Next-generation sequencing provides an added value in determining drug resistance and viral tropism in Cameroonian HIV-1 vertically infected children

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Abstract

With limited and low-genetic barrier drugs used for the prevention of mother-to-child transmission (PMTCT) of HIV in sub-Saharan Africa, vertically transmitted HIV-1 drug-resistance (HIVDR) is concerning and might prompt optimal pediatric strategies.

The aim of this study was to ascertain HIVDR and viral-tropism in majority and minority populations among Cameroonian vertically infected children.

A comparative analysis among 18 HIV-infected children (7 from PMTCT-exposed mothers and 11 from mothers without PMTCT-exposure) was performed. HIVDR and HIV co-receptor usage was evaluated by analyzing sequences obtained by both Sanger sequencing and ultra-deep 454-pyrosequencing (UDPS), set at 1% threshold.

Overall, median (interquartile range) age, viremia, and CD4 count were 6 (4–10) years, 5.5 (4.9–6.0) log_{10} copies/mL, and 526 (282–645) cells/mm^3, respectively. All children had wild-type viruses through both Sanger sequencing and UDPS, except for 1 PMTCT-exposed infant harboring minority K103N (8.31%), born to a mother exposed to AZT+3TC+NVP. X4-tropic viruses were found in 5 of 15 (33.3%) children (including 2 cases detected only by UDPS). Rate of X4-tropic viruses was 0% (0/6) below 5 years (also as minority species), and became relatively high above 5 years (55.6% [5/9], P = .040). X4-tropic viruses were higher with CD4 ≤ 15% (4/9 [44.4%]) versus CD4 > 15% (1/6 [16.7%]), P = .580; similarly for CD4 < 200 (3/4 [75%]) versus CD4 > 200 (2/11 [18.2%]) cells/mm^3, P = .077.

NGS has the ability of excluding NRTI- and NNRTI-mutations as minority species in all but 1 children, thus supporting the safe use of these drug-classes in those without such mutations, henceforth sparing ritonavir-boosted protease inhibitors or integrase inhibitors for the few remaining cases. In children under five years, X4-tropic variants would be rare, suggesting vertical-transmission with CCR5-tropic viruses and possible maraviroc usage at younger ages.

Abbreviations: 3TC = lamivudine, ABC = abacavir, AZT = zidovudine, DRRMs = drug resistance mutations, EFV = efavirenz, ETR = etravirine, HAART = highly active antiretroviral therapy, HIV-1 = human immunodeficiency virus type 1, HIVDR = HIV-1 drug-resistance, NGS = next-generation sequencing, NNRTI = non-nucleoside reverse transcriptase inhibitors, NRTI = nucleoside reverse transcriptase inhibitors, NVP = nevirapine, PCR = polymerase chain reaction, PI/r = protease inhibitors boosted with ritonavir, PMTCT = prevention of mother-to-child transmission, PR = Protease, RLS = resource-limited setting, RPV = rilpivirine, RT = reverse transcriptase, RT-PCR = reverse transcriptase polymerase chain reaction, Sd-NVP = single dose nevirapine, SSA = sub-Saharan Africa, UDPS = ultra-deep 454-pyrosequencing, VF = virological failure.

Keywords: children, coreceptor usage, HIV-1 drug-resistance, next-generation sequencing, PMTCT, sanger sequencing
1. Introduction

Despite increasing coverage (to about 61%) in prevention of mother-to-child transmission (PMTCT), human immunodeficiency virus type 1 (HIV-1) vertical-transmission remains consistent in sub-Saharan Africa (SSA). More so, although progress in PMTCT (from single-dose nevirapine [sd-NVP] to option-B+) has been reducing HIV-1 vertical-transmission, infected children stand at higher risks of HIV-1 drug resistance (HIVDR) to antiretrovirals administered pre-, peri-, or post-partum. This is particularly true in SSA because of wide use of low generic-barrier drugs, recurrent stock-outs, impaired-adherence, inadequate monitoring, HIV-1 diversity and, importantly, limited pediatric highly active antiretroviral therapy (HAART) options. All these factors lead to delayed detection of HAART failure and HIVDR accumulation even beyond 80%. As the footprint of long-term HAART depends largely on the effectiveness of first-line drugs in sustaining viral suppression, establishing adequacy between pediatric HAART and DR-mutations (DRMs) would be clinically relevant. In this line, we earlier reported low- and high-HIVDR, respectively, in naive and HAART-failing children, with successful switch to second-line. From these observations, we postulated that minority DRMs in HAART-naive children might grow-up through selective drug-pressure and populate plasma in a short-frame, herein justifying the rapidly emerging DRMs we observed at failure. Although not yet clinically endorsed, pediatric minority DRMs might be more concerning in the context of PMTCT, henceforth underscoring an unmet clinical need. Coupled to previous knowledge on the detection of DRMs by next-generation sequencing (NGS), we thus hypothesized that using NGS to assess DRMs in vertically infected HAART-naive children would contribute in designing long-term HAART strategies for SSA-children.

Current pediatric HAART-regimens consist of lamivudine (3TC), abacavir (ABC), or zidovudine (AZT), associated to ritonavir-boosted lopinavir (LPV/r) or NVP. LPV/r is recommended to overcome PMTCT-resulting non-nucleoside reverse transcriptase inhibitor (NNRTI) resistance, whereas NVP matches with postnatal prophylaxis. As HAART would be reaching 1.5 million children by 2020, as high as 20% virological failure (VF) is expected, favored by high-viremia and poor adherence in children. Without optimal strategies, VF would quickly overcome HAART success, maintaining children vulnerable.

Moreover, pediatric HAART options are limited in SSA, urging the quest for a wider therapeutic portfolio. Although not yet approved for under 16 years, the CCR5 antagonist—maraviroc—might represent a suitable antiretroviral alternative for children, pending proof-of-concept towards relevant pediatric clinical trials. Particularly, there are limited evidence on the potential effectiveness of maraviroc for SSA-children in PMTCT, initial-HAART and/or following treatment-failure.

With rising concerns of minority variants on response to several classes of antiretrovirals, a genuine delineation of HIV-1 tropism, considering both minority and majority quasi-species, could rationalize maraviroc suitability for pediatric HAART-policies in SSA.

Based on these assumptions, we aimed to ascertain DRMs and HIV-1 co-receptor usage, in majority and minority viral populations, from children according to maternal PMTCT-exposure in a resource-limited setting (RLS).

2. Study design

2.1. Sampling and setting.

A comparative study was conducted in 2015 among 18 HIV-1 vertically infected Cameroonian children, all HAART-naive, stratified according to maternal antiretroviral exposure during pregnancy: control-group (11 children from mothers without antiretroviral exposure) versus case-group (7 children from mothers exposed to reverse transcriptase inhibitors [RTIs]). For each child, a plasma sample was collected to perform both Sanger- and 454 ultra-deep pyrosequencing (UDPS).

2.2. Sanger sequencing.

Protease (PR/RT Sanger sequencing was performed as previously described. Briefly, viral RNA was extracted from plasma using QIAamp Viral RNA minikit (Qagen, Milan, Italy), following manufacturer’s instructions. PR/RT-containing region was then reverse-transcribed and amplified using SuperScript One-Step for long templates reverse transcriptase polymerase chain reaction (RT-PCR) of Invitrogen kit (Foster City, CA), with an eventual second-round seminested PCR. Direct sequencing was then performed using 7 overlapping primers.

V3 loop Sanger sequencing was performed as previously described. Briefly, viral RNA containing the V3-loop region was reverse-transcribed and amplified using an RT/Taq mix, with an eventual second-round seminested PCR. Direct sequencing was then performed using 4 overlapping primers.

2.3. Amplification of PR/RT region for UDPS

Ten milliliters of viral RNA was reverse transcribed and amplified using 1-step RT-PCR system containing 25 mL reaction mix (2×), 8 mL MgSO4 (5 mmol/L), 2.8 mL H2O DNase RNase free, 1 µL forward primer (10 µmol/L), 1 µL reverse primer (10 µmol/L), 1 mL RNase Out (40 U/µL Invitrogen) and 1.2 mL RT/Taq, for a final volume of 50 µL. RT-PCR conditions were the following: 1 cycle 50°C, 30 minutes; 1 cycle 94°C, 2 minutes; 40 cycles (94°C, 30 seconds; 51°C, 30 seconds; 68°C, 2 minutes); a final extension 68°C, 10 minutes. Forward and reverse primers were respectively 5’GACAGGCTAATTATTAGGG3’ (2075–2094 bps, gag) and 5’GATAAATTTGATATGGC3’ (3555–3576 bps, pol).

Nested-mid PCR was then performed with the Fast Start HiFi PCR system (Roche Diagnostics, Mannheim, Germany) using 5 pairs of barcoded-modified forward and reverse primers for each amplicon (Table 1). Based on band’s size from eurusafe (Euroclone) agarose gel, 31.1 µL in water diluted cDNA was mixed per tube with 3.75 µL PCR buffer (10×), 0.75 µL dNTPs (12.5%), 0.75 µL forward primer (10 µmol/L), 0.75 µL primer (10 µmol/L) and 0.4 µL Taq, under the following conditions: 1 cycle 94°C, 3 minutes; 30 cycles (94°C, 30 seconds; 94°C, 2 minutes; 94°C, 2 minutes); a final extension 72°C, 7 minutes.

2.4. Amplification of V3 loop region for UDPS.

Ten microliters viral RNA were reverse transcribed and amplified using 1-step RT-PCR system containing 25 µL reaction mix (2×), 8 µL MgSO4 (5 mmol/L), 2.8 µL H2O DNase RNase free, 1 µL forward primer (10 µmol/L), 1 µL reverse primer (10 µmol/L), 1 mL RNase Out (40 U/µL Invitrogen) and 1.2 mL RT/Taq, for a final volume of 50 µL. RT-PCR conditions were the following: 1 cycle 50°C, 30 minutes; 1 cycle 94°C, 2 minutes; 40 cycles (94°C, 30 seconds; 51°C, 30 seconds; 68°C, 2 minutes); a final extension 68°C, 10 minutes. Forward and reverse primers were respectively 5’GACAGGCTAATTATTAGGG3’ (2075–2094 bps, gag) and 5’GATAAATTTGATATGGC3’ (3555–3576 bps, pol).

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2.4. Amplification of V3 loop region for UDPS.
minutes. A nested mid-PCR was then performed with the Fast Start HiFi PCR system (Roche Diagnostics, Mannheim, Germany) as previously described.[26]

2.5. Amplicon purification and UDPS reaction.

PR/RT PCR products (5 fragments of 436, 387, 468 and 488 bps) and V3 loop (one fragment of 367 bps) were purified using Agencourt AMPure PCR purification beads (Beckman Coulter, Brea, CA) and quantified with Quant-iT Picogreen double-stranded DNA assay kit (Life Technologies, Eugene, OR) on a GloMax multidetection system (Promega, Madison, WI).

Pooled purified PCR products were clonally amplified by emulsion PCR and pyro-sequenced on the 454 GS junior platform (Roche Applied Science, Mannheim Germany) as previously described.[26] Phylogenetic analyses excluded any possible sample contamination (data not shown).

2.6. Bioinformatics analyses of PR/RT and V3 sequences.

The entire PR (amino acid position: 1–99), RT (1–251) and the entire V3 loop (1–35) sequences obtained after 454-pyrosequencing were de-multiplexed and then quantified using the SFF tool Roche. Using a home-made Perl script and SHORAH package 0.5.1, sequences were manually checked for insertion or deletion in homopolymeric errors and aligned against HIV-1 consensus B. Final alignments were filtered and corrected for homopolymeric region-associated errors and aligned against HIV-1 consensus B. Final alignments were considered concordant if viral-tropism was identical from both sequencing technologies. Viral-tropism was explored according to age and CD4 count.

All statistical analyses were performed using the statistical open source environment R.v.3.1.1. P values <0.05 were considered statistically significant.

2.10. Ethical considerations.

Ethical clearance was obtained from the Cameroon National Ethics Committee (Ref.N°034/NEC/SE) proxy-informed consent was provided, unique identifiers were used for privacy and confidentiality, and a material transfer agreement was established.

3. Results

3.1. Characteristics of children analyzed.

Overall, median (interquartile range [IQR]) age, viremia, and CD4 count were 6 (4–10) years, 5.5 (4.9–6.0) log_{10} copies/mL, and 326 (282–645) cells/mm³, respectively, without any significant difference between the 2 groups (data not shown).

In the control, neither children nor their mothers had any antiretroviral exposure. Antiretroviral history of children belonging to the case-group, considered at higher risk of HIVDR, is described in Table 2.

3.2. HIV-1 subtype distribution.

HIV-1 subtyping revealed 50% CRF02_AG (9/18), 33.3% F (6/18), 11.1% CRF01_AE (2/18), and 5.6% CRF11.cpx (1/18).

3.3. HIV-1 drug resistance in the children analyzed.

PR/RT sequences were successfully obtained both through Sanger sequencing and UDPS for 17/18 children. The median

### Table 1

| Amplicon | Primers | Primer sequences for each amplicon (HXB2 nt. position) | Annealing temperature\(^1\) | Amplicon’s size, bp\(^2\) |
|----------|---------|--------------------------------------------------------|---------------------------|------------------------|
| 1        | Forward | 5’AGACAGGCTTTTTTCTAGGATC3’ (nt. 2074–2095)            | 56.5°C                    | 436                    |
|          | Reverse | 5’GAACATGCCTGACGCAGAAGTAC3’ (nt. 2509–2487)            | 57°C                      | 387                    |
| 2        | Forward | 5’GCAGCAGACATGACGATAGCTAGA3’ (nt. 2329–2353)          | 56.5°C                    | 476                    |
|          | Reverse | 5’GAATAGGTTTCAACCCGCTTTT3’ (nt. 2703–2694)            | 51°C                      | 468                    |
| 3        | Forward | 5’TAAAGCCGCAAGATGATG3’ (nt. 2583–2602)                | 56.5°C                    | 476                    |
|          | Reverse | 5’GGTCTCAGCTTTTGCAGTC3’ (nt. 3058–3037)               | 51°C                      | 468                    |
| 4        | Forward | 5’GGCGAATCAGTCAATTGGAAT3’ (nt. 2811–2830)             | 53°C                      | 488                    |
|          | Reverse | 5’GGCCTGTACTGTCCATT3’ (nt. 3278–3262)                | 53°C                      | 488                    |

\(^1\) Annealing temperature was used for each amplicon during the nested MID-PCR for 454-UDPS.

\(^2\) Amplicon’s size corresponds to the number of nucleotides for each of the generated amplicons.

2.8. HIV-1 subtyping

Subtyping was performed through phylogenetic analysis, by aligning all PR/RT Sanger-sequences in Bio-Edit compared to reference sequences of HIV-1 subtypes and circulating recombinant forms (CRFs) available at http://www.hiv.lanl.gov as previously described.[28]

2.9. Statistical analysis

HIV-1 DRMs and coreceptor usage were compared between the two PMTCT-groups. Coreceptor results by Sanger sequencing and UDPS were considered concordant if viral-tropism was identical from both sequencing technologies. Viral-tropism was determined by a home-made pearl script, respectively. Using a quantitative interpretation, viruses were considered CXCR4-tropic (X4-variants) by UDPS when \( \geq 2\% \) viral species had a false-positive rate (FPR) \( \leq 3.5\% \),[27] or by Sanger sequencing when FPR was \( <10\% \), describing the probability of classifying an R5-virus falsely as an X4-variant.[25]
UDPS coverage was of 1642 (IQR: 1269–5193) reads. In the entire covered PR/RT regions, the 2 sequencing technologies showed total concordance in variants detection, and all UDPS variants with frequencies <20% were not detected by Sanger sequencing (Table 3).

By using Sanger sequencing, all 17 children had a wild type virus. Only E138A (5.9%), an accessory polymorphism weakly selected under etravirine (ETR) and rilpivirine (RPV), was found in a child aged 8 years from the control group.

By using UDPS, 1 (aged 1 year) of 7 children (14.3%) from the case-group harbored viruses with K103N (8.3% prevalence; mutational load: 190,567 copies/mL), a nonpolymeric mutation causing high-level resistance to NVP and efavirenz (EFV). This infant was born from an RTI-treated mother (AZT + 3TC + NVP). Thus, Sanger sequencing and UDPS were performed also for the mother (ID-18613). UDPS revealed a virus harboring 2 transmitted minority DRMs (K103N), known to be associated with resistance to NNRTIs/NRTIs in those children under 5 years (also as minority species at 1% the threshold), and became significantly higher as from 5 years and above (55.6% [5/9], P = .040). As expected, X4-tropic viruses were higher with CD4 <15% (4/9 [44.4%]) versus CD4 >15% (1/6 [16.7%], P = .580); similarly for CD4 ≤200 (3/4 [75%]) versus CD4 >200 (2/11 [18.2%]; cells/mm^3, P = .077). No statistical difference was found in X4-variants between the 2 PMTCT-groups: 2 of 7 (28.6%) case group versus 3 of 8 (37.5%) control group, P = 1.000.

### 4. Discussion

Sustaining HAART success remains challenging for children in a long term, especially in a context where adherence and drug options are limited.\[2,4,5\] Thus, novel strategies are required to limit the spread of preventable HIVDR and provide alternative therapeutics with utmost potency for SSA children.\[29,30\]

In this high CRF02_AG-infected population,\[6,9,31,32\] HAART-naive children appeared with wild-type viruses at population-levels, confirming the low-level of HIVDR previously reported of this target-group.\[8,33\] Interestingly, a vertically transmitted minority DRM (K103N), known to be associated with resistance to NNRTIs used both for PMTCT and first-line HAART in SSA, was found in a PMTCT-exposed infant, thus suggesting NNRTI-sparing regimens for such children.\[7,30,34\] Discrepancy in DRMs between mother and infant would be due to sample collection later after delivery (at the moment of infant HIV diagnosis), with possible selection following prophylaxis/breastfeeding; as previously reported in similar RLS (Kyela, Tanzania).\[33\] This infant (aged 1 year), compared to the median age of the study population (6 years), suggests that circulating DRMs might have fade-up with increasing age.\[7,33\] NNRTI mutations (E138A and V179D), found in children without PMTCT-exposure, are known as polymorphisms with little or no effect on drug susceptibility or virological response.\[29\] The ability of NGS in excluding minority RTI-mutations (in all but one children) supports the safe use of NNRTIs/NRTIs in those older ages and lower CD4 cells, suggesting limited vertical

### Table 2

| Patient ID | ARV exposure | Duration | ARV exposure Period | PMTCT mothers |
|------------|--------------|----------|---------------------|---------------|
| 7171       | None         | —        | sd-NVP              | Pregnancy     |
| 10155      | None         | —        | 3TC + AZT + EFV     | Life-long HAART |
| 10351      | None         | —        | sd-NVP              | Pregnancy     |
| 10430      | None         | —        | sd-NVP              | Pregnancy     |
| 11621      | None         | —        | 3TC + D4T + NVP     | Life-long HAART |
| 12042      | None         | —        | sd-NVP              | Pregnancy     |
| 18737      | AZT          | 1 mo     | 3TC + AZT + NVP     | 1 month       |

3TC = lamivudine, ARV = antiretroviral, AZT = zidovudine, D4T = stavudine, EFV = efavirenz, HAART = highly active antiretroviral therapy, NVP = nevirapine, PMTCT = prevention of mother-to-child transmission, RTI = reverse-transcriptase inhibitor, sd-NVP = single-dose nevirapine.
| Patient ID | Read coverage (+/-Std) | PR region | RT region | Patient ID | Read coverage (+/-Std) | PR region | RT region |
|------------|------------------------|-----------|-----------|------------|------------------------|-----------|-----------|
| PNL43\(^3\) (B) | 2789 (±989) | 018E (1.72%) | None | K102Q (100.0%) | K102Q (100.0%) |
| 7949 [10 y] \(02\_AG\) | 1353 (±70) | L10I (100.0%) | K20I (100.0%) | K102Q (100.0%) | K102Q (100.0%) |
| 9470 [0 y] \(02\_AG\) | 1060 (±435) | L10I (100.0%) | M36I (100.0%) | L63I (92.9%) | L63I (99.7%) |
| 10196 [6 y] \(02\_AG\) | 2025 (±735) | L10I (100.0%) | K20I (100.0%) | M36I (100.0%) | L63I (100.0%) |
| 10232 [12 y] \(F1\) | 2161 (±920) | K20I (100.0%) | M36I (100.0%) | L63I (100.0%) | L63M (99.0%) |
| 10696 [5 y] \(02\_AG\) | 1467 (±527) | K20I (100.0%) | M36I (100.0%) | L63I (100.0%) | L63M (100.0%) |
| 10965 [6 y] \(F2\) | 1762 (±1267) | K20I (100.0%) | M36I (100.0%) | L63I (100.0%) | L63M (100.0%) |
| 11442 [12 y] \(02\_AG\) | 1520 (±721) | L10I (100.0%) | M36I (100.0%) | L63I (100.0%) | L63M (100.0%) |
| 11538 [12 y] \(F2\) | 5287 (±1678) | L10I (100.0%) | M36I (100.0%) | L63I (100.0%) | L63M (100.0%) |
| 12227 [3 y] \(11.0\) | 1060 (±435) | L10I (100.0%) | M36I (100.0%) | L63I (92.9%) | L63I (99.7%) |
| \(\text{continued}\) | | | | | | |
Table 3
(continued)

| Patient ID [age] (subtype) | PR region | RT region | Patient ID [age] (subtype) | PR region | RT region |
|---------------------------|-----------|-----------|---------------------------|-----------|-----------|
|                          | Reads coverage (+/- Std) | UDPS | Sanger |                          | Reads coverage (+/- Std) | UDPS | Sanger |
|                          |           |           |               |           |           |               |           |           |
| UDPS                     | Sanger    |         | UDPS         | Sanger    |         |
|                          |           |           |               |           |           |
| 12481 [8 y] (02_AG)      | 1269 (+474) | L10V (77.1%) | L10V | 11269 (±474) | L10V | K219N (30.1%) |
|                          |           | L89M (100.0%) | L89M |           | L89M | V181C (96.7%) |
|                          |           |           |               |           |           |
| 12481 [8 y] (02_AG)      |           | L63V (5.3%) | L89M | 1269 (±474) | L10V | None |
|                          |           | L89M (100.0%) |     |           | K210l | M36I (100.0%) |
|                          |           |           |               |           |           |
| 12481 [8 y] (02_AG)      |           | V106I (2.3%) | V106I | 11269 (±474) | V106I | None |
|                          |           | V108I (2.3%) | V108I |           | V108I | L89M (100.0%) |
|                          |           |           |               |           |           |
| 12481 [8 y] (02_AG)      |           | None | L89M | 1269 (±474) | None | None |
|                          |           | L89M (100.0%) |     |           | L89M | None |

* Analysis performed on 17 of 18 children with available sequences obtained through both Sanger sequencing and UDPS.
† PNL4.3 has been used as control plasmid.
‡ ID-18737: infant born to the mother ID-18613.
§ ID-18613: Mother of the infant ID-18737.

Median (interquartile range) coverage of UDPS dataset was 1642 (1269–5193) reads. In bold are all major DRMs from the Stanford HIVdb list (Updated March 9, 2015). Major DRMs with high-level reduced susceptibility are underlined. DRM = drug resistance mutation, Std = standard deviation, UDPS = ultra deep 454-pyrosequencing. Percentages (in brackets) represent the proportion of sequences harboring the mutation within the viral population of an individual.

Table 4
Viral-tropism according to sequencing technologies.

| Patient ID | Age range, y | CD4 cells/mm³ (%) | PVL copies/mL | Age, y | Subtype | Sanger 10% FPR | UDPS % seq ≤3.5 FPR G2P |
|------------|--------------|--------------------|---------------|--------|---------|---------------|-------------------------|
| 18737      | Children (<5 y) | 139 (3%) | 2,295,996 | 1 | 01_AE | R5 (40.6%) | R5 (0.000) |
| 10696      | 282 (8%) | 4,556,020 | 5 | 02_AG | R5 (92.2%) | R5 (0.000) |
| 10430      | 640 (8%) | 4,764,653 | 4 | 02_AG | R5 (30.1%) | R5 (0.031) |
| 12062      | 730 (16%) | 318,890 | 3 | 02_AG | R5 (56%) | R5 (0.000) |
| 7171       | 1213 (32%) | 350,062 | 2 | 02_AG | R5 (19.5%) | R5 (0.000) |
| 10351      | 1328 (40%) | 5,462,764 | 2 | F2 | R5 (28.9%) | R5 (0.000) |
| 11004      | Children (>5 y) | 5 (0%) | 310,116 | 12 | 01_AE | X4 (5.1%) | X4 (15.710) |
| 10155      | 196 (10%) | 945,993 | 10 | F2 | X4 (3.7%) | X4 (63.783) |
| 11036      | 165 (8%) | 377,711 | 6 | 02_AG | X4 (70.7%) | X4 (96.170) |
| 11538      | 347 (24%) | 31,713 | 12 | F2 | X4 (9%) | X4 (45.253) |
| 12481      | 466 (8%) | 463,191 | 8 | 02_AG | X4 (82.4%) | R5 (0.000) |
| 11442      | 502 (13%) | 832,928 | 8 | 02_AG | X4 (60.1%) | R5 (0.260) |
| 11021      | 544 (14%) | 17,619 | 6 | 02_AG | X4 (37.5%) | R5 (0.000) |
| 7949       | 586 (24%) | 62,283 | 10 | 02_AG | X4 (22.3%) | R5 (0.000) |
| 10965      | 645 (15%) | 281,713 | 6 | F2 | R5 (37.7%) | R5 (0.000) |
| 11621      | 544 (14%) | 163,458 | 24 | 01_AE | X4 (7.8%) | X4 (4.692) |

| Infant born from mother ID-18613. |
| Mother of infant ID-18737. |

In bold are highlighted the 2 children with discordant viral tropism results. FPR = false-positive rate, G2P = geno2pheno, PVL = plasma viral load, UDPS = ultra deep 454-pyrosequencing.
transmission by CXCR4-tropic viruses, and later appearance of X4-variants with chronicity, immunological impairment,[36,37] as well as a baseline FPR <60 as previously demonstrated.[18,39] Further investigations might help in establishing novel public health strategies for an eventual usage of maraviroc in children.[19,40] As current PMTCT-practice might not be an independent factor for viral-tropism (i.e., similar distribution in X4-variants irrespective of PMTCT-history), CCR5-antagonist (maraviroc) could be a useful therapeutic weapon for pediatric HAART.[15,18,40]

Of the two children showing discordant results between the two sequencing techniques, the added value of UDPS in detecting X4-tropic minority variants is in accordance with previous reports.[13,39] Interestingly, by detecting minority insertions associated with a complete discrepant result on Sanger sequencing, UDPS appears very useful in validating tropism determination for non-B subtypes.[41]

Therefore, UDPS might provide additional information in detecting DRM and viral-tropism, confirming the added value of this technology for both clinical diagnostics and management of non-B HIV-infected children.[12,22,41]

In spite of this added value of UDPS, implementing NGS is more challenging in RLS (costs, technical complexity, maintenance), suggesting the need for simpler and affordable approaches integrating minority variants (point-of-care or pragmatic sequencing).[32,43]

A potential study limitation could be the relatively small sample size, which makes the study probability relatively large. Also, in the PMTCT-exposed group, only 3 of 7 were exposed to triple ART, calling for subsequent investigations with scale-up of option B+. Moreover, HIV-1 variants were investigated only in plasma compartment, suggesting the need for exploring HIV variability in several compartments (cellular reservoirs, central nervous systems, among others) and the impact on treatment and monitoring strategies in SSA.[12,13,44–46] This study therefore provides relevant data to be used as base for further/enlarged studies.

In a nutshell, NGS could help in identifying PMTCT-exposed children harboring minority NNRTI-DRMs, therefore serving for a timely switch of treatment and limiting failure rate. NGS also reveals a possible absence of X4-variants among children below 5 years, thus suggesting possible public health approaches using maraviroc. These preliminary evidences, generated on a small sample of mainly CRF02_AG-infected individuals, merit further investigations for improved pediatric-HAART strategies in RLS.

Author contributions

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Acknowledgements

We are appreciative to our institutional staff that participated locally in the enrolment and in sample processing. We thank Domenico Di Carlo for statistical analyses.

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