Genetic Diversity and Transmitted Drug Resistance of HIV-1 Subtypes in Blood Donors Candidates in Northern Mozambique

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Abstract

In order to evaluate the genetic diversity of HIV-1 subtypes and transmitted drug resistance (TDR) mutations in northern Mozambique, we analyzed 120 plasma specimens obtained from drug naive blood donors candidates, who tested positive during routine HIV screening in three blood banks in that region. The genotyping for HIV-1 resistance was performed using Trugene Genotyping System™. HIV-1 genetic subtypes were defined based on entire PR and partial RT gene regions, using REGA HIV-1 Subtyping Tool version 3.0 algorithm and confirmed by phylogenetic inference, using NJ algorithm. The majority of genotyped samples were classified as HIV-1 subtype C (80.0%), followed by subtype A1 (10.5%), subtype D (3.2%) and subtype G (2.1%). The inter-subtype recombinant forms (A1/C, A1/D and C/D) were identified in four participants (4.2%). TDR mutations associated with nucleoside and non-nucleoside reverse transcriptase inhibitors (K219E, G190A, K101E and K103N) were observed in five (5.3%) subjects.

Although a large proportion of HIV-1 subtype C was observed, non-C and recombinant forms together correspond to 20% in this study, which is different from what was described in central and southern regions, where subtype C was almost 100%. However, the profile mutations from this study correlate with the ARVs used both for first line schemes and PMTCT in Mozambique. This data reinforces the necessity of continuous surveillance of HIV-1 diversity, TDR and routes of spread, through inner cities of Mozambique to understand better the dynamics of the HIV-1 epidemic and support national public health policies in the country.

Keywords: Mozambique; HIV-1 genetic diversity; Resistance mutation; Blood donors

Introduction

Global HIV infection reached unprecedented levels in the last 25 years with the number of people living with HIV/AIDS estimated to be at 36.9 million in 2010 [2]. The UNAIDS estimates that 1.4 million people were newly infected with HIV in 2012 [3]. In Africa, the region most affected by HIV, the number of adults infected with HIV/AIDS is estimated at 25.9 million in 2010 [4]. Mozambique, a Sub-Saharan country faces serious HIV epidemics, with an overall prevalence of 11.5% (1.6 million people) [1] within the adult population. However, HIV prevalence varied from 2.5% in the northern region to 4.8% in the central region. Phylogenetic analysis has revealed an extensive genetic diversity of HIV in Africa, composed of types 1 and 2, groups M, N, O and P, subtypes A-K, circulating recombinant forms (CRFs) and many unique recombinant forms (URFs) [2-5]. Mozambique, a Sub-Saharan country faces serious HIV epidemics, with an overall prevalence of 11.5% (1.6 million people) [1] within the adult population. However, the profile mutations from this study correlate with the ARVs used both for first line schemes and PMTCT in Mozambique. This data reinforces the necessity of continuous surveillance of HIV-1 diversity, TDR and routes of spread, through inner cities of Mozambique to understand better the dynamics of the HIV-1 epidemic and support national public health policies in the country.

Materials and Methods

Ethical considerations

This study was approved by the National Committee for Bioethics in Health (CNBS), Mozambique. Reference number 189/CNBS, May, 26th, 2009. Written informed consent was obtained from each participant prior to conducting any study procedure. The study was conducted according to good clinical and laboratory practices.

Study population and HIV screening

The study was conducted in three out of four blood banks (Hospital Central de Nampula, Hospital Geral de Nacala-Porto and Hospital Provincial de Pemba) in the northern region, between November 2009 and June 2010. The blood bank of Lichinga in Niassa Province was excluded for the study because during the study period the ELISA plate reader used for HIV screening in the laboratory was broken. Initially, all enrolled participants provided a total of 500 ml of whole blood. A rapid diagnosis of HIV in Mozambique is limited to the city of Maputo and Beira. Although in 2010, 94% of the adult population from the northern and central regions of Mozambique were surveyed, only 70% of the people tested positive for HIV [1]. However, the profile mutations from this study correlate with the ARVs used both for first line schemes and PMTCT in Mozambique. This data reinforces the necessity of continuous surveillance of HIV-1 diversity, TDR and routes of spread, through inner cities of Mozambique to understand better the dynamics of the HIV-1 epidemic and support national public health policies in the country.

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blood in a blood bag. The specimens were screened for HIV according to the national algorithm, using Genescreen™ HIV-1/2 version 2, (Bio-Rad, France) and Determine HIV-1/2 (Abbott, USA). For the HIV positive samples, 50 µl of whole blood was used to prepare Dried Blood Spots (DBS) and 6.0 ml of plasma were prepared for each participant. All plasma and DBS specimens were sent to the reference laboratory of the Instituto Nacional de Saúde in Maputo, Mozambique where a confirmatory test was performed in plasma specimens, using Genescreen™ HIV-1/2 version 2, (Bio-Rad, France); Murex HIV Ag/Ab combination, (Abbott, UK) and Amplicor HIV-1 DNA test, version 1.5 (Roche Diagnostics, Germany) was performed in DBS specimens. For the specimens that were confirmed positive at the reference laboratory, viral load was measured in plasma specimen using COBAS Ampliprep/TaqMan 96 version 2.0, (Roche Diagnostics, Germany). The remaining sample portion was stored and later was used to prepare Proficiency Testing Panel at National Institute of Health Reference Laboratory. The permission for it was asked from the participants during the enrolment for the study.

RNA extraction, RT-PCR and sequencing for genotyping

HIV RNA was extracted from 140 ul of plasma, using QIAamp RNA Mini Kit, according to manufacturer’s instructions (QIAGEN, USA). The RT-PCR, one-step PCR and sequencing was performed using Trugene HIV-1 Genotyping System, (Siemens Diagnostics, US), according to manufacturer’s instructions and the sequences were analyzed using OpenGene DNA software.

HIV-1 subtyping, phylogenetic and resistance mutation analysis

For subtype analysis the sequences were submitted to the REGA HIV-1 Subtyping Tool version 3.0 [14,15] and confirmed with phylogenetic analysis, where the nucleotide pol gene sequences from this study were aligned together with representative reference sequences (group M subtypes A, B, C, D and G) retrieved from Los Alamos HIV Sequence Database [16], using ClustalW algorithm, available through MEGA version 5.0 software [17]. Tamura-Nei was the best fitting substitution model [18] with 100 replicates bootstrap and group O was used to root the phylogenetic tree. Neighbor-joining (NJ) algorithm [15] was used for phylogenetic inferences of all pure subtypes from this study.

Furthermore, to investigate routes of transmission of subtype C, based on virus similarity, the Subtype C Maximum Likelihood (ML) tree was performed using PhyML 3.0 algorithm [19], GTR was the best fitting substitution model [20], and SPR was applied to search the tree space. Recombination analysis was performed by bootscanning, using SimPlot.

Transmitted Drug Resistance mutations were analyzed using the Calibrated Population Resistance Tool (CPR) [21].

Results

Study population

From November 2009 through June 2010, a total of 200 blood donors were included in this study. Of these, 120 (60%) were serologically reactive for HIV-1 and confirmed by Amplicor HIV-1 DNA test version 1.5.

The median age of participants was 29 years old, and, in terms of gender distribution, all were males donating blood for the first time. None of the participants knew their HIV status. The main reason for donation was to replace the stock at the blood bank due to a transfusion that had to be done for a relative. Plasma viral load was high in most infected subjects with a mean of 5.0 log 10. Ninety two (76.7%) participants were from Nampula city, seventeen (14.2%) from Pemba, and eleven (9.2%) from Nacala Porto.

HIV-1 genetic diversity

Ninety five (79.2%) out of 120 samples were successfully genotyped for HIV-1 resistance, generating good quality sequences from HIV pol gene (entire Protease and partial RT). Among the 95 sequences submitted for phylogenetic analysis, 91 (95.8%) were classified as HIV-1 group M pure subtypes (C, A1, D and G) and 4 (4.2%) were classified as recombinant forms (A1/C, A1/D and C/D). Boot scanning analysis of concatenated genes, using Simplot confirmed the recombination events (Figure 1).

According to the REGA HIV-1 Subtyping Tool analysis, confirmed by phylogenetic analysis, 76 (80.0%) sequences clustered with subtype C, ten (10.5%) with subtype A1, three (3.2%) with subtype D and

Figure 1: Four isolates from the study: NPL200, NPL 209, NPL 123 and NPL070, did not cluster in HIV pure subtypes. They were aligned with HIV-1 subtype A, B, C, D and G. Simplot analysis were performed for each isolate and are illustrated above. Red line indicates subtype A strains, yellow subtype C and blue subtype D.
two (2.1%) sequences with subtype G (Table 1). In addition, for the Maximum Likelihood analysis of subtype C a large dataset of 1786 subtype C reference sequences from southern and east African countries, available in full or near full length genome in the Los Alamos HIV Database, were aligned together with 75 pure subtype C sequences from our study. In this analysis was observed that the subtype C sequences from Mozambique, including references, were segregate in small phylogenetic sub clusters, showing an almost uniform distribution along the phylogenetic tree (Figure 2). The largest cluster involving Mozambican strains were observed mainly with South African sequences. In contrast, subtype C strains recovered in eastern African countries (Uganda, Tanzania, Kenya and Burundi) seems to be relatively more diverse and clustered in a separate group.

Drug resistance mutations and other polymorphisms

Among the 95 blood donors analyzed, 19 (20.0%) have shown at least one major or minor mutation in the RT or PR region. The CPR analysis identified four mutations in five subjects (5.3%) as surveillance drug resistance mutation (SDRM) [21], occurring alone or combined each other. Four out of five subjects with SDRM were carrying NNRTIs mutations (K103N, n=1; K101E, n=2; G190A, n=2) and one NRTI (K219E) was found in one subject (Table 2). The mutations G190A and K103N are known to confer high-level resistance to Efavirenz (EFV) and Nevirapine (NVP), while K101E causes intermediate-level resistance to NVP and low-level to EFV. On the other hand, the Timidine analogue mutation (K219E) reported in one subject is known to be associated with decreased susceptibility to Zidovudine (AZT) and possibly to Stavudine [22,23]. Beside the SDRM, were also observed in the RT region, the minor mutations (K101H and V106I) in one subject each and E138A in nine subjects). Both K101H and E138A are common accessory polymorphic mutations associated with decreased susceptibility to two second generation NNRTIs (Rilpivirine and Etravirine) [22]. Additionally, K101H is also associated with decreased response to NVP and EFV [22,24]. No major mutations associated with resistance to protease inhibitors (PIs) was identified in our study. However, five minor PI mutations were observed (L10I, n=1; K20I, n=2; V11I, n=1; T74S, n=6; L10F, n=1). The combination of T74S with E138A observed in two subjects could cause low-level resistance to Rilpivirine and Nelfinavir, while two participants harboring only T74S, display low level resistance to Nelfinavir [22]. In one participant, the mutation K20I was associated with V11I, causing low-level resistance to Nelfinavir and potential low-level was observed when combined with L10I (Table 2).

Although not considered in WHO SDRM list, several polymorphisms listed in the International Antiviral Society (IAS) (2014) panel were observed in the PI region, of which M36I (83.2%); H69K (94.7%); L89M (83.2%) and I93L (74.7%) (Figure 3).

Discussion

HIV-1 genetic diversity

In this study we found that HIV subtype C is the most frequent in northern Mozambique. However, other subtypes (A1, D, G) and recombinant forms were also detected. The finding of subtype C in three different cities from northern Mozambique reinforce the hypothesis that this subtype has been circulating as a major lineage, not only in the northern of Mozambique but also in the south and center of the country [8,9]. In fact, there is strong evidence of subtype C epidemic spread throughout the country [8-10,12,25], as well as in other southern African countries [26,27].

Additionally, in a separated phylogenetic analysis of pure subtype C, strains from Mozambique clustered mainly with South African strains, which suggest an epidemiological linkage between these countries. These findings are in line with Msimanga et al., who discuss the impact of migratory movements in the spread of epidemics [26]. Our result is also consistent with Baltazar et al. who reported 22.3% HIV prevalence in Mozambican mine workers in South Africa [28]. These findings are consistent; given the intensive migratory flux due to labor necessity mainly of Mozambican mine workers in South Africa. In addition, the intensive flux of truck drivers through the Maputo corridor, which connects South Africa to Maputo Port in the south and to the central provinces (Manica, Tete, Zambézia), may be a favorable occasion of virus spread in the region [26].

In our study, HIV-1 non-C subtypes (A1, D, G) together with recombinants, were reported in a relatively high frequency (20%). Moreover, in Pemba, non-C subtypes from our study represented 50% (6 of 12) of the total subjects analyzed. This finding is in line with Abreu et al. who reported high diversity of HIV in the same region in 2002 [13]. However, our results are different from previous studies, which reported almost 100% subtype C in central and southern Mozambique [8,9,25]. These findings emphasize that although the HIV-1 epidemic in Mozambique is dominated by subtype C, non-C subtypes (A1, D, G) and inter-subtype combination are prominent features to be considered in northern region. The HIV epidemic pattern reported in northern Mozambique may somehow reflect the influence of A1 and D subtypes epidemic in east African countries (Kenya, Rwanda, Tanzania.

<table>
<thead>
<tr>
<th>Subtypes</th>
<th>Nampula</th>
<th>Nacala porto</th>
<th>Pemba</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>67 (83.8%)</td>
<td>2 (66.7)</td>
<td>6 (50.0%)</td>
<td>75</td>
</tr>
<tr>
<td>A1</td>
<td>6 (7.5%)</td>
<td>-</td>
<td>-</td>
<td>10</td>
</tr>
<tr>
<td>D</td>
<td>2 (2.5%)</td>
<td>1 (33.3)</td>
<td>-</td>
<td>3</td>
</tr>
<tr>
<td>G</td>
<td>-</td>
<td>-</td>
<td>2 (16.7%)</td>
<td>2</td>
</tr>
<tr>
<td>URFs</td>
<td>5 (6.3%)</td>
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<td>-</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>80</td>
<td>3</td>
<td>12</td>
<td>95</td>
</tr>
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</table>

Table 1: HIV-1 Subtypes distribution per cities.

Figure 2: Phylogenetic analysis of HIV-1 subtype C sequences from blood donors retrieved in Nampula, Nacala Porto and Pemba, northern Mozambique. 75 pure subtype C sequences of HIV-1, entire protease and partial RT genes (codon 40-247) from our study, were aligned with 1711 pure subtype C from southern, central and east Africa. Colors were used to identify sequences from same region or country. Group O was used as out group. The Maximum Likelihood tree was constructed using PhyML 3.0. GTR substitution model was applied, with 100 replicates bootstrap.
Drug resistance mutations and other polymorphisms

In this study, we found evidence of HIVDR mutations for the first time in the northern region, and it is the first report of HIVDR mutations in blood donor candidates in Mozambique. For both RT and PR, SDRM level was low. This finding is in line with Bila et al. who reported in naïve pregnant women in 2009 a low level (<5.0%) of TDR in southern, while an intermediate level (5.0-15.0%) was reported in central Mozambique [9]. On the other hand, data available, from naive pregnant women did not report any TDR in southern, central, and northern Mozambique in 2002, but high molecular diversity of HIV-1 subtypes was described in the northern region [13]. All together this data shows that TDR has been emerging in Mozambique. The trend of TDR elucidated in this study is consistent to global studies which indicate low prevalence of TDR in sub-Saharan Africa when compared with North America, Latin America and Europe [31-33], but with increasing trend from 2004 to 2010 [33,34]. The mutations K103N and K101E observed in our study were described in previous study in Uganda) [27]. In fact this finding is plausible given the historical relationship, which allows intensive flux of people between countries.

Overall, results from phylogenetic analysis of subtype C, A1, D and G from this study, suggest that Mozambique has an epidemiological linkage with most southern African countries, characterized by multiple introductions and circulation of HIV-1 strains in the country. This finding was also observed by Bartolo et al. in southern Mozambique in 2009 [25]. However, advanced phylogenetic analysis and continued monitoring of the different strains is necessary to understand better the origin of the strains circulating in northern Mozambique and also the possible spread to the central and southern regions of the country. The introduction of additional subtypes may lead to the formation of novel recombinant forms and the emergence of variants with biological and epidemiological property changes. The CRF01_AE is one of the most studied and has shown to be more pathogenic than pure subtype B in Singapore. A changing of molecular epidemiology profile may have important consequences on diagnostic testing, therapeutic response to treatment and vaccine candidates design [29,30].

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Subtype</th>
<th>PR</th>
<th>RT</th>
<th>Potential Low</th>
<th>Low</th>
<th>Intermediate</th>
<th>High</th>
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<td>G</td>
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<td>K101E, G190A</td>
<td>ETR, RPV</td>
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<td>T74S</td>
<td>E138A</td>
<td>NFV, ETR, RPV</td>
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<td>K219E, E138A</td>
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<td>ETR, NFV, RPV</td>
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<td>K103N</td>
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<td>EFV, ETR,</td>
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</table>

Table 2: HIV-1 drug resistance profile mutation.
in pregnant women naive to ARVs in Beira and E138A in Beira and Maputo in 2009 [9]. The occurrence of K103N and G190A will inhibit the use of Nevirapine and Efavirenz drugs, which are the backbone of ART first-line schemes in Mozambique. Thereby mutations will force inclusion of PI inhibitors in first-line schemes, which will further limit therapeutic schemes options for the second line. Additionally, the occurrence of E138A can compromise the use of NNRTIs second generation (Rilpivirine and Etravirine) in first line schemes. This scenario may compromise treatment strategies, especially in resource-poor countries. Furthermore, the mutation T74S, reported in this study is a common polymorphism in subtype C, which is known to be associated with restoring fitness of multi-drug resistant virus in vitro [35].

The polymorphisms M36I, H69K, L89M and 91L reported in our study were also observed in high frequency in other studies evolving non-B subtypes and are considered as polymorphic signatures for non-B subtypes virus [9,25].

Our study is a strong evidence of TDR mutation circulation in northern Mozambique. However, continued surveys with nationally representative pre-treatment patients could provide more details to inform first-line schemes guidelines in a national context, leading to an increase of ARV access in the country.

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References


