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Interlaboratory quality control of total HIV-1 DNA load measurement for multicenter reservoir studies

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Background: Viral reservoirs represent an important barrier to HIV cure. Accurate markers of HIV reservoirs are needed to develop multicenter studies. The aim of this multicenter quality control (QC) was to evaluate the inter-laboratory reproducibility of total HIV-1-DNA quantification.

Methods: Ten laboratories of the ANRS-AC11 working group participated by quantifying HIV-DNA with a real-time qPCR assay (Biocentric) in four samples (QCMD).

Results: Good reproducibility was found between laboratories (standard deviation ≤ 0.2 log_{10} copies/10^6 PBMC) for the three positive QC that were correctly classified by each laboratory (QC1<QC2<QC3).

Conclusions: Results of this QC validate the feasibility of multicenter studies using this standardized assay.
1 INTRODUCTION

The ultimate treatment goal in HIV infection would be to eradicate the virus from its reservoirs, which are the main obstacle to curative therapy. To further study these reservoirs and to evaluate new cure strategies, several markers have been proposed but none has been evaluated for using in multicenter clinical studies. Among them, total cell-associated HIV-1 DNA presents the advantage to be feasible within large series of samples and permits longitudinal studies in clinical trials (OPTIPRIM, SPARTAC…). This biomarker has been widely used in both adult and pediatric cohorts. HIV-DNA load correlates with values obtained in the quantitative viral outgrowth assay. It provides useful information at different stages of the infection, irrespective of combined antiretroviral therapy, and with a variety of clinical samples (blood, rectal mucosa, breast milk, and semen). Indeed, it has independent predictive value for progression to AIDS and death. Total HIV-1 DNA load in peripheral blood mononuclear cells (PBMC) correlates with biomarkers of inflammation and immune activation and residual plasma viremia during antiretroviral therapy. Total HIV-1 DNA load could be an accurate, clinically relevant marker for therapeutic decision-making. For example, some treatment de-escalation may not be appropriate for patients with high HIV-1 DNA levels, as stated in recent French guidelines. Baseline HIV-1 DNA levels might prove helpful for choosing patients who will be the best candidates to some therapeutic strategies. It might also prove useful for estimating HIV-1 reservoir to evaluate therapeutic strategies in multicenter trials. Finally, HIV-1 DNA can provide useful information, along with other markers, for HIV cure research.

Several home-made HIV-1 DNA quantification assays by PCR have been used worldwide with different regions amplified and using various standards for quantification and various units making difficult comparisons between studies. One real-time qPCR LTR HIV-1 DNA assay has been developed by the AC11 working group of the Agence Nationale de Recherches sur le SIDA et les hépatites virales (ANRS). This assay is now commercialized (Generic HIV DNA CELL, Biocentric, Bandol, France) and available for the scientific community.

In the current context of therapeutic trials to reduce the HIV reservoir, it is essential to be able to develop multicenter studies. The aim of this national multicenter quality control was to evaluate the inter-laboratory reproducibility of total HIV-1 DNA quantification in ten French laboratories using the same assay (Generic HIV DNA CELL, Biocentric).

2 MATERIAL AND METHODS

Ten laboratories of the AC11 Viral Quantification Working Group of ANRS participated in the qualitative HIV-1 DNA international QCMD quality control (www.qcmd.org/) in 2016. Four coded samples of freeze-dried PBMC pellets were used as the blind test panel and sent to each laboratory. It contained three samples (QC1, QC2, QC3) with HIV-1-infected PBMC, and one (QC4) with PBMC from an uninfected individual. QCMD indicated that QC1, QC2, and QC3 were obtained by dilutions of the same sample. Cell-associated HIV-1 DNA was searched and quantified by real-time PCR with the same Generic HIV DNA CELL assay kit (Biocentric) in the ten laboratories. The performances of this assay have been described previously. Notably the 95% and 50% detection thresholds for the assay were set at six and

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Procedural characteristics used in the ten laboratories and HIV-1 DNA loads obtained for the quality controls</th>
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<tbody>
<tr>
<td>Extraction method</td>
<td>Method to determine the total DNA concentration</td>
</tr>
<tr>
<td>Lab 1 Automatic Qiagen (Qiasymphony)</td>
<td>Albumin qPCR</td>
</tr>
<tr>
<td>Lab 2 Automatic Qiagen (Qiace)</td>
<td>Nanodrop</td>
</tr>
<tr>
<td>Lab 3 Manuel Qiagen (Qiamp DNA Blood)</td>
<td>Nanodrop</td>
</tr>
<tr>
<td>Lab 4 Automatic Qiagen (EZ1 Advanced XL)</td>
<td>Nanodrop</td>
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<tr>
<td>Lab 5 Manuel Qiagen (Qiamp DNA Blood)</td>
<td>Nanodrop</td>
</tr>
<tr>
<td>Lab 6 Automatic Qiagen (Qiasymphony)</td>
<td>Nanodrop</td>
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<tr>
<td>Lab 7 Automatic Qiagen (Qiace)</td>
<td>Nanodrop</td>
</tr>
<tr>
<td>Lab 8 Manuel Qiagen (Qiamp DNA Blood)</td>
<td>Nanodrop</td>
</tr>
<tr>
<td>Lab 9 Automatic Qiagen (Qiasymphony)</td>
<td>Albumin qPCR</td>
</tr>
<tr>
<td>Lab 10 Manuel Macherey-Nagel (Nucleospin Blood)</td>
<td>Nanodrop</td>
</tr>
</tbody>
</table>

Median | (110) 2.0 | (647) 2.8 | (2451) 3.4 |
| Standard deviation | 0.2 | 0.2 | 0.2 |

ND, not detected.
three copies/PCR, that is, 40 and 20 copies/10⁶ cells, respectively (1.6 and 1.3 log). When studying 1 µg total DNA per PCR, all laboratories obtained a good quality score of 0 with the qualitative analysis performed by the QCMD staff. So we decided to collect quantitative results to analyze inter-laboratories reproducibility.

DNA extraction procedure consisted of the automated Qiagen method (Courtabœuf, France) in six laboratories, the manual QIAamp DNA minikit (Qiagen) in three laboratories, and the manual NucleoSpin Blood kit (Macherey-Nagel, Hoerdt, France) in one laboratory (Table 1). The final eluate volume ranged from 50 to 100 µL. The total DNA concentration was determined by fluorescence reading at 260 nm by the Nanodrop method (Labtech, Ringmer, UK) in eight laboratories, and by concurrently amplifying a housekeeping gene by real-time PCR in two laboratories (Table 1). In the Generic HIV DNA CELL kits, the 8E5 cell line was used as standard for HIV-1 DNA quantification. Following the manufacturer's instructions, four dilutions of an extract of 8E5 cells were performed in order to deposit 6000 copies/PCR, 600 copies/PCR, 60 copies/PCR, and 6 copies/PCR. Quantitative results were normalized in number of copies per million cells as described previously. Quantitative data reported by the 10 laboratories in Log₂ copies/million cells were then compared. Logs were used as usually performed for real-time PCR assays and as discussed by Kotton et al. Median and standard deviation were calculated for each positive sample of the quality control (QC1, QC2, QC3).

3 RESULTS

None of the 10 laboratories obtained a positive result with the negative sample (QC4).

Assay performance was analyzed with the three samples of HIV-1-infected PBMC (QC1, QC2, QC3). All three samples were positive and measurable in all 10 laboratories, with the exception of QC1, which was detected but lower than the quantification threshold (70 copies/million PBMC) in one laboratory owing to unsuccessful QIAcube extraction (<2 µg/mL).

The ranges were 1.7-2.3 log₁₀ copies/10⁶ PBMC for QC1, 2.5-3.1 log₁₀ copies/10⁶ PBMC for QC2, and 3.1-3.7 log₁₀ copies/10⁶ PBMC for QC3 (Fig. 1A). The median values and standard deviations (SD) were 2.0 ± 0.2 log₁₀ copies/10⁶ PBMC for QC1, 2.8 ± 0.2 log₁₀ for QC2, and 3.4 ± 0.2 log₁₀ for QC3, and a 95% confidence interval of 0.12 log for each QC. There was no noteworthy influence of the HIV-1 DNA level on the dispersal of values obtained in the 10 laboratories. The results of the ten laboratories showed that the three positive quality controls were correctly classified by each of them: QC1<QC2<QC3. The study of variations of HIV-DNA loads between laboratories showed that the load for each control quality was comprised in the range (median −0.3 log; median + 0.3 log) for each laboratory (Fig. 1B).

Noteworthy, the 10 laboratories quantified HIV-DNA with one of three different batches of the Generic HIV DNA CELL assay kit. The results were similar regardless of the batch used. Despite the use of different extraction methods and the different amounts of DNA analyzed per well (between 0.1 and 1 µg of DNA per well, depending on the concentration of total DNA in the extract), results were reproducible within a small range (Fig. 1B).

4 DISCUSSION

Several markers have been proposed to estimate the size of HIV reservoir, but none is definitely considered as the best one. Among them, total cell-associated HIV DNA load is likely to become an increasingly important marker for the management of HIV-1 infection for several reasons: it is easy to perform in parallel to HIV RNA on the same clinical samples, it is clinically relevant at the different stages of HIV infection, and can be used in research protocols for clinical trials and studies concerning HIV pathogenesis and HIV cure.

The reproducibility of the real-time qPCR LTR HIV-1 DNA assay (Generic HIV DNA CELL, Biocentric) has been previously evaluated in one laboratory for a high-level positive-control sample with mean and standard deviation of 3.30 log copies/10⁶ leukocytes and 0.11 log, respectively, and for a moderate-level positive-control sample with 2.40 log copies/10⁶ leukocytes and 0.18 log.
In this study, we describe the first inter-laboratory external quality control of HIV-1 DNA quantification, implicating ten French virological laboratories using the same commercial assay. No such study has ever been performed for the other markers proposed to estimate the HIV reservoir size. This multicenter evaluation showed that there was a good agreement of HIV-1 DNA loads obtained in ten laboratories by using the standardized Generic HIV DNA CELL assay. Limit of this study is the low number of QCMD samples. However, QCMD is an external international control, bringing the opportunity to evaluate all laboratories in the same conditions. Interestingly, the three samples were at clinically relevant HIV-1 DNA levels. All laboratories detect and quantitate HIV-1 DNA loads with a low standard deviation of 0.2 log. As indicated in Fig. 1, the differences between median of each quality control and result of each laboratory is lower than 0.5 log, as recommended by Kotton et al. Transplantation 2010. Actually Kotton indicated that the precision of quantitative nucleic acid tests to determine viral load are such that changes in values should be at least threefold (0.5 log) to represent biologically important changes. Whatever the extraction protocol and its variable efficiency, the quality of extraction and the normalization of results by measuring the amount of total DNA explored in the real-time PCR permitted to have comparable results of HIV-DNA expressed per million cells between laboratories. The use of different batches of the Generic HIV DNA CELL assay on different thermocyclers did not compromise the results of this inter-laboratory quality control for HIV-1 DNA quantification. No reference assay is available and commercialized for HIV-1 DNA quantification so we could not perform comparison with one gold standard.

In conclusion, the inter-laboratory reproducibility reported here with this quality control is a key step for future applications in multicenter studies and clinical practice. Interestingly we showed the reproducibility of the standardized quantitative total HIV-1 DNA assay developed by the ANRS working group to quantify the HIV-1 DNA load as a marker of HIV reservoir.

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CONFLICTS OF INTEREST

The authors have no conflicts of interest.

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