Title: Improved Single Copy Assays for Quantification of Persistent HIV-1 Viremia in Patients on Suppressive Antiretroviral Therapy

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Running Title: Improved Plasma HIV-1 RNA Single Copy Assays

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Abstract

A quantitative real-time PCR (qRT-PCR) assay with single-copy sensitivity targeting HIV-1 gag RNA (gSCA) has been used widely to quantify plasma viremia below the limit of detection of clinical assays in patients on effective antiretroviral therapy (ART), but viral RNA in 15-30% of samples amplifies inefficiently because of primer/probe mismatches. We sought to develop improved single-copy assays with increased sensitivity by improving nucleic acid recovery, designing qRT-PCR primers and probe for a highly conserved region of integrase in the HIV-1 pol gene (iSCA), and increasing the plasma volume tested (Mega-iSCA). We evaluated gSCA versus iSCA in paired plasma samples from 10 consecutive patients with viremia >1000 copies/mL and 25 consecutive patients on suppressive ART. Three of 10 viremic samples amplified inefficiently with gSCA compared to the Roche Cobas Ampliprep/TaqMan 2.0, whereas all 10 samples amplified efficiently with iSCA. Among 25 samples from patients on suppressive ART, 8 of 12 samples that were negative for HIV-1 RNA by gSCA had detectable HIV-1 RNA by iSCA, and iSCA detected 3-fold or higher HIV-1 RNA levels compared to gSCA in 10 of 25 samples. Large volume plasma samples (>20 mL) from 7 patients were assayed using Mega-iSCA, and HIV-1 RNA was quantifiable in 6, including 4 of 5 that were negative by standard volume iSCA. These improved assays with superior sensitivity will be useful for evaluating whether in vivo interventions can reduce plasma viremia and for assessing relationships between residual viremia and other virologic parameters including the inducible proviral reservoir.
Introduction

Infection with human immunodeficiency virus type 1 (HIV-1) leads to high levels of cell-free virions in plasma within days following infection(1). Intensive study of HIV-1 plasma viremia has contributed to the current understanding of HIV-1 pathogenesis and has guided the development of effective antiretroviral therapy (ART). In chronically-infected untreated patients, the amount of virus in plasma reflects the number of productively infected cells and is a strong independent predictor of HIV-1 disease progression to AIDS and death(2, 3). Following the discovery of potent antiretroviral drugs, the decay dynamics of plasma viremia and productively infected cells was possible down to the limit of detection of available HIV-1 RNA assays(4-6). More sensitive qualitative assays revealed the persistence of viremia on combination antiretroviral therapy (ART) (7), indicating that ART alone does not eliminate plasma viremia.

Further studies using a two-step quantitative real-time polymerase chain reaction (qRT-PCR) assay with single copy sensitivity targeting HIV-1 gag (gSCA)(8), revealed that plasma viremia between 1 and 3 copies of HIV-1 RNA per milliliter of plasma persists in most patients independent of the ART regimen used to suppress viremia(9). Additional studies revealed three phases of plasma HIV-1 RNA decay, reflective of HIV-infected populations of cells with different half-lives, followed by a fourth phase of decay with an infinite half-life during which HIV-1 RNA levels average 1 copy per milliliter for at least 7 years following initiation of ART(10). Whether the persistent viremia on ART represents infectious virus remains unknown, but in some instances, viral sequences in plasma may match sequences of infectious virus recovered from resting CD4+ T cells(11).

gSCA has also been used in a number of settings to evaluate whether clinical interventions can affect residual viremia. Several investigators have used gSCA as an
endpoint in ART intensification studies to evaluate whether addition of raltegravir or
other antiretroviral agents to suppressive ART would reduce persistent low-level plasma
virema, and found that there was no effect on plasma viremia(12-14). In a recent study,
incomplete adherence to ART was also found to be associated with higher levels of
residual viremia(15). Another study evaluated cellular HIV-1 RNA and plasma viremia
following \textit{in vivo} administration of vorinostat (SAHA), and found that cellular HIV-1 RNA
was increased without a concomitant increase in plasma viremia as measured with
gSCA(16). Levels of plasma viremia as measured by gSCA have also been evaluated as
a potential biomarker of the size of the infectious reservoir in resting CD4+ T cells(17).
Despite its widespread use, the original gSCA assay has limitations. First, the
extraction method to isolate HIV-1 RNA published in 2003 may not maximize the
recovery of nucleic acid from plasma. Second, the qRT-PCR primers and probe were
designed to bind a conserved region of \textit{gag} from HIV-1 subtype B variants from the
United States based on sequences available in 2001-2002, but new sequences in the
Los Alamos database indicate that there are more highly conserved regions elsewhere
in the HIV-1 genome. Third, the volume of plasma extracted in the original description of
the assay was limited to 7 milliliters. In theory, sensitivity could be improved by assaying
larger plasma volumes, although this has not been shown. We sought to address these
limitations of gSCA by improving nucleic acid recovery, designing new primers and
probe in a more conserved region of the HIV-1 genome, and increasing the volume of
plasma assayed to potentially reduce censoring of data below the limit of detection,
thereby improving the ability to detect an effect of interventions on residual viremia.
Materials and Methods

Clinical Specimens

All blood donors were patients at the University of Pittsburgh AIDS Center for Treatment, and provided written informed consent. The University of Pittsburgh Institutional Review Board approved the study. Donors were not specifically selected for subtype B infection, and were recruited consecutively from the Center. Two groups of blood donors were recruited for this study: 1) donors with plasma viremia >1000 copies/mL as measured by Roche COBAS AmpliPrep/COBAS TaqMan, v2.0 (Roche TM2.0); and 2) donors with viremia suppressed to <20 HIV RNA copies/mL on combination antiretroviral therapy (Roche TM2.0). Donors underwent phlebotomy (100-180 mL) and blood was processed within 4 hours. Plasma was separated by centrifugation of whole blood at 400 x g for 10 minutes, followed by removal of the plasma layer and a second centrifugation of plasma at 400 x g. Aliquots of 1.5 milliliters of plasma were stored at -80°C.

Isolation of Nucleic Acid

Total nucleic acid was isolated from plasma samples using the previously reported method (v1.0)(8) or using modifications (v2.0) reported by Venneti et al.(18). A total of 4 aliquots of plasma (~6 milliliters) from each donor were thawed in a room temperature water bath. Samples from the same patient were pooled, split into two separate samples for gSCA and iSCA, and a known quantity of internal standard (Replication Competent ALV LTR with a Splice adaptor(8, 19, 20), RCAS) was spiked into each sample. The samples were then spun at 1600 x g for 15 minutes at 4°C to pellet debris (e.g. fibrin and other insoluble complexes; control experiments showed that this centrifugation improved HIV-1 and RCAS recovery). Plasma supernatants were then transferred to Seton Easy Seal tubes for ultracentrifugation, and the remaining volume in
the tube was filled with Tris-buffered saline. Virions were then pelleted by ultracentrifugation at 170,000 x g for 30 minutes in a Sorvall T-1270 rotor. Supernatants were then removed, and 100 μL 3M guanidinium hydrochloride (GuHCl) supplemented with 50 mM Tris-HCl, 1 mM CaCl₂ and 1 mg/mL proteinase K was added, followed by incubation at 42°C for 1 hour. Following this incubation, 400 μL of ~6M guanidinium thiocyanate (GuSCN) supplemented with 50 mM Tris-HCl, 1 mM EDTA and 600 μg/mL of glycogen was added, followed by incubation for an additional 10 minutes at 42°C. Next, 500 μL of 100% isopropanol was added to precipitate nucleic acids and the samples were centrifuged at 21,000 x g for 10 minutes. Supernatants were removed, 750 μL of 70% ethanol wash was added, followed by centrifugation at 21,000 x g for 10 minutes. The supernatant was removed, pellets were air-dried and resuspended in 55 μL of 5 mM Tris-HCl supplemented with 1 μM dithiothreitol and 1000 units/mL of RNasin. Test plasma samples were always processed in parallel with HIV negative plasma and low copy number plasma containing 5 HIV-1 RNA copies per milliliter (Rush University, Virology Quality Assurance Lab). HIV-1 RNA was not detected in any of the HIV-1 negative plasma samples (N >50) throughout the study.

Reverse transcriptase quantitative polymerase chain reaction

A two-step qRT-PCR reaction was used to quantify HIV-1 RNA in plasma. Complimentary DNA (cDNA) was reverse transcribed as previously described from 10μL of extract in triplicate in 30 μL reactions for HIV-1 RNA and duplicate reactions for RCAS; 5 μL of extract was used as input in the cDNA reaction mix without reverse transcriptase to confirm that HIV-1 DNA was absent. For real-time PCR, the LightCycler 480 Probes Master ready-made master mix was used at a 0.7x concentration in a total volume of 50 μL with either 400 nM forward and reverse primers and 200 nM probe (iSCA) or 600 nM forward and reverse primers and 100 nM probe (gSCA and RCAS).
Cycling parameters for real-time PCR were 95°C for 5 minutes, followed by 95°C for 10 seconds and 60°C for 1 minute, for 45 cycles of amplification. A standard curve was generated for each assay by serial 3-fold dilution of HIV-1 RNA transcripts that were characterized by optical density at 260 nM and serial end-point dilution to 1 copy per qRT-PCR reaction. Consistent with a theoretical Poisson distribution, we observed 40-70% of reactions being positive at 1 input copy of HIV-1 RNA transcripts. HIV-1 RNA transcripts were transcribed in vitro from a T7 expression plasmid containing a 1.7 kb insert containing the 3’ end of pol (bases 4186-5913 of the HIV-1 reference sequence HXB2) generated by standard PCR from the plasmid xxLAI(21) (GenBank accession number: K02013) using the primers forward: 5’-AACAAGTAGATAAATTAGTCA-3’ and reverse: 5’-TTACAATAGCAAGTGGTA-3’ at 600 nM each. Transcripts were purified and unincorporated nucleotides were removed as previously described(22). The primer and probe sequences have been previously reported for gSCA and RCAS(8) and for iSCA(22).

Mega-iSCA

To lower the limit of HIV-1 RNA detection, we assayed large volumes of plasma (>20 milliliters) by iSCA (Mega-iSCA). Plasma was divided into parallel replicate assays (4-5 mL of plasma being extracted for each replicate), and assayed as described above by iSCA. The total copy number in each sample was summed, multiplied by a correction factor to control for the proportion of the extract assayed for HIV-1 RNA, and divided by the total volume of plasma assayed to determine the number of copies of HIV-1 RNA per milliliter of plasma detected by Mega-iSCA.
Sequence and Statistical Analyses

To identify nucleotide sequence conservation in particular loci of HIV-1, full-length nucleotide web alignments of all subtypes (A-K without recombinants) containing only 1 sequence per patient were downloaded in FASTA form from the HIV Sequence Database at Los Alamos National Laboratory (http://www.hiv.lanl.gov/). To manipulate alignments, identify conserved regions, and generate sequence logos, the statistical program R(23) with the packages seqinr(24), Biostrings(25) and seqLogo(26) were used. Average pairwise distance was calculated using the package ape(27) in R.

Statistical analysis was performed using R, and all tests for significance were performed using a two-sided $\alpha=0.05$ as significant. For comparisons of assays on a continuous scale, negative HIV-1 RNA results were assigned a value equal to 50% of the assay detection limit. Figures were generated in R using the packages ggplot2(28) and scales(29).
Results

Analysis of HIV-1 Nucleotide Sequences for Highly Conserved Regions

The previously reported gSCA primers and probe were originally designed to bind to a conserved region of gag in subtype B HIV-1. Inefficient amplification of HIV-1 RNA in 15-30% of plasma samples, compared to FDA-cleared assays, led us to search for more conserved regions of HIV-1. Sequences (1360 total) from all HIV-1 subtypes (A-K) available in the Los Alamos Database (2012) were downloaded and a consensus sequence for subtype B was built by selecting the most prevalent nucleotides at each position from the 675 subtype B sequences available.

To identify the most highly conserved regions of the HIV-1 genome, the fraction of nucleotides at each position in 200 base pair fragments of individual sequences that matched the consensus was determined (Figure 1). The most highly conserved regions were in gag and at the 3’ end of pol (specifically, the region containing integrase), both of which had >95% sequence identity with the subtype B consensus. The integrase region (98.1%) was more highly conserved than the gag region (96.4%). Primer and probe binding sites in integrase were selected and sequence logos were generated (Figure 2) to compare the extent of mismatches for the new integrase primer/probe set versus the gag primer/probe set for their respective targets. The results of the sequence logos are in agreement with Figure 1 in that the primer/probe binding sites in the selected integrase region are more highly conserved than in the previously selected gag site, especially for the probe. In addition, the diversity (average pairwise distance, APD) calculated for the integrase primer/probe binding sites (2.6%) was significantly lower than for the gag primer/probe binding sites (3.1%; p-value between APDs <1x10^{-5}).

These analyses suggest that targeting the integrase region with a single-copy assay (iSCA) would result in a lower frequency of inefficient amplification compared with gSCA. When considering all subtypes, there was excellent agreement in sequence
conservation in the integrase region, with the exception of a one-nucleotide substitution present in subtype C present in the forward primer-binding site.

Improvement of Total Nucleic Acid Recovery

We next sought to improve the recovery of nucleic acid from plasma. The original gSCA nucleic acid extraction consisted of pelleting of virions by ultracentrifugation, digestion of proteins with proteinase K in water, and addition of the chaotropic agent guanidinium thiocyanate (GuSCN), followed by precipitation and washing of nucleic acid(8). We modified several steps in this extraction protocol as follows: i.) after ultracentrifugation, the sample was incubated with guanidinium hydrochloride (GuHCl) supplemented with Proteinase K, Tris-HCl and CaCl₂ at 42°C as described in the methods; ii.) GuSCN supplemented with Tris-HCl, EDTA and glycogen was then added; iii.) centrifugation times following isopropanol precipitation and ethanol wash were shortened to 10 minutes.

To determine whether the modified extraction method (v2.0) improves recovery of HIV-1 RNA from plasma compared with the original extraction method (v1.0), we assessed recovery of HIV-1 RNA from each extraction method in parallel for paired samples. To generate paired samples, plasma from an HIV-1 positive viremic donor was diluted 100-fold and 1000-fold with Tris-buffered saline containing 10% fetal bovine serum (Table 1). In replicate assays, HIV-1 RNA recovery was on average 3.0-fold higher at the 1:100 dilution and 2.0-fold higher at the 1:1000 dilution with extraction method v2.0 compared with v1.0. We also evaluated the recovery of the RCAS internal standard spiked into HIV-1 negative plasma or plasma containing 5 copies of HIV-1 RNA per milliliter (Virology Quality Assurance Lab, Rush University). In paired samples, we found that v2.0 recovered 1.4-1.8 fold more RCAS RNA and 1.6 fold more HIV-1 RNA from the VQA standards.
Performance of iSCA vs. gSCA in Viremic Patients

The performance of the gSCA and iSCA primer/probe systems was evaluated by quantifying HIV-1 RNA in duplicate plasma samples from 10 consecutive donors with plasma HIV-1 RNA values ranging from 1,202 to 382,189 copies/mL by Roche TM2.0. By gSCA, 1 of 10 patients had undetectable HIV-1 RNA and a total of 3 of 10 patients had HIV-1 RNA levels less than 10% of the value determined by Roche TM2.0 (Table 2), consistent with prior reports that HIV-1 RNA from 15-30% of patient samples does not amplify efficiently. By iSCA, all 10 patients had HIV-1 RNA levels within 10-fold of the Roche TaqMan platform (p=0.038, exact binomial test for difference in amplification efficiency between iSCA and gSCA). Levels of plasma viremia as measured with iSCA and gSCA did not correlate (Figure 3A; \( \rho = 0.54, p=0.11 \)), largely due to the 3 samples that were not efficiently amplified by gSCA.

Evaluation of iSCA vs. gSCA in Patients on Suppressive ART

The goal of this study was to improve the ability to detect and quantify HIV-1 RNA in patients on suppressive ART with plasma HIV-1 RNA below the limit of detection of clinical assays. To this end, we tested a panel of duplicate plasma samples from 25 consecutive donors on suppressive ART (Table 3, Table S1). HIV-1 RNA was detected in 17 of 25 samples by iSCA versus 13 of 25 samples by gSCA. A total of 8 samples had detectable HIV-1 RNA by iSCA alone, whereas 4 had detectable HIV-1 RNA by gSCA alone. HIV-1 RNA values were \( \geq 3 \)-fold by iSCA compared with gSCA in 10 of 25 samples, versus 5 of 25 being \( \geq 3 \)-fold higher by gSCA than iSCA (p=0.021, exact binomial test comparing fractions of samples that were \( \geq 3 \)-fold higher by either assay).

The median HIV-1 RNA level of all 25 samples determined by iSCA was 2.0 copies/mL (interquartile range [IQR] 0.35 to 4.9 copies/mL) versus 1.3 copies/mL (IQR: 0.30 to 3.3 copies/mL).
copies/mL) with gSCA (Table 3, Table S1). HIV-1 RNA values measured by iSCA and gSCA showed a trend toward a positive correlation but this was not significant (Figure 3B; \( p=0.36, p=0.077 \)). However, when considering all patient samples assayed (viremic and on suppressive ART), there was a significant correlation between iSCA and gSCA (Figure 3C; \( p=0.64, p<0.001 \)).

Development and Validation of Mega-iSCA

The input volume of plasma assayed currently limits the sensitivity of single copy assays, making it uncertain whether patients with HIV-1 RNA below the SCA detection limit are truly aviremic. We hypothesized that by increasing the volume of plasma assayed and thereby lowering the limit of detection, we would be able to detect viremia in more patients, including those with HIV-1 RNA below the limit of detection by standard-volume iSCA. To test this hypothesis, we assayed large volumes of plasma (20-35 mL) from 7 donors including 5 with undetectable HIV-1 by standard-volume (i.e., 3.0 mL of plasma) iSCA (Table 4). Assuming that 30 mL of plasma has been assayed, the theoretical limit of detection with large volume iSCA (Mega-iSCA) would be 0.06 copies of HIV-1 RNA per milliliter, about 10-fold less than standard iSCA. Using Mega-iSCA, HIV-1 RNA was detected in 6 of 7 donors, including 4 of 5 with standard-volume iSCA results below the limit of detection (Table 4).
Discussion

We have developed improved HIV-1 single-copy assays with improved sensitivity to quantify the persistence of plasma viremia in patients on suppressive ART. The original single-copy assay targeting HIV-1 gag sequences (gSCA) has been successfully applied in many studies to gain insight into the persistence of viremia on ART, although inefficient amplification of viral sequences in 15-30% of patients and consequent false negative results have been limitations. We have evaluated these limitations and modified the assay to improve amplification efficiency and lower the limit of detection.

First, we found that isolation of nucleic acid by sequential GuHCl/proteinase K incubation and GuSCN addition, followed by precipitation of nucleic acid recovered more RCAS and HIV-1 RNA in test panels. The higher recovery observed is likely due to the use of GuHCl/proteinase K during the initial incubation, after the virions have been pelleted, rather than using the original extraction method with water and proteinase K. Use of the chaotropic agent GuHCl with proteinase K likely increases the digestion of proteins in protein/nucleic acid complexes as compared with proteinase K in water alone. GuHCl and EDTA also likely inactivate RNases.

Second, to improve amplification efficiency, we targeted a more conserved region of the HIV-1 genome. The high rate of nucleotide substitution in HIV-1 has been extensively studied(30), and contributes to difficulties with sensitivity of nucleic acid-based diagnostics for HIV-1(31). Using sequences in the Los Alamos database, we uncovered a region at the 3’ end of the pol gene that was more highly conserved than gag and that was also amenable to primer and probe design for qRT-PCR. Selection and use of this integrase region of the HIV-1 genome for qRT-PCR was effective, in that we eliminated primer/probe mismatches in the test panel of 10 viremic patients.
With the improved performance of iSCA for samples from patients with viremia detectable by Roche TM2.0, we proceeded with a head-to-head comparison of iSCA versus gSCA in samples from patients on suppressive ART, and generally with HIV-1 RNA <20 copies/mL by Roche TM2.0. In the suppressed patients, iSCA again had better performance than gSCA as evidenced by: i) detecting viremia in more patients (17 of 25 detectable by iSCA versus 13 of 25 detectable by gSCA with 8 of 12 not detected by gSCA detected by iSCA); ii) measuring ≥3-fold higher HIV-1 RNA values in 10 of 25 with iSCA versus 5 of 25 with gSCA; and iii) trending towards measuring a higher mean level of residual viremia (2.0 cps/mL for iSCA vs. 1.3 cps/mL for gSCA). The median levels of HIV-1 RNA detected in samples from suppressed patients were low enough that stochastic variation in the number of virions between samples may have increased the variability in each assay. Other limitations of our study are the relatively small number of patients evaluated and the likelihood that only subtype B samples were tested. Nevertheless, the combination of a new extraction method and new primers and probe improved assay performance for iSCA compared with gSCA.

Finally, we assessed whether the sensitivity of iSCA could be improved by increasing the volume of plasma assayed. The typical limit of detection for gSCA/iSCA is 0.6 copies of HIV-1 RNA per milliliter of plasma, assuming that 3.0 mL of plasma have been assayed. It is possible that patients have levels of residual viremia that are below this limit of detection and thus score as undetectable by iSCA using the standard volume of plasma. Our data support this hypothesis, as we have shown that 4 of 5 patients with undetectable viremia by standard iSCA had detectable viremia by Mega-iSCA. Mega-iSCA is ≥10-fold more sensitive than iSCA, and will be useful for future evaluations of residual viremia and changes in residual viremia following interventions. Plasma volumes of 20-30 mL can be easily obtained from large-volume blood draws (75-120 mL), which is less blood volume than that required for an assay of inducible HIV-1.
reservoirs from resting CD4+ T cells (32). Furthermore, the same whole blood sample can be used for both assays. Two studies have reported a positive correlation between the levels of persistent viremia measured by gSCA and infectious virus recovery from resting CD4+ T cells (17, 33), but another study found no relationship between gSCA and infectious virus recovery (32). The correlation between gSCA and levels of infectious virus suggests that HIV-1 RNA in the plasma may originate from stochastic reactivation of latently infected cells, and that changes in levels of HIV-1 RNA in plasma following an intervention may be reflective of changes in the size of the latent reservoir. Future studies should seek to further elucidate the relationship between persistent plasma viremia and the infectious virus reservoir.

In summary, we have developed and validated an improved single-copy assay (iSCA) with an improved ability to detect and quantify low levels of plasma viremia. iSCA will undoubtedly be useful for future studies, such as an endpoint of clinical trials with interventions aimed at reducing the size of the HIV-1 reservoir. As the field of HIV-1 cure research continues to evolve, the ability to accurately and precisely quantify residual plasma viremia will be important to evaluate whether novel treatments have caused a change in viremia and whether HIV-1 RNA in plasma remains detectable by the most sensitive means possible after potentially curative interventions.
References


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**Figure Legends**

**Figure 1.** Proportion of nucleotides in 200 base pair fragments that match the HIV-1 2012 subtype B consensus sequence. A consensus sequence was generated from a total of 675 aligned full-length subtype B sequences from the Los Alamos Sequence Database, and the number of bases in 200 base pair fragments that matched the consensus sequence was calculated. The most highly conserved regions are present where the original gSCA primers were designed and in the 3' region of *pol*, where the iSCA primers and probe target.

**Figure 2.** Comparison of Sequence Logos for the forward (A), reverse (B) and probe (C) binding regions for iSCA and gSCA from 675 subtype B sequences from the Los Alamos Sequence Database. The iSCA system has fewer mismatches compared to the gSCA system, suggesting that the primers and probe from the iSCA system would efficiently amplify HIV-1 RNA in a higher proportion of patient samples compared to the gSCA system.

**Figure 3.** Relationship between plasma HIV-1 RNA measured with gSCA versus iSCA. (A) There is no significant correlation between HIV-1 RNA values measured by gSCA and iSCA in the 10 samples from viremic patients because of the 3 samples with inefficient amplification by gSCA. The dashed lines indicate the limit of quantification of Roche TM2.0 (20 cps/mL). (B) There is a non-significant trend toward a correlation between HIV-1 RNA values measured by iSCA vs gSCA in the 25 plasma samples from patients on suppressive ART. The dotted lines indicate the limit of detection of iSCA and gSCA (0.6 cps/mL). (C) HIV-1 RNA values are significantly correlated between iSCA and gSCA for all 35 patients evaluated in this study. The dashed lines indicate the
limit of quantification of Roche TM2.0 (20 cps/mL), and the dotted lines indicate the limit of detection of iSCA and gSCA (0.6 cps/mL).
Table 1. Improved recovery of plasma HIV-1 RNA using a new extraction method

<table>
<thead>
<tr>
<th>Sample</th>
<th>Extraction Method</th>
<th>Number of Replicates</th>
<th>Expected HIV-1 RNA (copies per milliliter)</th>
<th>Mean HIV-1 RNA Detected (copies per milliliter)</th>
<th>Mean Percent Expected HIV-1 RNA Detected</th>
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<td>Viremic Plasma Diluted 1:100</td>
<td>v1.0</td>
<td>4</td>
<td>271</td>
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<tr>
<td></td>
<td>v2.0</td>
<td>4</td>
<td>292</td>
<td>108%</td>
<td></td>
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<tr>
<td>Viremic Plasma Diluted 1:1000</td>
<td>v1.0</td>
<td>6</td>
<td>27.1</td>
<td>12</td>
<td>44%</td>
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<tr>
<td></td>
<td>v2.0</td>
<td>8</td>
<td>23</td>
<td>85%</td>
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Table 2. Quantification of plasma viremia with gSCA and iSCA in 10 patients with untreated HIV-1 infection

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Plasma HIV-1 RNA (Log_{10} HIV-1 RNA copies per milliliter of plasma)</th>
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<tbody>
<tr>
<td></td>
<td>gSCA</td>
</tr>
<tr>
<td>1</td>
<td>-0.52^b,c</td>
</tr>
<tr>
<td>2</td>
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<tr>
<td>3</td>
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<td>9</td>
<td>4.6</td>
</tr>
<tr>
<td>10</td>
<td>2.5</td>
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Median (IQR) 3.4 (1.9 to 4.6) 4.2 (3.3 to 4.5) 4.2 (3.4 to 4.7)

^aRoche COBAS Ampliprep/COBAS TaqMan version 2.0  
^bSample below the limit of detection of the assay  
^cSample with <10-fold recovery compared to Roche TaqMan
Table 3. Summary of iSCA versus gSCA in 25 patients on ART

<table>
<thead>
<tr>
<th></th>
<th>gSCA</th>
<th>iSCA</th>
</tr>
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<tbody>
<tr>
<td>Fraction of samples with</td>
<td>13 of 25</td>
<td>17 of 25</td>
</tr>
<tr>
<td>HIV-1 RNA Detected</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fraction of negatives</td>
<td>8 of 12</td>
<td>4 of 8</td>
</tr>
<tr>
<td>detected by alternative assay</td>
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<td></td>
</tr>
<tr>
<td>Median HIV-1 RNA copies/mL</td>
<td>1.3 (0.30 to</td>
<td>2.0 (0.35 to</td>
</tr>
<tr>
<td>(IQR)</td>
<td>3.3)</td>
<td>4.9)</td>
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<tr>
<td>Fraction with HIV-1 RNA</td>
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<td>10 of 25</td>
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<td>levels 3-fold higher than</td>
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<td>alternative assay</td>
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Table 4. Mega-iSCA can be used to increase assay sensitivity

<table>
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<tr>
<th>Patient ID</th>
<th>Plasma volume tested</th>
<th>Copies per milliliter</th>
<th>iSCA</th>
<th>Plasma volume Tested</th>
<th>Total Copies Detected</th>
<th>Copies per milliliter</th>
<th>Mega-iSCA</th>
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