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The development of a standardized quality assessment material to support Xpert HIV-1 viral load testing for ART monitoring in South Africa

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Abstract

The tiered laboratory framework for HIV viral load monitoring accommodates a range of HIV viral load testing platforms, with quality assessment critical to ensure quality patient testing. HIV plasma viral load testing is challenged by the instability of viral RNA. An approach using an RNA stabilizing buffer is described for the Xpert HIV-1 Viral Load (Cepheid) assay and was tested in remote laboratories in South Africa.

EDTA-plasma panels with known HIV viral titres was prepared in PrimeStore molecular transport medium for per-module verification and per-instrument external quality assessment. The panels were transported at ambient temperature to 13 testing laboratories during 2017-2018, tested according to standard procedures and uploaded to a web portal for analysis

A total of 275 quality assessment specimens (57 verification panels and two EQA cycles) were tested. All participants passed verification (n=171 specimens) with an overall concordance correlation (ρ_c) of 0.997 (95% confidence interval [CI]:0.996,0.998) and a mean log bias of -0.019log cp/mL (95% CI:-0.044,0.063). The overall EQA ρ_c (n=104 specimens) was 0.999 (95% CI:0.998,0.999), with a mean log bias of 0.03 log cp/mL 95% (CI:0.02,0.05).

The panels are suitable for use in quality monitoring of Xpert HIV-1 VL and are applicable to laboratories in remote settings.

Keywords: HIV viral load, external quality assessment, verification, quality, thermostable, PrimeStore MTM

1. Introduction

Several countries striving to attain their 2020 UNAIDS 90%/90%/90% targets for global HIV healthcare (1-4), struggle with the third 90% (virological suppression). To address this, fast-track targets were designed (5), aiming to increase the number of people living with HIV (PLWH) accessing treatment and achieving virological suppression. Current global estimates show that 25.4 million people, approximately 67% of PLWH, were accessing antiretroviral therapy (ART) by end-2019 (6), and monitoring needs are likely to increase over the next decade as more people access ART. A total of 5,231,809 (70%) patients currently access ART in South Africa alone (7), with the number expected to increase, as the remaining PLWH are reached. The recommended test for monitoring ART response is HIV viral load

(VL) quantification (8), which has historically been performed at centralised laboratories owing to the number of specimens requiring processing, the logistical needs of the available technologies, and the lack of accurate and cost-effective near patient VL technologies. South Africa has addressed the VL scale-up testing needs through a highly centralised model within the National Health Laboratory Service (NHLS), which is responsible for laboratory testing of ~80% of the population. The capacity of the 16 high throughput, centralised HIV VL laboratories has been further augmented through automation and instruments with increased throughput (9-17), most recently the cobas® 8800 (Roche Molecular, Pleasanton, CA, USA) and Alinity (Abbott Molecular, Des Plaines, IL, USA) systems.

Nonetheless, there are a number of PLWH who live in remote areas and whom are unable to access the centralised facilities, as highlighted during the current COVID-19 pandemic, either because no collection facilities exist within travelling distance or because specimen transport to the testing laboratories is limited by the stability of HIV RNA plasma (18, 19). While there are studies showing long-term stability of HIV in whole blood (20-22), the manufacturers of the VL technologies recommend testing within 24 hours, with separation of plasma within six hours and specimen refrigeration (23, 24), primarily to maintain the quality of low VL specimens and to overcome the extreme temperatures (>30°C) in many high HIV prevalence regions. The use of plasma preparation tubes (PPT; Becton Dickinson, USA) was introduced (22, 25, 26) to increase the specimen transport window to at least 24 hours (27, 28), although specimens should still be separated within six hours of collection and prior to transport (22). Alternative options to plasma based testing include the use of dried blood spots (DBS) and several countries have shown that this is a feasible option for remote collection and central testing (29-35). The DBS matrix, nonetheless, is challenged by inaccuracies at the clinically relevant range (1000 copies per millilitre (cp/mL)) as the VL at this threshold increases due to the contribution of cell associated RNA (36). Subsequently, while this remains the recommended level for virological failure (37), there is contention regarding the use of DBS at VL below 5000 cp/mL (38, 39). A decentralised model, utilising mobile or remote clinics, may address the needs of PLWH in remote areas through a tiered laboratory network (18, 19, 40), similar to that originally used for CD4 scale-up (41). As such, the NHLS National Priority Programme (NPP), in collaboration with the South African Department of Health, and through the Global Fund to Fight HIV, Tuberculosis and Malaria (Global Fund; Geneva, Switzerland), performed a pilot evaluation of the Xpert HIV-1 VL (Cepheid, Sunnyvale, CA, USA), in remote district laboratories. The Xpert HIV-1 VL assay was previously evaluated in collaboration with the NPP (3) and received World Health Organisation pre-qualification status in 2017 (42). In addition to being one of the few commercially available POCT HIV VL assays ready for implementation at the time of the study, this platform was selected due to the existing GeneXpert footprint in South Africa, through the Xpert MTB/RIF programme which comprises 207 tuberculosis testing sites, and the goal of integrated diagnosis and monitoring through multipurpose testing platforms.

As part of the HIV VL testing mandate, technologies selected for the NHLS laboratories must be verified (“fit for purpose”) upon installation and prior to testing clinical specimens, regardless of placement within the testing framework. Verification material is frequently sought by the testing laboratory (laboratory networks) from residual patient’s specimens, but it is often difficult to obtain sufficient volumes for paired (duplicate/split) testing, and is not always possible for remote testing sites. Participation in EQA programs, such as the global

Virology Quality Assurance program (VQA, supplied by the Department of AIDS (National Institute of Health, Atlanta, GA, USA)) or the National External Quality Assessment Service (NEQAS, United Kingdom) HIV-1 RNA quantitation programme, does provide assurance to an accredited laboratory for pathology services, but does not address pre-testing verification. Furthermore, these panels require expensive shipment, are only available at times of the annual panel testing cycles, and comprise limited numbers of specimens ($n \sim 5$). In addition, the World Health Organisation has published considerations for POCT, including the need for instrument verification as 'fit for purpose' and external quality assessment at least annually (43). Dried tube specimens (DTS) (44-46) were not considered, as it was desirable to minimise onsite processing, mimic plasma specimens as far as possible and ensure sufficient specimen volume for use with the Xpert HIV-1 VL assay (1.1mL).

In addition to the programmes described above, the South African Viral Load Quality Assessment (SAVQA) panel (47) was developed to address the need for scaled HIV VL services in centralised HIV VL laboratories. This panel provides an accessible option for the verification of newly installed HIV VL testing platforms, initially the RealTime HIV-1 (Abbott) and cobas® AmpliPrep/cobas® TaqMan® (CAP/CTM; Roche) assays, prior to testing clinical specimens, and is also used for the rapid evaluation of new HIV VL assays (3, 4, 48, 49). The SAVQA panel is a 42-specimen plasma panel, which is stored and shipped frozen, and only defrosted immediately prior to testing (47). The panel comprises 17 negative specimens and five repeats of five positive specimens with VL ranging from 2.7 log cp/mL to 5.0 log cp/mL, and was designed to measure accuracy, precision, carryover and limit of the blank (47). The SAVQA panel was readily available, but was not suitable in its existing format. The panel required adaption to avoid the need for cold-chain shipping and storage, with the remote testing sites having no refrigeration facilities, and also to include a smaller number of specimens, minimising the cost and time constraints since the GeneXpert is a modular, cartridge-based system designed for random access, single specimen testing. We hence designed a miniaturised, thermostable version of the SAVQA panel using a commercially available matrix, PrimeStore® Molecular Transport Medium (MTM; Longhorn Vaccines and Diagnostics LLM, Bethesda, MD, USA), to allow ambient temperature shipping and storage. This medium achieved US FDA approval in 2018 (50), and has been evaluated with a variety of mycobacterial (51-53) and viral (54-56) specimens, including HIV (57). In addition to use of MTM-stored specimens with PrimeMix® (52, 55, 56), MTM has been shown to be compatible with the Xpert MTB/RIF assay (Cepheid, Sunnyvale, CA, USA) (51) and the m2000 RealTime HIV-1 (57). Verification panels were developed alongside a web-based result reporting tool, which was based on the web portal (www.tbgxmonitor.com) previously developed for Xpert MTB/RIF quality monitoring (58). Following the successful verification rollout, an external quality assessment (EQA) panel was requested, and designed to measure pre- and post-processing analytics at these pilot laboratories. This manuscript aims to provide a detailed description of these pilot quality panels as an option for POCT HIV VL sites, using clinically relevant panel specimens, which can be prepared centrally and sent to remote sites. These panels were specifically designed to meet the needs of remote testing laboratories using the GeneXpert HIV-1 VL assay, notably limited cold-chain shipping and cold-storage facilities on site, low throughput testing platforms, the need for *ad hoc* verification products and, frequently, lower-skilled laboratory staff. The use of QA materials, particularly when evaluated between laboratories, ensures that

instruments are fit-for-purpose and that onsite processing is robust, thus ensuring best possible patient result quality within a tiered laboratory framework.

2. Materials and Methods

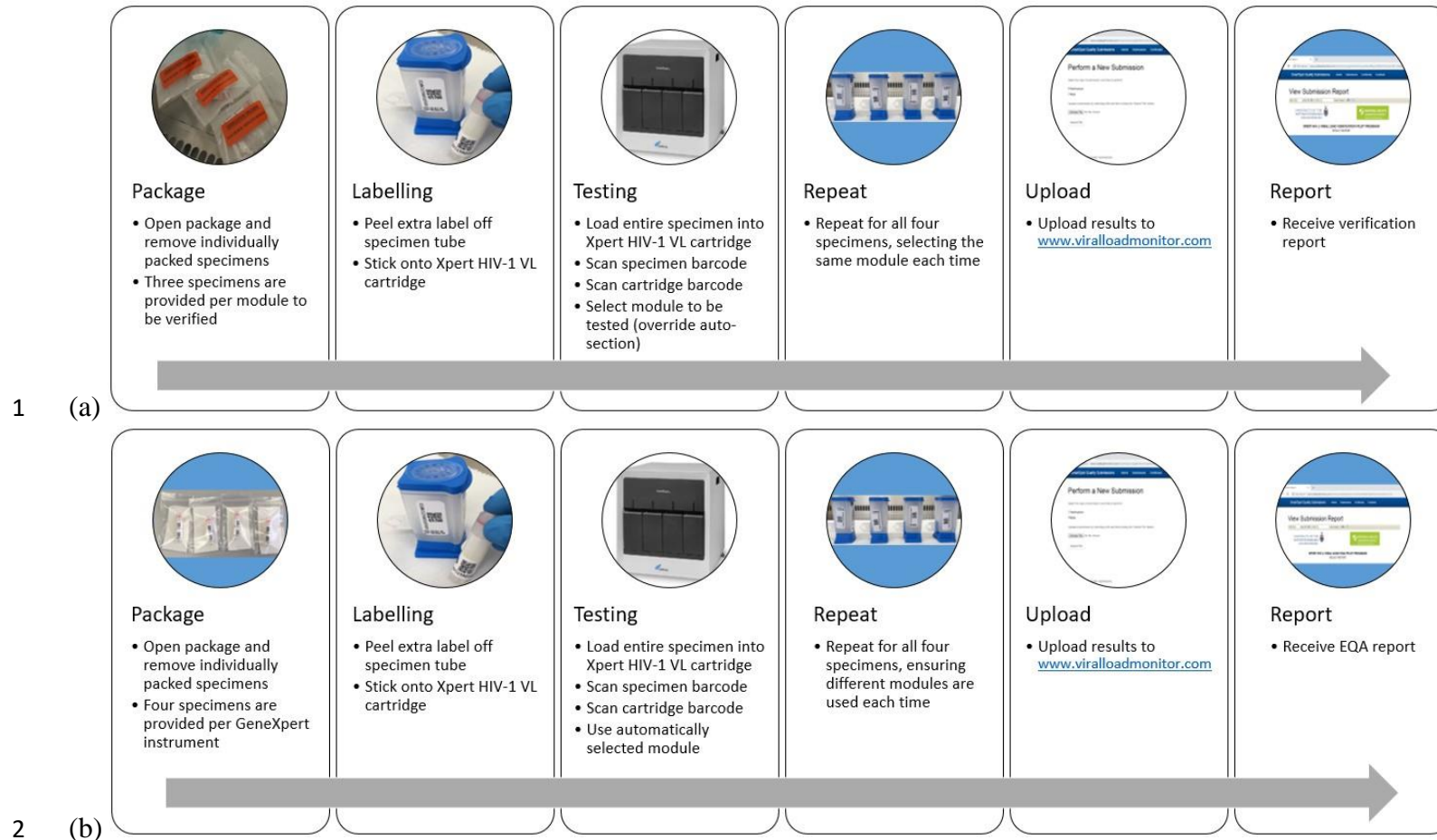
2.1 Panel material preparation

SAVQA plasma was removed from storage (-80°C) and defrosted at ambient temperature, followed by brief centrifugation (3000rpm, 1 minute). HIV-negative specimens (1.3mL) were not mixed with MTM to provide a clinically relevant specimen, overcoming the decreased viscosity/fat content of the MTM. The negative specimen is important to ensure that no cross-contamination occurs in either the reference laboratory or the testing laboratory during specimen preparation and loading. HIV-positive plasma specimens (300µl) with known VL were added to 1ml MTM (Longhorn Vaccines and Diagnostics LLC, Bethesda, MD, USA), giving a dilution factor of 4.3. To minimise the risk of leakage, each specimen was packaged individually in a sealed plastic bag with an absorbent pad and the complete panel was then placed into a second sealable bag. Specimens were shipped at ambient temperature using the routine NHLS specimen transport system.

Two panel formats were designed: (i) a verification panel (Figure 1a) and (ii) an EQA panel, (Figure 1b). The **verification panel** is used to ensure that an instrument is functioning correctly upon installation, instrument (module) replacement or instrument movement, and can also be used for staff training. The verification panel consists of three specimens per module tested: two specimens of known HIV VL stabilised in MTM buffer and one HIV-negative specimen (plasma only). The target ranges for the HIV-positive specimens were 2.7 log cp/mL (low), 3.0 log cp/mL (low), 4.7 log cp/mL (high) and 5.0 log cp/mL (high), with all sites receiving one low VL and one high VL specimen in addition to an HIV-negative specimen, as per testing organisation requirements. The **EQA panel** is necessary for ongoing monitoring of instruments and testing sites. Four specimens are provided per instrument tested, with an instrument being defined as “up to four” GeneXpert systems attached to one computer. The panel includes three specimens of a known HIV VL stabilised in MTM buffer with a target range of 3.0 log cp/mL, 3.7 log cp/mL and 4.7 log cp/mL and one HIV-negative plasma specimen. On preparation of either panel format, one specimen in each range was tested using the reference laboratory instrument (reference specimen; day 0).

2.2 Xpert HIV-1 VL quality panel testing

Both the verification and EQA specimens were processed according to the Xpert HIV-1 VL manufacturer’s instructions (Cepheid, Sunnyvale, CA, USA), using the liquid panel in place of clinical plasma. Briefly, the Xpert HIV-1 VL cartridge was opened and the entire specimen volume (1.3mL) was transferred into the Xpert HIV-1 VL cartridge using a precision pipette or 1mL Pasteur pipette (supplied by Cepheid as part of the kit). The specimen barcode and cartridge number were scanned and the specimen was tested using the Xpert HIV-1 Viral Load assay definition file. The original SOP did not include centrifugation instructions, but this was amended after the first verification panel was analysed to ensure that every specimen was briefly centrifuged (3000rpm, 1 minute) prior to processing.



3 **Figure 1: Processing of Verification and EQA panels**

4 (a) Verification panel: same module must be used for each set of specimens. Verification panels are labelled with orange labels to remind users
 5 of this. (b) EQA panel: different modules must be used for each specimen.

2.3 Result return and performance scoring

A web portal (www.viralloadmonitor.com), based on the original TBGxMonitor website (58) for upload of both verification and EQA results and report generation, was created in collaboration with SmartSpot Quality (Johannesburg, Gauteng, South Africa). Post specimen panel testing, users were required to upload the comma-separated values (CSV) run files (automatically produced by the GeneXpert software) of the Xpert HIV-1 VL panel specimens using a USB device. Results were converted using the dilution factor and this was applied within the website logic as part of the scoring algorithm. The acceptable difference between the test specimen and the Xpert HIV-1 VL reference specimen (described above) was defined as <1.0 log cp/mL difference. Retrospective analysis at <0.5 log cp/mL difference and <0.3 log cp/mL difference, in line with standard VL variation (59), was also performed. Further, the Xpert HIV-1 VL reference VL was compared to the pooled mean VL achieved by the 13 testing sites, ensuring that the reference laboratory instrument was performing acceptably and that the reference result was suitable for use as the standard. Each specimen tested received a score out of two: correct result (2/2); error, invalid, >1.0 log cp/ml quantifiable result bias (1/2); incorrect result (e.g. HIV positive reported as HIV negative: 0/2). Each panel was then scored out of six for verification and out of eight for EQA. Scoring logic is detailed in Table 1. The overall panel performance across all sites was measured by the mean, median, range and standard deviation (SD) of the quantifiable viral loads, which were calculated using Microsoft® Excel® 2016 (Microsoft Corporation, Redmond, WA, USA). Regression, the concordance correlation coefficient (P_c) (60, 61), including a Pearson correlation coefficient (p ; measure of precision) and a bias correction factor (C_b ; measure of accuracy), and Bland-Altman (62, 63) analyses were performed and graphically represented using MedCalc Statistical Software version 18.11 (MedCalc Software bvba, Ostend, Belgium; <http://www.medcalc.org>; 2018).

Table 1: Summary of scoring logic

Specimen Score	Results	Outcome
2/2	Correct result	Pass
1/2	Error, Invalid, No result >1.0 log cp/ml quantifiable result bias	Acceptable
0/2	Incorrect result (e.g. HIV positive reported as HIV negative)	Concern
Verification Score	Percentage Performance	Outcome
6/6	100%	Pass
5/6	83.3%	Acceptable
$\leq 4/6$	66.7%	Unacceptable
EQA Score	Percentage Performance	Outcome
8/8	100%	Pass
7/8	87.5%	Acceptable
6/8	75.0%	Concern
$\leq 5/8$	62.5%	Unacceptable

Each specimen generates a score out of two. Verification of a module generates a score out of six (three specimens per module) and EQA of an instrument generates a score out of eight (four specimens per instrument, run over different modules). If an unacceptable score is obtained, the site is required to conduct a root cause analysis and corrective action, and to test a second verification or EQA panel. Site trainers or monitors may provide further interventions (e.g. staff training, instrument calibration).

2.4 Verification and EQA pilot field evaluation

The pilot evaluation was nested within a field trial of near-patient VL testing, overseen by the NHLS NPP (Johannesburg, South Africa). Briefly, thirteen district laboratory facilities were selected and provided with a GeneXpert IV (Cepheid, Sunnyvale, CA, USA). The laboratories were located in remote areas across six provinces (Eastern Cape: n=2, Northern Cape: n=4, Western Cape: n=3; Free State: n=1, Limpopo: n=2; North West Province: n=1). Technicians were recruited and received training on the GeneXpert platform and the Xpert HIV-1 VL assay. The verification and EQA material was designed to meet requirements of the NPP to ensure that the instruments were fit-for-purpose and that specimen processing was being correctly performed.

Verification panels (n=4 per site) were provided to all sites in September 2017, following instrument installation and prior to patient testing. Further verification panels (n=5) were provided on an ad hoc basis as modules were replaced. EQA panels (n=1 per site) were provided to the sites in June and November 2018. For the pilot evaluation, the automatically generated reports were manually checked prior to release, but the website has the capacity automatic report release to the sites.

2.5 Stability testing

Prior to initial supply to sites, verification specimens (2.7 log cp/mL; 5.0 log cp/mL) were prepared and tested in duplicate at days 7, 14, 21 and 28 (as per process described above) to determine stability compared to the day 0 reference result. Extra EQA panels (3.0 log cp/mL, 3.7 log cp/mL and 4.7 log cp/mL) were prepared at the same time as those sent to the sites and tested at days 24, 43, 84 and 150 post manufacture to determine longer term stability. All specimens were stored at ambient temperature in sealed plastic bags with desiccant.

3. Results

3.1 Verification panel performance

All sites tested and uploaded results to the website within three days of panel receipt. Result scores and outcomes are summarised in Table 2 and Figure 2, with detailed information provided in supplementary table S1. Quantifiable VL results were within acceptable limits for verification (<1.0 log cp/mL difference from the reference VL) and all reference results were within 0.3 log cp/mL of the pooled mean VL of the specimens tested, although it was noted that the VL bias was high in the 5.0 log cp/mL reference specimen (0.22 log cp/mL). In addition, the sites' verification VL results were compared to the mean VL (data not shown) and this was comparable to analysis using the reference VL values. The p_c across all sites (n=151 specimens) was 0.997 (95% confidence interval (CI): 0.995, 0.998), with a p of 0.997 and a C_b of 0.999. The mean log bias was -0.02 log cp/mL (95% CI: -0.046, 0.006), with a coefficient of determination (R^2) value of 0.9940.

The error rate (20/171; 11.7%) for the verification panels was higher than expected, and was primarily a result of processing errors (55% of errors). Seven errors (35%) were linked to the internal probe failures, two to syringe pressure (10%) and eleven relating to input volume (errors 2096 (35%) and 2097 (20%)). The majority of errors reported (13/20; 65%) occurred in the clinically relevant negative specimen, indicating laboratory processing errors. It was determined, on discussion with the programme manager, that the specimens were not being centrifuged prior to testing and that incorrect pipetting procedures may have contributed. Changes were implemented to the standard operating procedure, to centrifuge all specimens prior to use (as would be required for clinical specimens), and staff retraining was performed if necessary. Once these changes were implemented, the error rate (over ad hoc verification and EQA) decreased to 1.7% (2/119 further tests), indicating that correct operating procedures were being observed.

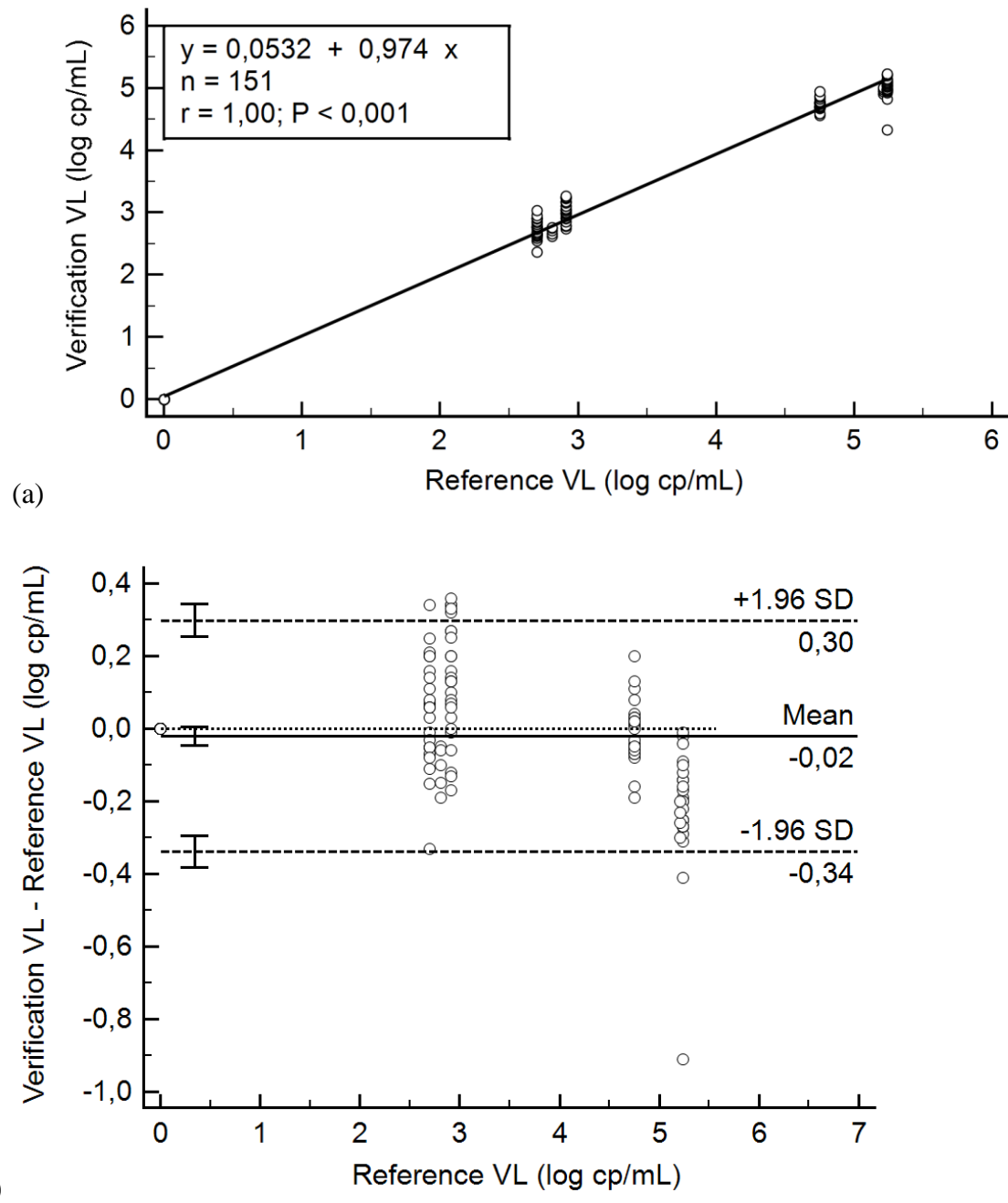


Figure 2: Verification panel VL variation (log cp/mL) across different testing sites (n=13)

(a) Regression analysis for all verification panels tested between September 2017 and November 2018. (b) Bland-Altman agreement of the viral load results, compared to the reference result obtained at panel preparation. One outlier specimen (4.33 log cp/mL; -0.91 log cp/mL difference from reference VL) was noted in the 5.0 log cp/mL category, but was within the acceptable range for the pilot panels (<1.0 log cp/mL).

Table 2: Site Verification Summary: September 2017-November 2018 (compared to reference VL)

Panel	Expected Viral Load (log cp/mL)	Reference Viral Load (log cp/mL)	Tested (n)	Result obtained (n (%))	Viral Load Bias (mean (median) range) (log cp/mL)	Standard Deviation of Mean Bias (log cp/mL)	Error (n)	Invalid (n)	Reference vs Mean (log cp/mL)
1	Negative	Negative	52	39 (75.0)	0	0	12	1	0
2 ^c	Negative	Negative	5	5 (100)	0	0	0	0	0
1	2.70	2.70	26	23 (88.5)	0.04 (0.06) -0.33, 0.34	0.15	2	1	-0.04
2 ^c	2.70	2.81	5	5 (100)	-0.11 (-0.10) -0.19, -0.06	0.06	0	0	0.11
Overall (log 2.70)			31	28 (90.3)	0.02 (-0.02) -0.33, 0.34^b	0.15	2	1	-
1	3.00	2.91	26	25 (96.2)	0.13 (0.13) (-0.17, 0.36)	0.16	1	0	-0.14
1	4.70	4.75	26	25 (96.2)	-0.01 (0.00) -0.19, 0.20	0.09	1	0	0.01
1	5.00	5.24	26	25 (96.2)	-0.22 (-0.20) -0.91 ^a , -0.01	0.18	1	0	0.22 ^a
2 ^c	5.00	5.21	5	4 (80.0)	-0.25 (-0.25) -0.30; -0.20	0.04	0	1	0.25
Overall (log 5.00)			31	29 (93.6)	-0.23 (-0.22) -0.91; -0.01	0.17	1	1	-
Overall (57 verification panels)			171	151/171 (88.3) Quantified: 107/114 (93.9)	-0.02 (0.00) (-0.91, 0.36)	0.16	17 9.9%	3 1.8%	0.07

^a increased variability owing to one outlier specimen (4.33 log cp/mL). If this specimen is excluded, the mean bias increases to -0.19 log cp/mL with a range of -0.41 to -0.01, and the difference between the reference and the pooled mean decreases to 0.19 log cp/mL.

^b variation around the median >0.30 when two panels are combined, but remains <0.03 log cp/mL in the individual panels.

^c Verification panel 2 numbers are low (n=5), so values lack robustness, but are similar to the larger panel 1.

3.2 Pilot EQA performance

Two cycles of EQA (E18V1, E18V2) were shipped to 13 sites (18 June 2018, 12 November 2018) and results were uploaded within seven days (mean: 4.1 days). All sites showed acceptable performance across both EQA panels; the programme performance is summarised in Table 3 and Figure 3, and complete site results are detailed in supplementary table S2. Viral loads were within acceptable limits for EQA (<1.0 log cp/mL bias), and all negative specimens were reported as not detectable (no carryover). The p_c for the EQA pilot panels (two EQA panels, $n=102/104$ specimens) across all sites was 0.9985 (95% CI: 0.9978, 0.9990), with a p of 0.9987 and a C_b of 0.9998. The mean log bias was 0.03 (95% CI: 0.02, 0.05). The error rate was 1.9% (2/104 tests) and was caused by volume loading (user) errors.

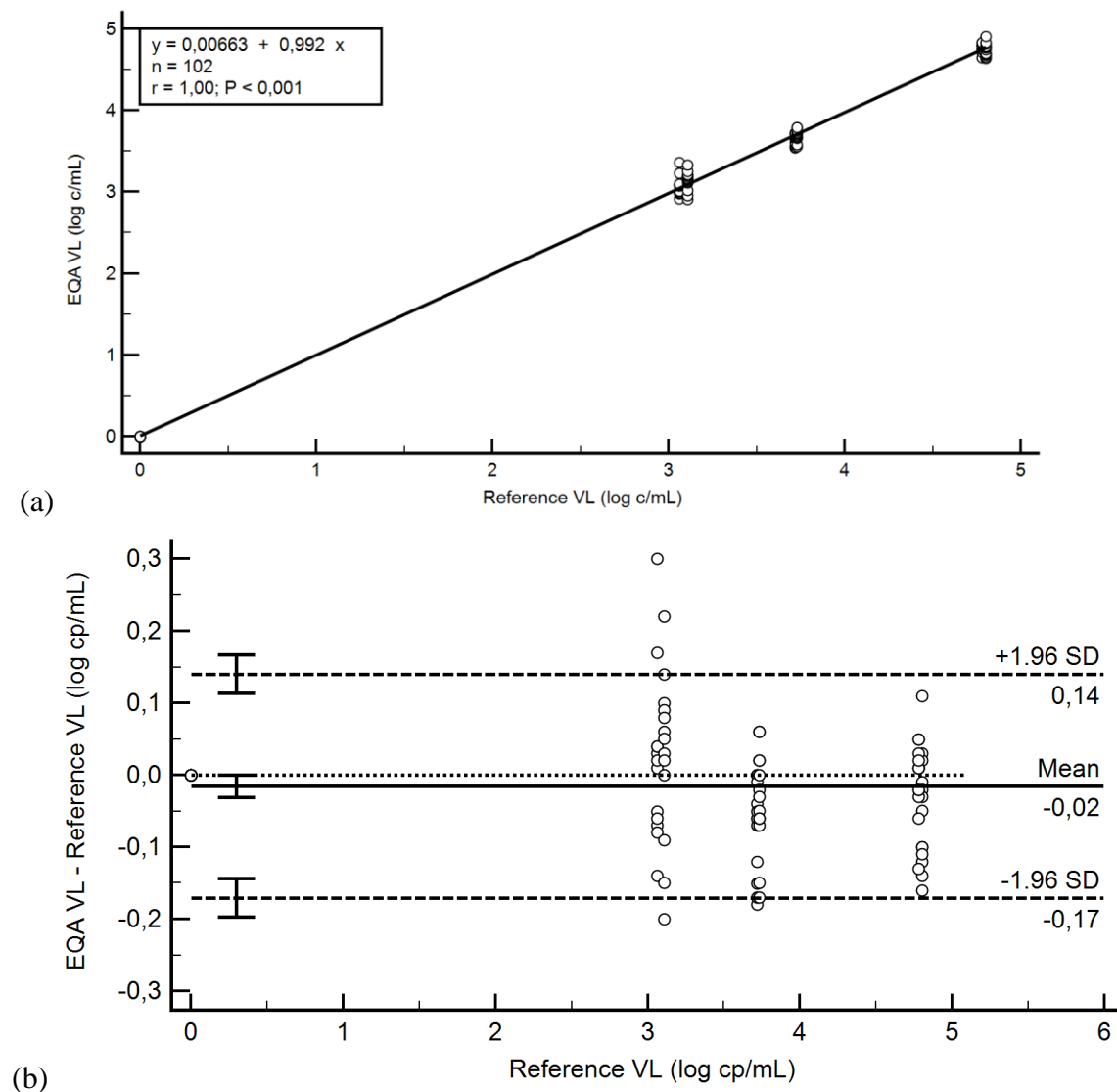


Figure 3: EQA Panel VL variation (log cp/mL) across different testing sites ($n=13$) and EQA panels ($n=2$)

(a) Regression analysis for EQA Panels 1 and 2 ($n=102/104$ specimens). (b) Bland-Altman agreement of the viral load results ($n=102/104$ specimens), compared to the reference result obtained at panel preparation.

Table 3: Site EQA Summary: September 2017-November 2018

Panel	Expected Viral Load (log cp/mL)	Reference Viral Load (log cp/mL)	Tested (n)	Result obtained (n (%))	Viral Load Bias (mean (median) range) (log cp/mL)	Standard Deviation of Mean Bias (log cp/mL)	Error (n)	Reference vs Mean (log cp/mL)
1	Negative	Negative	13	13 (100)	0	0	-	0
2	Negative	Negative	13	13 (100)	0	0	-	0
1	3.00	3.06	13	12 (92.3)	0.02 (0.02) -0.14, 0.30	0.11	1	-0.02
2	3.00	3.11	13	13 (100)	0.02 (0.04) -0.20, 0.22	0.12	-	0.05
Overall (log 3.00)		3.09	26	25 (96.2)	0.02 (0.02) -0.20, 0.30	0.11	1	-
1	3.70	3.72	13	13 (100)	-0.06 (-0.06) -0.18, 0.05	0.07	-	0.06
2	3.70	3.73	13	13 (100)	-0.04 (-0.04) -0.17, 0.06	0.07	-	0.01
Overall (log 3.70)		3.73	26	26 (100)	-0.05 (-0.05) -0.18, 0.06	0.07	-	-
1	4.70	4.80	13	13 (100)	-0.05 (-0.04) -0.16; 0.11	0.08	-	0.05
2	4.70	4.78	13	12 (92.3)	-0.01 (0.01) -0.13, 0.05	0.05	1	-0.02
Overall (log 4.70)		4.79	26	25 (96.2)	-0.03 (-0.02) -0.16; 0.11	0.07	1	-
Overall (26 EQA panels panels)			104	102/104 (98.1)	-0.02 (-0.02) -0.20, 0.30	0.09	2 1.9%	-

3.3 Retrospective result analysis

Retrospective analysis of the verification and EQA results was performed after the pilot evaluation, in order to accommodate acceptable VL biases (59). Amongst 107 quantifiable verification results, ten (9.3%) showed a bias of >0.3 log cp/mL (range: 0.36, -0.91), with only one outlier specimen (4.33 log cp/mL) displaying a bias >0.5 log cp/mL. This specimen has a bias of -0.91 log cp/mL compared to the reference VL and -0.69 log cp/mL compared to the pooled mean VL. This specimen was part of the 5.0 log cp/mL group, where the reference VL (5.24 log cp/mL) was notably higher than the pooled mean VL (5.02 log cp/mL). A second outlier (4.83 log cp/mL) in this group had a VL bias of -0.41 log cp/mL compared to the reference VL, with an acceptable bias of -0.19 log cp/mL compared to the pooled mean VL. Only three specimens (2.8%) had a bias of >0.3 log cp/mL compared to the pooled mean VL. All quantifiable EQA VL (n=76) results showed a bias of <0.3 log cp/mL compared to the reference VL.

3.4 Specimen stability

Stability of the specimens stored in MTM was evaluated prior to panel design and supply, with stability shown up to 28 days (Figure 4a), with specimen stability acceptable up to four weeks. Testing of EQA panels in the reference laboratory between weeks 4 and 20 (Figure 4b), showed stability of all specimens at week 6 (day 43) and extended stability of the higher VL range (4.7 log cp/mL) specimens until week 12 (day 84). However, by week 12, a decrease of ~ 0.5 log cp/mL was noted in the lower (3.0 log cp/mL) VL range. Errors were noted in the 3.0 log cp/mL specimen at day 24 (error 2126; module reset) and in the 3.7 log cp/mL specimen at day 84 (invalid, error 5016; probe check error). These relate to the instrument and the cartridge, rather than the specimen. Retesting was not possible due to limited specimen availability. By Day 150, all VL exceeded >0.5 log cp/mL difference from baseline (day 0), with both the 3.7 log cp/mL and 4.7 log cp/mL specimens showing a VL decrease of >1.0 log cp/mL. Bland-Altman analysis of the reportable VL results (n=14/16) over the weeks, including day 84, when a VL decrease was noted, but excluding day 150, when VL were no longer relevant, gave a mean log bias of -0.06 log cp/mL with a lower limit of -0.34 log cp/mL (95% CI: -0.89, -0.21) and an upper limit of 0.23 log cp/mL (95% CI: 0.10, 0.77). Including day 150 (n=18/20) gave a mean log difference of -0.20 log cp/mL with a lower limit of -0.97 log cp/mL (95% CI: -2.11, -0.62) and an upper limit of 0.58 log c/mL (95% CI: 0.23, 1.72), beyond acceptable limits for supply to sites.

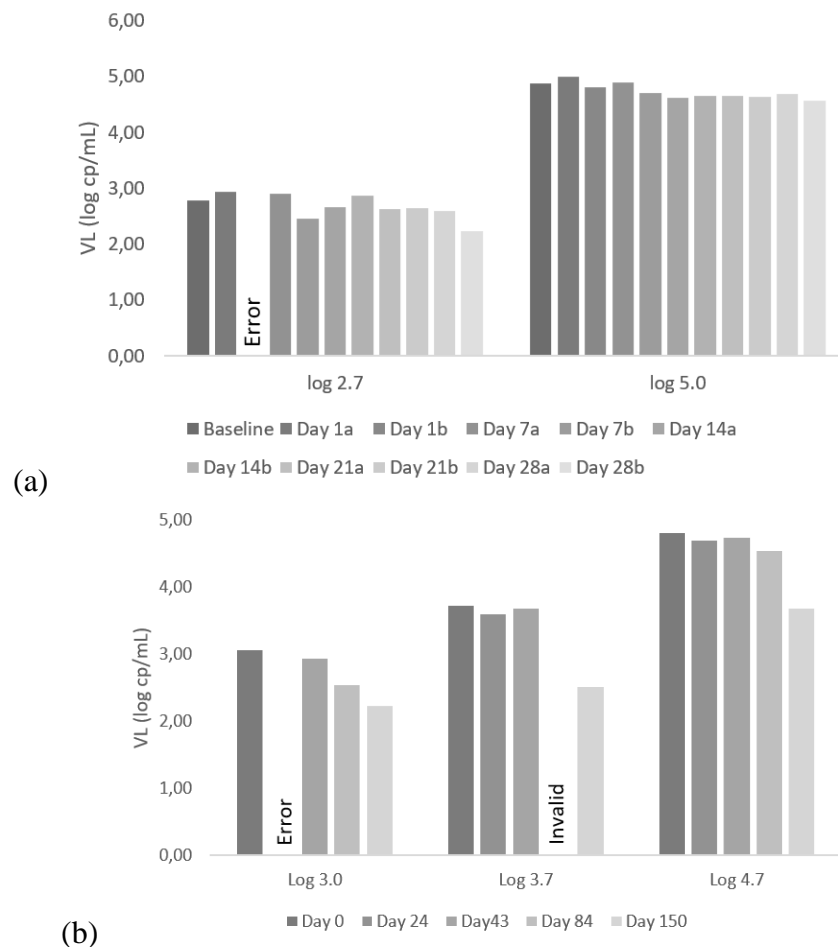


Figure 4: Stability of EQA Pilot Panel Baseline to Day 150

(a) Bar chart showing VL from Day 0 to Day 28, with specimens tested in duplicate. There is little VL variability. (b) Bar chart showing VL from Day 0 to Day 150. There is a decrease in VL between Day 84 and Day 150. The VL remains within 0.2 log cp/mL of the expected VL for the log 3.7 and log 4.7 specimens until Day 84. There is a decrease at Day 84 for the log 3 specimen, but it remains within 0.5 log cp/mL of the expected VL. By Day 150, all VL exceed >0.5 log cp/mL difference from day 0, with both the log 3.7 and log 4.7 specimens showing a VL decrease of greater than 1.0 log cp/mL.

4. Discussion

Laboratory quality monitoring is vital to ensure ongoing patient result testing accuracy. Instruments must be evaluated prior to implementation, verified before use in the field and monitored on an ongoing basis. Similarly, staff competency should be evaluated through training, observation and participation in quality programmes. Evaluation can be performed on existing specimens (e.g. frozen plasma), prospective specimens (against a reference instrument currently in use) or on well-described quality panels (e.g. NEQAS, SAVQA). EQA, through supply of standardised specimens for testing and through continuous quality monitoring (CQM; e.g. analysis of central data repositories), enables programme managers to identify potential instrument or staff deficiencies for correction. Participation in EQA programmes has been shown to improve participant performance (45). CQM of assays and instruments is becoming standard practice for many connected diagnostics. Operational

dashboards, such as C360 (Cepheid), provide assay and instrument quality information on errors, utility, and various result parameters on a module/instrument/laboratory and location basis, and can be utilized for daily and monthly monitoring to identify quality issues, without waiting for EQA panel cycles (64). CQM, through the C360 platform, was successfully applied during the near-patient testing pilot into which this evaluation was nested, but is beyond the scope of this manuscript. EQA is complimentary to CQM, ensuring ongoing pre- and post-analytical performance monitoring, which is particularly important where staff turnover is high.

The Xpert HIV-1 VL assay was previously evaluated, using both the SAVQA panel and clinical specimens (3), and has since been extensively evaluated in the field (65, 66), meaning that the assay did not require further evaluation prior to implementation. However, before the implementation pilot could commence, verification of the modules was required and was complicated by the remote placement of the instruments, as residual plasma specimens were not readily available. Alternative options for instrument verification were thus needed. This manuscript describes the design and pilot evaluation of quality panels used for POCT HIV VL. The panels were designed to meet specific requirements: (i) specimen processing needed to be as similar as possible to actual specimens; (ii) thermostable transport and storage; (iii) reproducible VL results, such that processing or instrument issues could be detected during verification and ongoing EQA, and (iv) safe during transport. While initially designed for module verification, the panels were easily adapted for ongoing EQA. This program is based on similar principles to the Xpert MTB/RIF program (67, 68), which has been used successfully throughout the NHLS to monitor 207 Xpert MTB/RIF testing sites, as well as internationally (28 countries), and is expected to provide similar rigorous quality monitoring to Xpert HIV-1 VL sites.

It is notable that the panels were supplied in a liquid format and that no processing was required beyond direct addition of specimen into the Xpert HIV-1 cartridge, mimicking routine patient specimen testing. This is in contrast to dried tube specimens (DTS), which have been used throughout sub-Saharan Africa for EQA (44-46). DTS were not selected for this programme as the NPP preferred to minimise specimen processing variability during specimen reconstitution by using a liquid panel, although DTS meet all other requirements described. Furthermore, similarly to the original SAVQA panel, the verification programme was designed for rapid deployment using local resources, decreasing reliance on scheduled schemes (47). This design can be adapted by reference laboratories aiming to supply similar panels, adapted to individual country needs, to remote sites within a tiered laboratory system to ensure continued quality POCT HIV VL testing, although this would need to be evaluated on an individual basis. It is often difficult for smaller programmes to realise cost savings through in-house QC offerings, as the costs of the MTM buffer, plasma (if purchased), staff time required to manufacture the panels and to collate the results, post-manufacture quality testing and shipping must be considered. Similarly, if this quality material was provided commercially, the cost and feasibility of scaled manufacture should be investigated for implementation uptake. Furthermore, this product was designed and optimised specifically for the GeneXpert platform placed at POC, and its compatibility with alternative HIV VL assays must be determined prior to use on alternative platforms, particularly as it has been observed that the MTM buffer can interact negatively with certain assays (personal communication, Dean Sher, SmartSpot Quality Inc.). Commercially available panels,

particularly for EQA, should not be disregarded, particularly for smaller programmes where globally standardised specimens may provide more rigorous quality measures. A further consideration for using commercial EQA panels is to free up the time of the programme managers from producing panels and evaluating results, so as to use this time to assist the laboratories which the EQA identifies as needing help, to identify root-causes and implement corrective actions. Ultimately, whichever option is selected, the goal is to ensure quality testing to positively impact on patient care and management.

Shipping of liquid specimens is potentially problematic, given the risk of leakage. Panels were well packaged and no leakage of the specimen from the tube into the protective packing was observed. However, the extra packaging, as described above, is recommended for similar panels going forward to minimise risk to transport personnel and to meet IATA requirements (69). The infectivity of HIV when stored in MTM was not tested in this pilot, but existing studies have shown that pathogens are fully inactivated on addition to the buffer (52, 54, 70), while RNA integrity is simultaneously preserved (52, 55, 56), including HIV-1 RNA (57). Thermostability of the panels, with little VL variation, was shown for a minimum for a minimum of twelve weeks from manufacture. Earlier studies have shown that viral RNA (e.g. influenza) can be reliably detected for up to 196 days (54) and quantified for up to 23 days (56). This study has shown longer-term stability on HIV RNA. However, it must be noted that stability testing was performed in Johannesburg during the South African winter and spring, with temperatures ranging from 8°C to 23°C, but with minimal humidity. More recent studies performed during the hotter months (maximum temperature 31°C) and with increased humidity showed decreases of >1log cp/mL by 10 weeks (personal communication, Dean Sher, SmartSpot Quality Inc.). Further stability evaluations in humid and warmer settings are recommended to ensure similar stability in such settings. In this evaluation, the QA specimens were made to order and generally tested within one week. It should also be noted that the dilution factor was applied to this pilot in order to allow comparison with the original SAVQA data.

In order to determine if specimen variability (71) affects the performance of sites compared to the reference VL, the specimen VL from all sites and the reference VL were compared to the pooled mean VL of all sites. In all cases, the mean VL and the reference VL were similar (-0,02 log cp/mL mean difference), although the reference instrument did produce a higher VL (5.24 log cp/mL vs 5.02 log cp/mL) than all sites in the 5.0 log cp/mL range. This was not clinically significant and did not affect site performance outcome. The bias of the single outlier specimen described (-0.91 log cp/mL bias) was acceptable for verification in terms of the panel design, but unacceptable in the retrospective analysis. However, the site still achieved a module score of 5/6 in the retrospective analysis and patient specimens testing could commence. The benefit of a quality program across multiple sites is that multiple instruments are tested concurrently and panels can be compared to the pooled mean VL rather than only the reference VL; this provides an additional quality control of the reference instrument and can highlight unexpected instability of the quality material. Retrospective analysis of the specimens showed that they potentially can be evaluated at 0.3 and 0.5 log cp/mL bias, and this should be implemented when using this quality panel further.

Ongoing quality monitoring at all levels of a tiered laboratory network is paramount to ensure that patient results are accurate. This can be difficult for POCT instruments placed in remote settings, where quality management options used in centralised laboratories are not feasible,

but where quality monitoring is vital. The quality panels described in this manuscript provide simple and convenient verification and/or EQA options for countries aiming to implement Xpert HIV-1 VL.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Supplementary Table S1: Detailed Site Verification Summary: September 2017-November 2018; Supplementary Table S2: Detailed Site EQA Summary: September 2017-November 2018

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