonucleic acid (DNA) intermediate. This DNA intermediate is integrated into the host genome where it is referred to as proviral DNA. Unintegrated DNA is present in cells in linear and circular forms. It has a short half-life and disintegrates soon after it is formed [1].

HIV replicates in an error-prone manner that generates a mutation virtually every time the virus replicates. This ongoing mutation allows the emergence of different variants in the host, such as drug-resistant strains or immunological escape mutants. HIV is categorized into HIV type 1 and HIV type 2. HIV-1 is further divided into groups; the major (M) group, the more divergent outlier group (O) group; the non-M, non-O group (N) and the P group. Most HIV infections occur within group M, which is differentiated into subtypes A, B, C, AE, AG, H, J and K. All subtypes and most Circulating Recombinant Forms (CRFs) are found in sub-Saharan Africa. Subtype B is the predominant strain in the US, Europe, Canada and Australia but the prevalence of non-B subtypes in these countries is increasing [1].

The most sensitive FDA approved HIV Nucleic Acid Test (NAT) on the market today is the Abbott Real Time HIV-1 Assay. It has an analytical sensitivity of approximately 25 copies/ml for the 1 ml application. It is approved for the detection of HIV RNA in plasma samples [1]. This assay is not suitable for detecting ultra-low HIV-1 DNA and RNA within host cellular compartments.

Resting memory CD4+ T cells have the ability to harbor latent HIV infection and have been established as an HIV reservoir. The gold standard assay for measuring the frequency of resting memory CD4+ T cells containing latent but replication-competent virus is a viral outgrowth assay that involves harvesting large volumes of blood from an infected patient, sorting purifying resting memory CD4+ T cells and activating limiting dilutions of the cells in culture with phytohemagglutinin (PHA). The cells are co-cultured with CD4+ T lymphoblasts from an HIV-negative donor to amplify any virus released from the cells. A p24 Enzyme-Linked Immunosorbent Assay (ELISA) is used to measure infectious units per million cells (IU/ml) after two to three weeks of culture. The assay is expensive and labor intensive. It requires large volumes of blood, highly skilled staff and specialized laboratory equipment. The assay has a wide coefficient of variation and cannot be performed with tissue biopsies. The assay may not perform well for eradication approaches that produce only small (1 log) reductions in the size of the latent reservoir [2,3].

There are an increasing number of ultra-sensitive laboratory developed PCR-based assays in use that are capable of detecting lower concentrations of HIV RNA and are capable of detecting HIV DNA. The main advantage of most of the PCR-based assays is that they can be performed on small volumes of fresh and frozen samples including blood and tissue. They are relatively faster and simpler to perform when compared to the gold standard assay. These assays are playing an increasing role in HIV cure-related research. A system needs to be devised for their evaluation and standardization.

Currently, most Taqman PCR assays designed to quantify HIV-1 DNA are optimized for Subtype B and may not be suitable for non-B subtypes. HIV-1 molecular assays do not detect HIV-2. There is a lack of sero-conversion panels for non-B HIV-1 and HIV-2 infections [4].

The most recent ultra-sensitive NATs reported in the literature are addressing this problem by basing oligonucleotide sequences on the Long Terminal Repeat (LTR) region of the HIV genome where sequence conservation across subtypes is at its greatest. Examples include a whole blood leukocyte assay - pbs-rtPCR - that is reported to have 100% sensitivity for 2 input copies of DNA even in the presence of high amounts of genomic DNA (1μg) [5]. Another recently reported assay utilizes a non-traditional 13mer probe with a Locked Nucleic Acid (LNA™) modification. This novel nucleic acid analogue incorporates a 2′-O, 4′-C-methylene bridge that restricts flexibility of the ribofuranose ring and locks it into a rigid C3-endo confirmation (Exiqon). LNA™ bases have improved hybridization affinity and biostability effectively raising the melting temperature of an oligonucleotide by 3 to 8°C for each LNA™ base. This allows for the design of shorter Taqman PCR probes that allow researchers to target very short cross-subtype-conserved sequences within the HIV-1 genome and allows for the development of assays that have broader subtype specificity [4]. Yet another recently reported assay targets the LTR region of the HIV genome and uses a Major Grove Binding (MGB) Probe to achieve greater cross-subtype specificity [6].

*Corresponding author: Catherine Kibirige, Johns Hopkins Bloomberg School of Public Health, Baltimore, MD, USA, E-mail: kibirigeck@gmail.com

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Ultra-Sensitive Molecular Assays in HIV Cure-Related Research

There are currently two general approaches to HIV cure-related research. One approach involves strategies such as very early initiation of HAART and the use of agents that reverse latent infection and thus reduce the latent reservoir. The second approach involves the use of agents that re-activate the latent reservoir and then target and reduce the reactivated virus. The successful outcomes of these potential treatments are classified into “sterilization cures” or “functional cures”.

A sterilization cure occurs when HIV-1 DNA and RNA becomes undetectable in samples from the treated patient when analyzed using ultra-sensitive assays and the patient remains free from disease over a prolonged period without antiretroviral treatment. The first reported case of a sterilization cure from HIV was in 2009 [7,8]. A German HIV positive patient with acute myeloid leukemia received a bone marrow transplant from an HIV negative donor homologous for a deletion in the CCR5 gene. The patient ceased HIV treatment soon after the procedure. There was a reconstitution and great improvement in the patient’s immune profile. He currently remains undetectable for HIV nucleic acids and asymptomatic for HIV-related disease [2,7,8].

A functional cure occurs when low grade HIV-1 DNA and/or RNA is detectable in the treated patient but the patient remains free of HIV-associated disease over a prolonged period of time. Elite controllers of HIV infection are considered prototype cases for a functional cure against HIV [2].

Ultra-sensitive viral load or Single Copy Assay (SCA)

The Ultra-sensitive viral load assay or SCA was one of the first ultra-sensitive PCR-based viral load assays developed against HIV that is successfully able to detect down to 1 copy of HIV-1 RNA per ml of plasma [9-11]. It has been used extensively to characterize persistent viremia in patients receiving Highly Active Anti-Retroviral Therapy (HAART). The assay requires 7ml of fresh patient plasma, is labor-intensive, and requires highly skilled technicians and specialized laboratory equipment. Some studies have shown a lack of association between residual viremia as detected by the SCA and viral persistence or immune activation [12-14]. It is therefore generally believed that alternative, more reliable methods that do not require large amounts of fresh patient blood and are thus more versatile will have more utility in HIV eradication and other research efforts.

Total HIV DNA assays

Assays that measure total (unintegrated and integrated) HIV DNA have been in clinical use for a number of years. Standardized International quality controls are available for them. They quantify replication competent and incompetent virus. They are currently the most feasible tool available for large-scale clinical trials and cohort studies [2,15-20].

Despite a general correlation between total DNA and integrated proviral DNA, there are studies that have shown discordance between the two measurements. Some researchers argue that specifically measuring integrated DNA is required as it is a better prognostic marker for treatment failure and success [21].

Proviral HIV DNA assays

Measuring proviral DNA in sort purified resting CD4+ T cells or cells derived from other potential HIV reservoirs is considered a marker for latency. Original assays used linker ligation [22], inverse PCR [23] and nested Alu-PCR [24-27]. Various modifications of these assays now exist including more sensitive methods that employ repetitive sampling to maximize on assay sensitivity [28-30].

These assays can be labor intensive. Multiple methods are used in different studies and the reproducibility across multiple laboratories is unknown [2,25,31-33]. Many HIV-1 genomes within cells are defective and as the assays measure both replication competent and incompetent virus, the assays cannot fully ascertain that a viral reservoir is present [3].

Viral reactivation assays

Detection of low grade viral activation will be important for strategies that rely on the reactivation and targeting of previously latent/recently reactivated infected cells [34-36]. Various assays are available that could be used to detect or assess viral reactivation.

2-Long Terminal Repeat (LTR) HIV DNA assays: A transient increase in 2-LTR circle detection in peripheral lymphocytes is interpreted as an indication of ongoing viral replication [2,34,37-39] and could be used to detect recent reactivation. There is, however, considerable controversy as to whether these unintegrated viral DNA forms play a significant role in the HIV replication cycle [29]. There is skepticism about the reliability of the findings as results vary considerably depending on the laboratory, reagents and context under which the assays are performed [40]. The relationship between 2-LTR circles and residual viral replication has, however, been clearly shown in treatment intensification studies using the integrase inhibitor Raltegravir [41]. This treatment intensification results in a specific though transient increase in episomal DNA in a large number of patients. 2-LTR circle quantification in patients and in vitro models may therefore be a valid marker for virus reactivation and expansion. Standardization of laboratory protocols is required.

Cell-associated HIV RNA assays: These assays measure HIV transcription in productively and latently infected cells. No extracellular HIV RNA is expected in latently infected cells if viral replication has been blocked by HAART or another means. There is a block in nuclear export of multiply spliced RNA and inefficient production of unspliced RNA in these cells. These assays could be used to quantify any increase in transcription following HIV latency reactivation treatment strategies. There are currently very few published studies using these techniques and reproducibility across multiple laboratories is therefore unknown. Assaying for HIV RNA in latent cells may, however, serve as a better marker for reactivation as compared to 2-LTR circles [2,23,37,42-50].

Droplet digital PCR (ddPCR)

Droplet digital PCR (ddPCR) is a nucleic acid detection method that uses the same primers and probes as traditional Taqman RT-PCR. The aqueous reaction mixture is emulsified into thermostable oil and micro-partitioned into picoliter droplets that can contain a single copy or less of target DNA. Following PCR amplification, enumeration of both fluorescing and non-fluorescing droplets allows absolute quantitation of target sequences without relying on the use of standard curves. This greatly reduces the amount of assay “background noise” and allows for greater accuracy and precision. Using 96-well plates, 2 million PCR reactions can be performed simultaneously [51,52]. Emerging data show a significant improvement in precision (~5 fold decrease in assay coefficient of variation) and a greater than 20 fold accuracy improvement in the detection of 2-LTR circles [52]. Some current limitations include unexplained false positives when used in an endpoint PCR format [52] and a maximum sample input
volume of 7.5ul compared to the larger volumes that can be included in traditional PCR [51]. ddPCR therefore requires higher concentrations of DNA substrate in order to maximize sensitivity[51]. Partitioning the reactions into droplets may reduce the inhibition usually experienced when large amounts of DNA are used in PCR, but there is a limit to the amount of target DNA copies that can be loaded per droplet without loss of linearity [51]. Overtly concentrated samples therefore require dilution for best results. Another limitation of this assay format is that, though it can very accurately detect total DNA, it may not be able to detect integrated HIV-1 DNA as precisely. Current PCR-based methods for specifically detecting integrated HIV require a 2-step amplification protocol, the initial step being, amplification across the HIV DNA and host chromosomal DNA junction. ddPCR could conceivably be used at the second step PCR to quantify first-round amplification products but a standard curve would be required and would thus limit precision [51].

This technology provides a viable alternative to traditional PCR and should be optimized and validated further for use in HIV cure-related research.

Endpoint versus kinetic PCR assays

Both endpoint and kinetic PCR protocols have been applied to these assays. Generally endpoint PCR is more sensitive but kinetic or quantitative / real-time PCR provides more accurate quantification. Kinetic PCR can control for false positive signals better than endpoint PCR [28].

Standardizing the Evaluation of HIV Cure-Related Research

PCR-based assays and protocols are allowing for greater and greater sensitivity. Assays can now detect down to 1 input copy of HIV DNA or RNA with greater and greater confidence. The main limitation in sensitivity now lies in sample processing and the number of cell equivalents that can be incorporated into each test, i.e., the number of copies of DNA that can be detected per million cells.

Chun et al. observed a patient with undetectable HIV DNA both in blood and in tissue who nonetheless experienced viremia rebound after ART cessation [53]. This shows that undetectable DNA by molecular assays does not rule out infection [34]. Standardized protocols and testing algorithms therefore need to be developed for evaluating HIV cure-related research.

Some of the considerations that will need to be addressed are as follows:

- Sample collection and handling can greatly affect Nucleic Acid Test Results. It is imperative that specimens be handled, stored and processed correctly and that standardized protocols be devised.
- Specimen concentration and enrichment protocols will need to be utilized to ensure maximum assay sensitivity.
- Researchers need to arrive at a consensus on the possible reservoir sites within a patient and standardize the protocols for the sampling and processing of cells from these sites.
- Specialized staff training and competency assessment will be required when implementing sample collection and ultra-sensitive viral load testing.
- There will be a need for dedicated laboratory equipment with pre and post-amplification areas, reagents and equipment to minimize specimen carry-over and cross-contamination.

- False-positive results and specificity vary depending on the assay used. Low positive results will need to be repeated and assays may need to be used in combination to improve specificity.
- A universal reagent program that provides standardized reagents and panels should be devised.
- A standardized validation protocol needs to be devised for testing the laboratory developed assays that are to be used to evaluate potential treatment strategies.
- The HIV strains circulating in the population will need to be considered when selecting the assays.
- A standardized protocol for re-testing after a certain number of months or years needs to be devised that incorporates the gold standard assay whenever appropriate and possible.
- Clear definitions of what will be considered a “functional” versus a “sterilization” cure should be devised with wording that takes into account the assay limitations.
- As in clinical diagnosis and monitoring of HIV, different settings may require different assay protocols and algorithms.

References

1. Rosenberg ES, Brennan CA, Claessens C, Constantine NT, Murphy G, et al. (2011) Criteria for Laboratory Testing and Diagnosis of Human Immunodeficiency Virus Infection; Approved Guideline. Clinical and Laboratory Standards Institute, Wayne, Pennsylvania 31: 1-89.
2. Leunin SR, Rouzioux C (2011) HIV cure and eradication: how will we get from the laboratory to effective clinical trials? AIDS 25: 885-897.
3. Elisee E, Siliciano RF (2012) Redefining the viral reservoirs that prevent HIV-1 eradication, Immunity 37: 377-388.
4. Li P, Ruel T, Fujimoto K, Hatano H, Yuki S, et al. (2010) Novel application of Locked Nucleic Acid chemistry for a Taqman assay for measuring diverse human immunodeficiency virus type 1 subtypes. J Virol Methods 170: 115-120.
5. Casabianca A, Gori C, Orlandi C, Forbici F, Federico Perno C, et al. (2007) Fast and sensitive quantitative detection of HIV DNA in whole blood leukocytes by SYBR green I real-time PCR assay. Mol Cell Probes 21: 368-378.
6. van der Sluis RM, van Montfort T, Centlivre M, Schopman NC, Cornelissen M, et al. (2013) Quantitation of HIV-1 DNA with a sensitive TaqMan assay that has broad subtype specificity. J Virol Methods 187: 94-102.
7. Hütter G, Nowak D, Miossern M, Gamepola S, Müssig A, et al. (2009) Long-term control of HIV by CCRR Delta32/Delta32 stem-cell transplantation. N Engl J Med 360: 692-698.
8. Hütter G, Schneider T, Thiel E (2009) Transplantation of selected or transgenic blood stem cells - a future treatment for HIV/AIDS? J Int AIDS Soc 12: 10.
9. Palmer S, Maldarelli F, Wiegand A, Bernstein B, Hanna GJ, et al. (2008) Low-level viremia persists for at least 7 years in patients on suppressive antiretroviral therapy. Proc Natl Acad Sci U S A 105: 3879-3884.
10. Shiu C, Cunningham CK, Greathough T, Muresan P, Sanchez-Merino V, et al. (2009) Identification of ongoing human immunodeficiency virus type 1 (HIV-1) replication in residual viremia during recombinant HIV-1 poxvirus immunizations in patients with clinically undetectable viral loads on durable suppressive highly active antiretroviral therapy. J Virol 83: 9731-9742.
11. Maldarelli F, Palmer S, King MS, Wiegand A, Polis MA, et al. (2007) ART suppresses plasma HIV-1 RNA to a stable set point predicted by pretherapy viremia. ProcNatlAcadSciUS A 104: 9971-9975.
12. Muir D, White D, King J, Verlander N, Pillay D (2000) Predictive value of the ultrasensitive viral load assay in clinical practice. J Med Virol 61: 411-416.
13. Hatano H, Jain V, Hunt PW, Lee TH, Sinclair E, et al. (2012) Cell-Based Measures
of Viral Persistence Are Associated With Immune Activation and Programmed Cell Death Protein 1 (PD-1)-Expressing CD4+ T Cells. J Infect Dis.

14. Vallejo A, Gutierrez C, Hernandez-Novoa B, Diaz L, Madrid N, et al. (2012) The effect of intensification with raltegravir on the HIV-1 reservoir of latently infected memory CD4 T cells in suppressed patients. AIDS 26: 1885-1894.

15. Chun TW, Nickle DC, Justement JS, Meyers JH, Roby G, et al. (2008) Persistence of HIV in gut-associated lymphoid tissue despite long-term antiretroviral therapy. J Infect Dis 197: 714-720.

16. Avetland-Fenoll V, Chaix ML, Blanche S, Burgard M, Floch C, et al. (2009) LTR real-time PCR for HIV-1 DNA quantitation in blood cells for early diagnosis in infants born to sero-positive mothers treated in HAART area (ANRS CO 01). J Med Virol 81: 217-223.

17. Bouony O, Mannioui A, Sellier P, Roucair C, Durand-Gasselin L, et al. (2010) Effect of a short-term HAART on SIV load in macaque tissues is dependent on time of initiation and antiviral diffusion. Retrovirology 7: 98.

18. Denton PW, Garcia JV (2005) Novel humanized murine models for HIV research. Curr HIV/AIDS Rep 6: 13-19.

19. Beloukas A, Paraskevis D, Haida C, Syssas V, Hatzakis A (2009) Development and assessment of a multiplex real-time PCR assay for quantification of human immunodeficiency virus type 1 DNA. J Clin Microbiol 47: 2194-2199.

20. Domadula G, Zhang H, Shetty S, Pomerantz RJ (1999) HIV-1 virions produced from replicating peripheral blood lymphocytes are more infective than those from nonproliferating macrophages due to higher levels of intravirion reverse transcripts: implications for pathogenesis and transmission. Virology 253: 10-18.

21. Carr JM, Cherry KM, Coolen C, Davis A, Shaw D, et al. (2007) Development of methods for coordinate measurement of total cell-associated and integrated human immunodeficiency virus type 1 (HIV-1) DNA forms in routine clinical samples: levels are not associated with clinical parameters, but low levels of integrated HIV-1 DNA may be prognostic for continued successful therapy. J Clin Microbiol 45: 1288-1297.

22. Van de Graaff N, Kumar R, Burrell CJ, Li P (2001) Kinetics of human immunodeficiency virus type 1 (HIV) DNA integration in acutely infected cells as determined using a novel assay for detection of integrated HIV DNA. J Virol 75: 11253-11260.

23. Chun TW, Carnall J, Finzi D, Shen X, DiGiuseppe JA, et al. (1997) Quantification of latent tissue reservoirs and total body viral load in HIV-1 infection. Nature 387: 183-188.

24. Butler SL, Hansen MS, Bushman FD (2001) A quantitative assay for HIV DNA integration in vivo. Nat Med 7: 631-634.

25. O'Doherty U, Swigard WW, Jeyakumar D, McGain D, Malim MH (2002) A sensitive, quantitative assay for human immunodeficiency virus type 1 integration. J Virol 76: 10942-10950.

26. Brussel A, Deleclu X, Sonigo P (2005) Alu-LTR real-time nested PCR assay for quantifying integrated HIV-1 DNA. Methods Mol Biol 304: 139-154.

27. Brussel A, Sonigo P (2003) Analysis of early human immunodeficiency virus type 1 DNA synthesis by use of a new sensitive assay for quantifying integrated provirus. J Virol 77: 10119-10124.

28. Yu J, Wu TL, Liszewska MK, Dai J, Swigard WW, et al. (2008) A more precise HIV integration assay designed to detect small differences find lower levels of integrated DNA in HAART treated patients. Virology 379: 78-86.

29. Kumar R, Van de Graaff N, Mundy L, Burrell CJ, Li P (2002) Evaluation of PCR-based methods for the quantitation of integrated HIV-1 DNA. J Virol Methods 105: 233-246.

30. Agosto LM, Yu J, Dai J, Kalentsis R, Monie D, et al. (2007) HIV-1 integrates into resting CD4+ T cells even at low inoculums as demonstrated with an improved assay for HIV-1 integration. Virology 360: 60-72.

31. Graf EH, Meexas AM, Yu J, Shaheem F, Liszewski MK, et al. (2011) Elite suppressors harbor low levels of integrated HIV DNA and high levels of 2LTR circular HIV DNA compared to HIV+ patients on and off HAART. PLoS Pathog 7: e1001300.

32. Sonza S, Maerz A, Deacon N, Meanger J, Mills J, et al. (1996) Human immunodeficiency virus type 1 replication is blocked prior to reverse transcription and integration in freshly isolated peripheral blood monocytes. J Virol 70: 3863-3869.

33. Lewis SR, Murray JM, Solomon A, Wightman F, Cameron PU, et al. (2008) Virological determinants of success after structured treatment interruptions of antiretrovirals in acute HIV-1 infection. J Acquir Immune Defic Syndr 47: 140-147.

34. Frater J (2011) New approaches in HIV eradication research. Curr Opin Infect Dis 24: 593-596.

35. Rasmussen TA, Schmelz Sggaard O, Brinkmann C, Wightman F, Lewis SR, et al. (2013) Comparison of HDAC inhibitors in clinical development: Effects on HIV production in latently infected cells and T-cell activation. Hum Vaccin Immunother 9.

36. Shan L, Deng K, Shroff NS, Durand CM, Rabli SA, et al. (2012) Stimulation of HIV-1-specific cytolytic T lymphocytes facilitates elimination of latent viral reservoir after virus reactivation. Immunity 36: 491-501.

37. Sharkey ME, Teo I, Greenough T, Sharova N, Luzuriaga K, et al. (2000) Persistence of episomal HIV-1 infection intermediates in patients on highly active anti-retroviral therapy. Nat Med 6: 76-81.

38. Sharkey M, Triques K, Kuritzkes DR, Stevenson M (2005) In vivo evidence for instability of episomal human immunodeficiency virus type 1 DNA. J Virol 79: 5203-5210.

39. Sharkey M, Babic DZ, Greenough T, Gulick R, Kuritzkes DR, et al. (2011) Episomal viral cDNAs identify a reservoir that fuels viral rebound after treatment interruption and that contributes to treatment failure. PLoS Pathog 7: e1001303.

40. Durand CM, Blankson JN, Slicciaro RF (2012) Developing strategies for HIV-1 eradication. Trends Immunol 33: 554-562.

41. Buzón MJ, Massanella M, Libbre JM, Esteve A, Dahl V, et al. (2010) HIV-1 replication and immune dynamics are affected by raltegravir intensification of HAART-suppressed subjects. Nat Med 16: 450-455.

42. Pasternak AO, Jurriaans S, Bakker M, Prins JM, Berkhout B, et al. (2009) Cellular levels of HIV unspliced RNA from patients on combination antiretroviral therapy with undetectable plasma viremia predict the therapy outcome. PLoS One 4: e8490.

43. Vesana M, Markowitz M, Cao Y, Ho DD, Saksela K (1997) Human immunodeficiency virus type-1 mRNA splicing pattern in infected persons is determined by the proportion of newly infected cells. Virology 236: 104-109.

44. Zhang L, Ramratnam B, Temmer-Raczk K, He Y, Vesana M, et al. (1999) Quantifying residual HIV-1 replication in patients receiving combination antiretroviral therapy. N Engl J Med 340: 1605-1613.

45. Yukl SA, Gianella S, Sinclair E, Epling L, Li Q, et al. (2010) Differences in HIV burden and immune activation within the gut of HIV-positive patients receiving suppressive antiretroviral therapy. J Infect Dis 202: 1553-1561.

46. Burgard M, Iozep J, Duman B, Tamalet C, Descamps D, et al. (2000) HIV RNA and HIV DNA in peripheral blood mononuclear cells are consistent markers for estimating viral load in patients undergoing long-term potent treatment. AIDS Res Hum Retroviruses 16: 1939-1947.

47. Christopherson C, Kidane Y, Conway B, Krowka J, Sheppard H, et al. (2000) PCR-Based assay to quantify human immunodeficiency virus type 1 DNA in peripheral blood mononuclear cells. J Clin Microbiol 38: 630-634.

48. Lewis SR, Vesana M, Kostrikis L, Hurley A, Duran M, et al. (1999) Use of real-time PCR and molecular beacons to detect virus replication in HIV immunodeficiency virus type-1 infected individuals on prolonged effective antiretroviral therapy. J Virol 73: 8099-8103.

49. Yerly S, Perneger TV, Vora S, Hirschel B, Perrin L (2000) Decay of cell-associated HIV-1 DNA correlates with residual replication in patients treated during acute HIV-1 infection. AIDS 14: 2805-2812.

50. Schmid A, Gianella S, von Wyl V, Metzker JN, Scherrher AU, et al. (2010) Profound depletion of HIV-1 transcription in patients initiating antiretroviral therapy during acute infection. PLoS One 5: e13310.

51. Henrich TJ, Gallien S, Li LZ, Pereyra F, Kuritzkes DR (2012) Low-level detection and quantitation of cellular HIV-1 DNA and 2LTR circles using droplet digital PCR. J Virol Methods 186: 68-72.

52. Strain MC, Lada SM, Luong T, Rought SE, Gianella S, et al. (2013) Highly precise measurement of HIV DNA by droplet digital PCR. PLoS One 8: e55943.

53. Chun TW, Justement JS, Murray D, Hallahan CW, Maenzen J, et al. (2010) Rebound of plasma viremia following cessation of antiretroviral therapy despite profoundly low levels of HIV reservoir: implications for eradication. AIDS 24: 2803-2808.