Implementation of a Classic Nested PCR DNA for HIV Diagnosis in Kinshasa

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Abstract

Background: The interest in the technique of HIV diagnosis by Polymerase Chain Reaction (PCR) is in: (i) primary infection by the Human Immunodeficiency Virus (HIV), where it is necessary to identify the patient in time, (ii) screening of newborns for HIV-positive mothers, and (iii) the accuracy and reliability of the technique.

Objective: This study aims to assess the feasibility and the performance of the Nested PCR of Deoxyribose Nucleic Acid (DNA) for HIV diagnosis in Kinshasa in order to improve the detection and management of HIV infected patients.

Methods: The present study is a cross-sectional study in collaboration with 3 centers in Kinshasa. Our sample consisted of 171 individuals who have made voluntary testing for HIV. 5.0ml of blood was collected in tubes with anticoagulant, centrifuged for separation into 3 phases. 500μl of buffy coat was collected in a pre-labeled tube. The DNA extraction was made from 200μl of buffy coat using the QIAamp DNA Mini Kit from QIAGEN ® for DNA extraction. Classical HLA PCR and Nested PCR gag and pol were performed to determine the proviral DNA. Nested PCR on env was performed in cases of discordance of the results of gag and pol. The revelation was made under UV after electrophoretic migration on 1% agarose gel. The data were collected in confidentiality in the forms developed and pre-tested for the study, entered on Windows Excel 2007 and analyzed using SPSS 17.0 for Windows. The statistical significance was set at p <0.050.

Results: For Monkole, 22 samples were positive, 18 negative and 6 indeterminate for HIV. For Organization for Disease Prevention and Health Promotion (OPPS), we received 55 samples positive for HIV who served as a control for the study. For Foundation "La Grâce", 30 samples were positive, 25 negative and 15 indeterminate for HIV. This gives a total of 107 samples positive, 43 negative and 21 indeterminate for HIV. After amplification by Nested PCR DNA, we have a total of 112 samples positive and 59 HIV-negative. Of the 21
indeterminate samples, 3 were positive and 18 negative by PCR. 2 samples were negative by RDT but positive by PCR.

**Conclusion:** The Rapid Diagnostic Test (RDT) results and those of PCR DNA Nested allowed us to evaluate and validate the technic of diagnosis of HIV in Kinshasa by this new technic. This method provides a more reliable alternative for the diagnosis of HIV Kinshasa.

**Key words:** diagnosis, HIV, Nested PCR, Kinshasa

**Introduction**

The technique of amplification by Polymerase Chain Reaction (PCR) generated a lot of enthusiasm. Its "relatively simple" realization has led many teams to try its realization. Its principle is attractive since the shortcomings of previous methods can be related to the mode of virus replication, with a possible eclipse phase and especially the need for integration in the cell genome without the virus or its antigens are expressed necessarily continuously. It was extremely tempting to go "dig out" the provirus where he was hiding in the Deoxyribose Nucleic Acid (DNA), within the genome of the infected cell. But the relative ease of technical execution, are increasing difficulties of interpretation and training of appropriate staff.

The medical and scientific interest in the technique of HIV diagnosis by PCR lies in: (i) primary infection, where it is necessary to identify the patient in time, (ii) screening of newborns of mothers HIV-positive, and (iii) the accuracy and reliability of the technique.

This study aims to implement a Classic Nested PCR DNA for HIV diagnosis in Kinshasa in order to improve the detection and management of HIV-infected patients.

**Methods**

**Framework**

The present study is a cross-sectional study that was conducted in collaboration with the Department of Infectious Diseases (Monkole-3) Centre Hospitalier Monkole (CHM) which is a screening service and support for People Living with HIV (PLHIV), the non-profit Organization for Disease Prevention and Health Promotion (OPPS) and Foundation "La Grâce" which are associations that participate in voluntary HIV testing in the community, all in the city of Kinshasa. This study took place from 01 August to 27 October 2012.

**Patients**

Only people willing to voluntary testing for HIV were selected for this study. Our sample consisted of 171 individuals who have made voluntary testing for HIV by Rapid Diagnosis Test (RDT). Seventy (70) persons were screened through the Foundation "La Grâce", 55 through the OPPS and 46 through the CHM. All were willing and signed consent for screening with the respective institutions. No results for RDTs had been revealed before the PCR results.
**Blood samples**

A minimum of 5.0 ml of blood was collected into tubes containing EDTA from the respective testing centers. The collected blood was centrifuged at 3000 rotation per minute (rpm) for 10 minutes to obtain a separation into 3 phases: the supernatant (plasma), the intermediate zone (buffy coat) and the pellet. Five hundred microliters (500μl) of buffy coat was collected in a pre-labeled tube and stored at 4 °C before being sent to the laboratory of Molecular Biology of the Faculty of Medicine for analysis.

**Extraction and DNA amplification**

The DNA extraction was made from 200μl of buffy coat using the QIAamp DNA Mini Kit from QIAGEN® for DNA extraction. The DNA extract was stored at -20 °C.

After extraction, a Classic PCR on HLA and Nested PCR on gag and pol region were performed to determine the proviral DNA. Nested PCR on env region was performed about in cases of discordant results of gag and pol. The primers for amplification were HLA: GH26 and GH27 (Table 1). The primers for the first stage of the HIV PCR were: GAG1 and GAG4 for gag amplification, POLITG1 and POLITG4 for pol, and ENV1 and ENV4 for env (Table 1). The primers for PCR were Nested: GAG2 and GAG3 for gag, POLITG2 and POLITG3 for pol, and ENV2 and ENV3 for env (Table 1). PCR assays were carried out under the conditions described in Table 2.

**Interpretation of Results**

The revelation was made under UV after electrophoretic migration on agarose gel 1% prepared in 1X TAE (Tris / acetate / EDTA). For a sample to be counted positive for HIV by PCR, it should be positive for HLA amplification and positive amplifications for two proviral (gag / pol, gag / env and/or pol / env).

**Statistical Analysis**

The data were collected in confidentiality in the data developed and pre-tested to work. They were transcribed on Windows Excel version 2007 and analyzed on SPSS version 17.0 for Windows. The statistical significance was set at p <0.050.

**Results**

A total of 171 samples were received for Nested PCR DNA: 46 from CHM, 55 from OPPS and 70 from Foundation "La Grâce."

**RDT’s status**

For CHM, 22 samples were positive, 18 negative and 6 indeterminate for HIV. For the OPPS, we received 55 samples positive for HIV who served as a control for the study. For Foundation "La Grâce", 30 samples were positive, 25 negative and 15 indeterminate for HIV.
This gives a total of 107 samples positive, 43 negative and 21 indeterminate for HIV (Tables 3 and 4).

**DNA amplification**

After PCR amplification Nestled DNA, we have a total of 112 samples positive and 59 HIV-negative. Of the 21 indeterminate samples, 3 were positive and 18 negative by PCR. Two (2) samples were negative by RDT but positive PCR (Tables 3 and 4).

**Discussion**

This study was to assess the feasibility and the performance of the Nested PCR DNA for HIV diagnosis in Kinshasa. We had a total of 171 blood samples for this work from three different centers: Centre Hospitalier Monkole (CHM), which contributed 46 samples, the Foundation "La Grâce" with 70 samples and OPPS with 55 samples. All samples provided were previously tested for HIV using Rapid Diagnostic Tests (RDTs). The tests used, as by national recommendations, are: Determine ½, Uni-Gold and Double Check.\(^1\)

For CHM, we received on 46 samples, 22 were positive, 18 negative and 6 indeterminate by RDTs. After Nested PCR diagnosis, we had 22 positive and 24 negative samples. The six samples that were indeterminate with RDTs were clearly negative by PCR.

For Foundation "La Grâce", out of the 70 samples received, 30 were positive, 25 negative and 15 indeterminate. After Nested PCR diagnosis, we had 35 positive samples and 35 negative for HIV. Two samples that were negative with RDTs were positive by PCR. Of the 15 indeterminate samples by RDTs, 3 were positive and 12 were negative by Nested PCR. The two discordant samples as well as indeterminate by RDTs that turned positive by Nested PCR were tested by ELISA, as 3rd diagnostic method to confirm the results, as these were positive by Nested PCR.

For OPPS, the 55 samples received were all positive tests for HIV by RDTs. After Nested PCR, all 55 samples were confirmed positive for HIV.

This shows us how much the Nested PCR is more specific and more accurate than RDTs. The Indeterminate results with RDTs were classified positive or negative, and then confirmed by ELISA. Nested PCR gives us much more accurate results than the diagnostic techniques used so far in our centers and laboratories. ELISA p24 antigen reduces the diagnostic gap to four weeks after infection, but in some cases it may remain negative for more than 3 months.\(^12-15\) Commercial standardized tests, or RDTs, facilitated the diagnosis of HIV in their simplicity.\(^16\) However, they are still shadows hanging over the specificity and accuracy of these since they were designed based on specific markers for certain types of HIV.\(^12\) This confirms some of the deficiencies identified in our results RDTs and by Kleinman and colleagues.\(^17\) The shortcomings of the RDTs are mostly the causes of the false results in the diagnosis of the infection.

In case of indeterminate results, the patients are recommended to come back in 3 weeks to get tested again to see if there would be a more precise result.\(^11\) However, it is very unlikely for a indeterminate-patient to come back for testing afterward fearing the unpleasant result. This
implies that, in our case, 14.29% of this class would have gone with the infection without coming back for their result.

The principle of the Nested PCR DNA is that the product of the first PCR is used for the second amplification. Thus giving a much more precise result for a qualitative PCR. The \textit{env}, \textit{gag} and \textit{pol} regions are used for the amplification because they are the common regions for the virus genome.\textsuperscript{1,5,8-10} Every mature and functional virus does have these regions. Therefore, they are present at the integrated proviral DNA. Even at different period of the virus cycle, these regions can be found.\textsuperscript{8-10} These are some of the advantages of the technique; knowing that at any stage of the cycle, we can dig-out the virus from where it is hiding.

Although laborious, the Nested PCR technique has proven to be an accurate tool for diagnosing HIV.\textsuperscript{1,12,18} The feasibility of this PCR does not required very sophisticated material. It requires a classic thermocycleur for the amplification, a kit for the migration and an UV table for the visualization; most material that could be found in developing country.

HIV diagnosis by Nested PCR is a useful tool for management accurately and early. Especially in cases that are indeterminate or doubtful.

**Conclusion**

The RDTs results and those of Nested PCR DNA allowed us to evaluate the performance of this diagnosis method for HIV in Kinshasa. Discrepancies were confirmed by ELISA for the Nested PCR results. This method provides a more reliable alternative for the diagnosis of HIV Kinshasa. It should be recommended for the general laboratory to strengthen HIV-infected patients diagnosis.

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**Conflicts of Interest:** The authors declare that they have no competing interests.

**References**


**Table 1**: Primes and sequences for the PCR reactions

<table>
<thead>
<tr>
<th>PCR</th>
<th>Primers</th>
<th>Sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>HLA</td>
<td>GH26 forward</td>
<td>5'-GTGCTGCAGGTGTTAATACT-3'</td>
</tr>
<tr>
<td></td>
<td>GH27 reverse</td>
<td>5'-CACCGATCCGATGTTAATACT-3'</td>
</tr>
<tr>
<td>Gag</td>
<td>GAG1 forward</td>
<td>5'-GTGCACTACAGGCCATATCCACC-3'</td>
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</table>
Table 2: PCR Cycle (temperature per cycle)

<table>
<thead>
<tr>
<th></th>
<th>gag and pol</th>
<th>env and Nested pol</th>
<th>HLA and Nested gag</th>
<th>Nested env</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Denaturation</td>
<td>95°C : 9min</td>
<td>95°C : 9min</td>
<td>95°C : 9min</td>
<td>95°C : 9min</td>
</tr>
<tr>
<td>Denaturation per cycle</td>
<td>94°C : 1min</td>
<td>94°C : 1min</td>
<td>94°C : 1min</td>
<td>94°C : 1min</td>
</tr>
<tr>
<td>Hybridation per cycle</td>
<td>50°C : 1min</td>
<td>50°C : 1min</td>
<td>55°C : 1min</td>
<td>45°C : 1min</td>
</tr>
<tr>
<td>Elongation per cycle</td>
<td>72°C : 1min</td>
<td>72°C : 1min</td>
<td>72°C : 1min</td>
<td>72°C : 1min</td>
</tr>
<tr>
<td>Elongation finale</td>
<td>72°C : 10min</td>
<td>72°C : 10min</td>
<td>72°C : 10min</td>
<td>72°C : 10min</td>
</tr>
<tr>
<td>Hold</td>
<td>4°C</td>
<td>4°C</td>
<td>4°C</td>
<td>4°C</td>
</tr>
<tr>
<td>Number of cycle</td>
<td>35</td>
<td>30</td>
<td>25</td>
<td>25</td>
</tr>
</tbody>
</table>

Table 3: Confirmation Diagnostic VIH RDT and PCR

<table>
<thead>
<tr>
<th>TD R</th>
<th>HLA</th>
<th>gag</th>
<th>pol</th>
<th>env*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Positive</td>
<td>107</td>
<td>106</td>
<td>1</td>
<td>105</td>
</tr>
<tr>
<td>Negative</td>
<td>43</td>
<td>40</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Indeterminate</td>
<td>21</td>
<td>20</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>171</td>
<td>166</td>
<td>5</td>
<td>110</td>
</tr>
</tbody>
</table>

*PCR env to confirm status after difference of results with gag and pol

Table 4: Confirmation of results per center

<table>
<thead>
<tr>
<th>Centre Hospitalier Monkole</th>
<th>Foundation «La Grâce»</th>
<th>OPPS</th>
<th>Total</th>
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</thead>
<tbody>
<tr>
<td>TDR+ =&gt; PCR+</td>
<td>22</td>
<td>30</td>
<td>55</td>
</tr>
<tr>
<td>TDR+ =&gt; PCR-</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>TDR- =&gt; PCR+</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>TDR- =&gt; PCR-</td>
<td>18</td>
<td>23</td>
<td>0</td>
</tr>
<tr>
<td>TDR± =&gt; PCR+</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>TDR± =&gt; PCR-</td>
<td>6</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>46</td>
<td>70</td>
<td>55</td>
</tr>
</tbody>
</table>

*TDR± = Result Indeterminate with RDT