Comparison of 3 Tests for Plasma HIV-1 RNA Quantitation of Non-B Subtypes in Patients Infected with HIV-1 in N’Djamena-Chad: Cobas AmpliPrep/Cobas TaqMan HIV-1 Test Version 2.0, Abbott m2000 RealTime and Generic HIV Viral Load® Assays

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Abstract

Introduction: Antiretroviral therapy is effective only when it maintains the plasma viral load at an undetectable level or below 50 copies of RNA/ml. Viral Load (VL) is a marker of therapeutic follow-up, particularly within the combination protocols of antivirals. Given the genetic diversity of HIV-1, in Resource-Limited Countries (RLCs), dubious conditions for collecting, conserving and analyzing samples, the choice of one technique over another implies its evaluation on all levels and in particular the cost/benefit ratio.

Objective: The objective of this study was to compare the efficacy of 3 techniques for the measurement of VL for HIV-1 non-B subtypes. The 3 techniques used were: Cobas AmpliPrep/Cobas TaqMan (CAP/CTM) V2.0, Abbott Real-Time and Generic HIV Viral Load®.

Methods: Sample collection was done at the National General Reference Hospital (NGRlH) between June and October 2013. A total of 116 samples were collected from People Living with HIV (PLHIV) and under treatment for at least 6 months. Measurements of VL were done at the AIDS Reference Laboratory at the University Hospital of Liège with the 3 techniques mentioned above.

Results: After amplification by the different techniques, 116 samples were compared with Cobas and Abbott and 42 samples were compared with all 3 techniques. This is due to the lack of plasma for some patients for the Generic HIV Viral Load®. A good correlation is obtained between CAP/CTM and Abbott with R²=0.96016 (p<0.05), while between Abbott vs Generic HIV Viral Load® on one hand and CAP/CTM vs Generic HIV Viral Load® on the other, Pearson correlations (R²) were good and were respectively 0.81064 and 0.72603. This difference with the Generic HIV Viral Load® assay is due to the fact that the plasma has been thawed more than twice. This confirms the fact that plasma freezing, and thawing has more than twice interferes with viral load.

Conclusions: Abbott Real time remains the recommended technique for resource-poor countries, particularly Chad, because of its sensitivity and variability in detecting different subtypes of HIV-1.

Keywords: Plasma viral load; Cobas ampliprep/cobas taqman; Abbott real-time; Generic HIV viral load; Resource-limited countries; HIV-1

Introduction

Antiretroviral Therapy (ART), regardless of the first line or subsequent lines, including after multiple failures, is administered to the patient with the objective to obtain and maintain undetectable plasma Viral Load (VL) [1]. Measurements of VL and resistance tests are used as markers of treatment efficacy. But for their usefulness, these methods require adequate equipment and qualified personnel. The complexity of these techniques and their high cost means that they are not used or at least rarely in Resource-Limited Countries (RLCs). Although there are patients’ follow-up structures such as hospitals and laboratories, most are limited and under-equipped to address the problem of diagnosis and monitoring of HIV infection. In addition, logistical difficulties such as the transport of blood samples and respect
for the cold chain make it difficult to monitor patients living in remote areas, far from specialized centers [2]. The diversity of the different subtypes in RLCs and the proliferation of recombinant forms favored by the emergence of HIV resistance to treatment, which requires the choice of a technique capable of detecting several subtypes of HIV-1, sensitive, reproducible and less expensive. It is for this reason that in this study we will evaluate the sensitivity of different techniques will be evaluated according to different subtypes of HIV-1 to recommend one of them for RLCs. Measurement of VL is used to assess the effectiveness of the treatment and to see the course of the disease. However, in order for the measurement to be significant, the difference between two results must be less than 0.5 log10, a difference from simple to triple. Thus, a VL of 100,000 RNA copies/ml (5 log10) and a VL of 250,000 RNA copies/ml (5.4 log10) are not considered to be significantly different [3,4]. Table 1 summarizes the 3 techniques used in this study [5,6], Therefore, the aim of the study was to evaluate the sensitivity of different VL measurement techniques for HIV-1 non-B subtypes in N’Djamena in Chad.

### Methods

One hundred and sixteen patients under ARV treatment, 78 women and 38 men representing respectively 67.24% and 32.76% recruited on the basis of specific inclusion criteria were enrolled in the study. The average age was 41 ± 6.87 years. Each patient signed an informed consent form after receiving sufficient information about the benefit of the study. The sampling used was consecutive and not exhaustive. The patients were recruited for 5 months (June-October 2013). A venous blood sample was taken at the elbow crease in two EDTA tubes of 5 ml each. The two tubes were stirred gently to mix the blood with the anticoagulant. The blood was centrifuged for 10 min at 2000 g. The plasma was aliquoted in three 2 ml cryotubes (2 to 3 cryotubes were used depending on the plasma volume). They are then put in a box and stored at -80°C and transported to the AIDS laboratory of Liège in Belgium for the analyzes. Three techniques were used to measure the VL on the collected samples of HIV-1 non-B subtypes infected patients. For CAP/CTM, plasma did not undergo freezing or thawing. But for the Abbott Real time and the Generic HIV Viral Load™ assay, the plasma was subjected respectively one and two freezing/thawing.

### Table 1: Comparative table of plasmatic viral load from the 3 techniques

<table>
<thead>
<tr>
<th>Assays</th>
<th>Equipment for amplification/ detection</th>
<th>Principle</th>
<th>Kits</th>
<th>Amplified regions</th>
<th>Types and subtypes detected</th>
<th>Quantitative threshold for detection (RNA copies/ml)</th>
<th>Number of tests/day (8 h per day)</th>
<th>Sample type</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAP/CTM V2.0</td>
<td>Cobas Tagman m2000rt</td>
<td>RT-PCR Real-Time</td>
<td>AmpliPrep/CobasTaQMan V2.0</td>
<td>gag and LTR</td>
<td><strong>M (A à G) + O</strong></td>
<td>20</td>
<td>144</td>
<td>Plasma and DBS</td>
</tr>
<tr>
<td>Abbott</td>
<td>Open platform or FluoroCycler</td>
<td>RT-PCR Real-Time</td>
<td>Abbott Real-time HIV-1 quantitative Assay: m2000system</td>
<td>intégrase, gene and pol</td>
<td><strong>M (A à H) + O + N</strong></td>
<td>40</td>
<td>96-144</td>
<td>Plasma and DBS</td>
</tr>
<tr>
<td>Generic HIV Viral Load</td>
<td>Open platform or FluoroCycler</td>
<td>RT-PCR Real-Time</td>
<td>PCR Real-Time in house</td>
<td>LTR</td>
<td><strong>M (A à H)</strong></td>
<td>390</td>
<td>192</td>
<td>Plasma and DBS</td>
</tr>
</tbody>
</table>

### Extraction

A volume of 1000 μl of plasma is used for the extraction of the RNA while the machine draws only 850 μl. CAP/CTM v2.0 uses automated sample preparation and extraction on the COBAS AmpliPrep instrument using a generic silica-based capture technique.

### Reverse transcription and PCR amplification

Reverse transcription and amplification are performed with the DNA polymerase of thermostable recombinant enzyme Thermus species (Z05). The reaction mixture is heated to allow the anti-sense primers to hybridize specifically to the HIV-1 target RNA and the HIV-1 QS RNA. In the presence of Mg++ and an excess of dNTPs, the polymerase Z05 lengthens the hybridized primers, thus producing strands of DNA complementary to the target RNA.

Cobas ampliprep/cobas TaqMan V2.0 (CAP/CTM)

COBAS® AmpliPrep/COBASTaqman® HIV-1 version 2.0 (CAP/CTM v2.0) is a test based on the in vitro amplification of HIV-1 RNA from plasma collected on EDTA tube. Introduced in 2009 [7,8] it seems to give better results than older versions, notably v1.0 and Amplicor v1.5 [9-11]. It can quantify 20 to 10,000,000 RNA copies/ml of HIV-1 M and O group. The reaction mixture designed to allow equivalent quantitative determination of HIV-1 M and O subtypes contains primers and Probes specific to both viral RNA and standard quantification RNA (QS), which uses reverse transcription and PCR amplification primers that define sequences in the highly conserved regions of the gag gene and the LTR region.

**Abbott real time test ref 2G3190**: The Abbott Real Time HIV-1 assay in vitro is an RT-Real Time (Reverse Transcription in Real Time). A RT-qCR is a conventional PCR after reverse transcription of the RNA into cDNA. In other words, a PCR carried out on a cDNA obtained from an RNA by the action of a Reverse Transcriptase (RTase). Test on plasma and on whole blood collected on blotting paper on adult and pediatric samples. This test allows the detection of subtypes HIV-1: group M (A-H), groups N and O. Real-time PCR is based on the amplification and detection of a fluorescent reporter. The amount of HIV-1 target sequences present at each amplification cycle is measured using fluorescence-labeled oligonucleotide probes on the Abbott m2000rt apparatus. The amplification cycle in which the fluorescent signal is detected by the m2000rt is proportional to the log of the HIV-1 RNA concentration present in the original sample. The detection threshold is 40 RNA copies/ml.

**Biocentric markets the kit for the AmpliPrep/Cobas TaqMan HIV-1 Test Version 2.0, Abbott m2000 RealTime**

**Generic HIV viral load assay (Biocentric, Bandol-France)**: It is of interest in the detection and quantification of HIV-1 RNA targeting the LTR gene, which is described as the least-changing region of the HIV-1 genome [12-14]. Biocentric markets the kit for the measurement of the VL described by Rouet and collaborators in the ANRS (Agence Nationale de Recherche sur le SIDA) HIV quantification working group. This Generic HIV Viral Load™ assay allows the detection of most HIV-1 subtypes in group M [15]. This assay uses, unlike the other assays, extracted RNA for PCR.
amplification. The extraction kit used is the QIAamp DNA mini kit (Qiagen, Courtaboeuf, France). Hundred and Forty microliter of the plasma preserved at -80°C is used for extraction to obtain an eluate of 60μl at the end. Ten microliters of control (RNA virus) is placed in all the samples before extraction. For RT-PCR, a total volume of 25 μl containing 20 μl of Master Mix and 5 μl of RNA extract is used. The primers (forward and reverse: final concentration of 500 nM each), the probe (concentration of 200 nM) and Taqman One-Step RT-PCR Master Mix (Applied Biosystems). The primer used is the HIV1MGF 5’-GCCTCAATAAAGCTTGCCTTGA-3’. The sequence of the antisense primer is HIV1MGR 5’-GGCGCCACTGCTAGAGATTTT-3’. The probe used is HIV1MGProbe 5’-AAGTAGTGTGGCCCGTCTGTTRTKTGACT-3’. The reporter for the probe is 5’-6-carboxyfluorescein and 3’ quencher: 6-carboxytetramethylrhodamine (Applied Biosystems, Foster City, CA). The probe is labeled 5’ with 6-FAM (Fluorochrome) and 3’ with TAMRA (Quencher). The enzyme used is TaqMan one step. The standard curve is made by dilutions of 10 on one control, the Cy5 RNA Detection Probe. The detection threshold is 390 RNA copies/ml for 250 μl of plasma. For each assay, negative and positive control samples were used for all techniques.

Results

Comparison cobas ampliprep/cobas TaqMan V2.0 and abbot real time (n=110)

The VL was measured on all 116 samples first on CAP/CTM v2.0 and then on Abbott Real Time. The comparison was made on 110 samples tested successfully in both automates. Here, the plasma has not undergone freezing or thawing. The results of the VL measurement are given in the following Table 2.

<table>
<thead>
<tr>
<th>Viral Load</th>
<th>Cobas V2.0</th>
<th>Ampliprep/TaqMan</th>
<th>Abbott RealTime</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Copies/ml</td>
<td>Log10</td>
<td>Copies/ml</td>
</tr>
<tr>
<td>Minimum</td>
<td>0,00</td>
<td>0,00</td>
<td>0,00</td>
</tr>
<tr>
<td>Maximum</td>
<td>966 000</td>
<td>5,98</td>
<td>885 196</td>
</tr>
<tr>
<td>Mean</td>
<td>43 301</td>
<td>4,64</td>
<td>43 460</td>
</tr>
<tr>
<td>Median</td>
<td>85</td>
<td>1,93</td>
<td>51</td>
</tr>
<tr>
<td>≥ 1000</td>
<td>48</td>
<td>48</td>
<td></td>
</tr>
<tr>
<td>&lt; 1000</td>
<td>31</td>
<td>29</td>
<td></td>
</tr>
<tr>
<td>Not detected</td>
<td>32</td>
<td>37</td>
<td></td>
</tr>
<tr>
<td>Invalid</td>
<td>5</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

Table 2: Viral load measurement for 3 techniques (n=116)

Invalid samples are due to the fact that the volume of the plasma is very insufficient, and the PLCs cannot read. Some have been supplemented with negative serum. It is important to remember that for two measurements of VL to be significant, the difference must be at least 0.5 log10. Figure 1 present the comparison of CAP/CTM v2.0 and Abbott Real Time. The distribution of points for Abbott and Cobas gives a satisfactory Pearson correlation coefficient (R2=0.96016). Therefore, these Abbott and Cobas measures are statistically similar (p<0.05) for α=5%. We noticed that two samples were detected with Abbott but were not detected with CAP/CTM v2.0. This would be due to a blip during sampling.
Comparison cobas ampliprep/cobas taqMan v2.0, abbott real time and generic HIV viral load assay (n = 42)

For measurement with the Generic HIV Viral Load® assay, 42 samples were tested and compared with the Abbott and Cobas techniques (Table 3).

<table>
<thead>
<tr>
<th>Viral Load</th>
<th>Cobas</th>
<th>Abbott</th>
<th>Generic Load®</th>
<th>HIV Viral Load®</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copies/ml</td>
<td>Log10</td>
<td>Copies/ml</td>
<td>Log10</td>
<td>Copies/ml</td>
</tr>
</tbody>
</table>

Table 3: Viral load measurement for 3 techniques (n=42)

**Figure 2:** Abbott vs Generic HIV Viral Load® correlation

This table gives information on the different values measured by the three different techniques. We noted that there is no difference in the values between the minimum and maximum values (p < 0.05). While for median and mean values, the Generic HIV Viral Load® measurement is different from at least 0.5 Log10. This would certainly be due to the fact that the plasma was thawed at least twice. The correlation between Generic HIV Viral Load® and Abbott is good with a Person correlation coefficient of R^2 of 0.81064 (p<0.05) (Figure 2).

Good correlation between Cobas and Generic HIV Viral Load® with a Pearson R^2 correlation coefficient of 0.72603 (p < 0.05) (Figure 3). Given that each test used at a specific sensitivity to the different HIV-1 groups and subtypes, it goes without saying that some subtypes can be detected by some tests while others cannot.

**Discussion**

In Resources-Limited Countries (RLCs), the assessment of virological failure is often late according to the WHO recommendations [16,17] this is sometimes the cause of accumulation of resistance mutations and reduced effectiveness of second-line drugs. Notwithstanding, some authors report an urgent need for virological monitoring by measuring VL [18-20]. For others, the monitoring of HIV-1 VL in RLCs faces multiple challenges [7,21]. The ANRS assay marketed by Biocentric under the name “Generic HIV Viral load assay” is used routinely in Ivory Coast, Cambodia, Vietnam, Gabon, Cameroon. Various concordance evaluations for this technique with the Cobas Monitor V1.5 were done by Rouet et al. in various countries with an average correlation coefficient of 0.79 [22]. A study published in 2015 presented a correlation coefficient of 0.9452 between Generic HIV Viral Load® assay and the CAP/CTM v2.0 using 39 samples of non-B subtypes of HIV-1 [23]. In this study, comparing the Generic HIV Viral Load® assay with CAP/CTM v2.0, there was a correlation coefficient of 0.72603 with CAP/CTM v2.0 despite the fact that the plasma had undergone two thawing measure with the Generic HIV Viral Load® assay.

Comparison between the two Abbott m2000 Real-Time and CAP/CTM V2.0 assays for the B and non-B subtypes, obtained a correlation coefficient of R^2=0.95 [24,25]. Wirden et al. compared Abbott and CAP/CTM on non-B subtypes and did not obtain a significant difference between the two with R^2=0.84 [25]. Van Rensburg et al. in two cohorts made in Africa and the USA did not find significant differences between the two techniques with a strong correlation of 95% [26]. All these results are in the same direction as this study on these Abbott and Cobas methods because the correlation coefficient obtained in our study (R^2=0.96016) corroborates well with
those of these authors for Abbott and CAP/CTM v2.0. In a study by Margariti et al. in 2016, comparing low VL (<200 RNA copies/ml) in patients infected with subtypes B and non-B, noted a significant difference and concluded that for virological monitoring, a single technique between the two should be used [27]. In this study, for the measurement between CAP/CTM v2.0 and Abbott Real Time, two samples not detected in Abbott but detected in Cobas with 2.23 and 2.68 Log10 respectively. This variation could be due to a blip on two samples. A "blip" is defined as a transient elevation of plasma HIV RNA, usually between 50 and 1000 RNA copies/ml, observed on a single sample, and does not justify the prescription of a Resistance test sensitive to subtypes A-H of group M. While Abbott Real Time detects a reliable measure of HIV-1 VL versus other assays used for genotype specificity and susceptibility [11].

Conclusion
The correlation between COBAS TaqMan/AmpliPrep and Abbott Real Time Viral Load is very good with R2 equal to 0.96016; while the measurement with the Generic HIV Viral Load assay of the Generic HIV Viral Load assay seems to be discordant of the first two but the difference of Log10 is not very significant. This discrepancy could be explained by the fact that for the measurement with the Generic HIV Viral Load assay, the plasma was frozen and thawed twice. Abbott Real time remains the recommended technique for resource-poor countries, particularly Chad, because of its sensitivity and variability in detecting different subtypes of HIV-1.

Acknowledgments
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References

As for the specificity and sensitivity of the techniques, we note that the CAP/CTM v2.0 values are better than those observed for Abbott Real Time. It must also be said that the sensitivity of the techniques differs from one subtype to another. Therefore, CAP/CTM v2.0 is only sensitive to subtypes A-H of group M. While Abbott Real Time detects the N, O groups and the A-G subtypes of the M group. Finally, the Generic HIV Viral Load assay detects the sub-types from A to H of the M group [28]. The Pearson coefficients obtained with CAP/CTM v2.0 and Generic HIV Viral Load assay (0.72603) are due to the freezing and thawing of the plasma more than once; this confirms the fact that plasma freezing, and thawing has more than twice interferes with the number of copies of the VL. The objective of the evaluation the performance of these different techniques of VL is precisely to have a great sensitivity and possibility of detection of the different subtypes in the countries with limited resources and notably in Chad.

Figure 3: CAP/CTM vs Generic HIV Viral Load correlation (n=42)


