Human immunodeficiency virus case detection and antiretroviral therapy enrollment among children below and above 18 months old: A comparative analysis from Cameroon

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Abstract

While pediatric human immunodeficiency virus (HIV) testing has been more focused on children below 18 months through prevention of mother to child transmission of HIV (PMTCT), the yield of this approach remains unclear comparatively to testing children above 18 months through routine provider-initiated testing and counselling (PITC). This study aimed at assessing and comparing the HIV case detection and antiretroviral therapy (ART) enrolment among children below and above 18 months of age in Cameroon. This information is required to guide the investments in HIV testing among children and adolescents.

We conducted a cross-sectional study where we invited parents visiting or receiving HIV care in 3 hospitals to have their children tested for HIV. HIV testing was done using polymerase chain reaction (PCR) and antibody rapid tests for children <18 months and those ≥18 months, respectively. We compared HIV case detection and ART initiation between the 2 subgroups of children and this using Chi-square test at 5% significant level.

A total of 4079 children aged 6 weeks to 15 years were included in the analysis. Compared with children <18 months, children group ≥18 months was 4-fold higher among those who enrolled in the study (80.3% vs 19.7%, \( P < .001 \)); 3.5-fold higher among those who tested for HIV (77.6% vs 22.4%, \( P < .001 \)); 6-fold higher among those who tested HIV+ (85.7% vs 14.3%, \( P = .24 \)), and 11-fold higher among those who enrolled on ART (91.7% vs 8.3%, \( P = .02 \)).

Our results show that 4 out of 5 children who tested HIV+ and over 90% of ART enrolled cases were children ≥18 months. Thus, while rolling out PCR HIV testing technology for neonates and infants, committing adequate and proportionate resources in antibody rapid testing for older children is a sine qua none condition to achieve an acquired immunodeficiency syndrome (AIDS)-free generation.

Abbreviations: AIDS = acquired immunodeficiency syndrome, ART = antiretroviral therapy, ASPA = active search for pediatric HIV/AIDS, bPITC = blanket provider-initiated testing and counselling, EID = early infant diagnosis, HIV = human immunodeficiency virus, PCR = polymerase chain reaction, PITC = provider-initiated testing and counselling, PLHIV = people living with HIV/AIDS, PMTCT = prevention of mother to child transmission of HIV, tPITC = targeted provider-initiated testing and counselling, WHO = World Health Organization.

Keywords: adolescent, child, human immunodeficiency virus infection, human immunodeficiency virus testing, infants, polymerase chain reaction
1. Introduction

The human immunodeficiency virus/acquired immunodeficiency syndrome (HIV/AIDS) pandemic remains a global health threat. According to the UNAIDS estimates, 37.9 million people were living with HIV/AIDS globally in 2018, including 1.7 million children (<15 years). In that same year, 770,000 persons died of HIV/AIDS in the world, including 100,000 children. Africa hosts 68% of people living with HIV/AIDS (PLHIV) worldwide and in 2018, 62% of AIDS related deaths were from this continent.[1] Despite this burden, transformative and remarkable progress has been made in the past decades with regards to the expansion of antiretroviral therapy (ART).[2–4] In 2018, there were 23,300,000 PLHIV receiving ART, representing 62% of patients in need of treatment. At global level, this treatment expansion has contributed in the reduction of HIV incidence and mortality by 16% and 33%, respectively, from 2010 to 2018.[11] However, children are not benefiting enough from this treatment expansion as there were 54% children on ART against 62% of adults as of June 2019.[5] Cameroon (West and central Africa) with an HIV prevalence of 3.6% has 540,000 PLVIH, including 43,000 children. In 2018, 14,000 people died of AIDS in this country, including 3600 children. The gap in pediatric ART is even wider in Cameroon as 24% of eligible children are on treatment against 55% of adults.[6]

Multiple factors account for current gaps in pediatric and adolescent ART coverage. However, poorly-implemented and/or ineffective case-finding strategies are major barriers to ART roll out in this sub-population.[7–9] More importantly, though achieving an AIDS-free generation requires the effective implementation of both prevention and treatment HIV services,[10] the global agenda in the past decades has been focused more but on prevention. In this regard, the prevention of mother to children transmission of HIV (PMTCT) program has received more attention and resources from donors, policymakers, practitioners.[11] Though the fourth prong of PMTCT include the provision of treatment to HIV-infected infants, this program provides services to infants only from birth up to 18 months.[12] Consequently, HIV testing and treatment services for older children (≥18 months) has received less attention in the past decades. Actually, out of the PMTCT program, older children (≥18 months) are expected to be screened for HIV in health facilities through the WHO routine provider-initiated counseling and testing (PITC).[13] Unfortunately, PITC implementation has been suboptimal across countries and this for several reasons including fear of stigma among parents/caregivers, lack of staff training, lack of HIV testing kits, poor commitment from facility leadership, and missed parental consent to test children.[14–16] These weaknesses have left many HIV infected children undiagnosed and untreated in the community to the extent that pediatric HIV has been considered as neglected disease by some authors.[17]

The target population for pediatric HIV infection are children from 0 to 15 years.[11] Thus, achieving an AIDS free generation requires investing adequate resources in HIV testing and treatment across this age group. As per WHO guidelines, children <18 months (in PMTCT) are HIV tested using polymerase chain reaction (PCR) while those older than 18 months (in PITC) are tested using antibody rapid tests.[18] While more focus and emphasis have been on PCR testing, the yield of this approach in terms of pediatric HIV case detection and ART initiation compared with antibody rapid tests remain unclear. This study intends to bridge this scientific gap and provide information needed to guide pediatric HIV testing investments. This should contribute in reducing the current gap in HIV treatment for children in Cameroon and beyond.

2. Methods

2.1. Design

This was a cross-sectional study, part of the “Active Search for Pediatric HIV/AIDS” (ASPA) project published elsewhere.[19] In the ASPA study, we invited all HIV infected parents receiving HIV care in 3 hospitals in Cameroon to have their children of unknown HIV status aged 6 weeks to 19 years tested for HIV (tPITC group). In the same hospitals, all parents/guardians who accompanied their sick children of the same age group for consultation at the outpatient departments were also counseled, and these children were invited to test for HIV irrespective of the presenting complaint (bPITC group).

2.2. Setting

The ASPA study was implemented at 3 public health facilities: Limbe Regional Hospital (LRH) in the South-West, Ndop District Hospital (NDH) in the North-West, and Abong-Mbang District Hospital (ADH) in the Eastern Region of Cameroon. These facilities provide comprehensive healthcare services, including HIV testing and treatment and were purposefully selected for inclusion of urban, semi-urban, and rural populations.

2.3. Study period and population

The ASPA study was implemented at LRH from July through December 2015, and from June through November 2016 at the 2 other sites. The study population in the tPITC group consisted of parents living with HIV/AIDS receiving care in the hospital and their children of unknown HIV status, aged 6 weeks to 19 years. Similarly, in the bPITC group, the study population consisted of parents/guardians and their sick children of the same age group who attended the hospital outpatient department for any reason. Children or parents critically ill (in vital distress) were excluded from the study. For this article the study population was limited to children.

2.4. Eligibility criteria

Were eligible to participate, children and adolescents of unknown HIV status, aged 6 weeks to 19 years, and enrolled in the study through parents in HIV care (tPITC group) or parents/guardians accompanying them to the hospital (bPITC group).

2.5. Sample size

The sample size was that of the aforementioned ASPA study reported elsewhere.[19]

2.6. Study procedures

2.6.1. Enrollment of participants and data collection. In the tPITC group, HIV-positive parents in care at the HIV treatment center (ART clinic) were counseled and invited by a trained counselor to participate in the study together with their children.
These parents were offered a testing opportunity for their biological children in either the hospital or at home (community testing). In the bPITC group, parents/guardians were also counseled and invited to have their sick children tested for HIV irrespective of the reason of consultation.

In both groups, all parents/guardians who consented to participate in the study were enrolled together with their children. Pre-tested and structured questionnaires were used by a trained data clerk to collect socio-demographic information of participants. For children, we collected information on age, sex, educational level, and HIV testing history (previous HIV testing and result).

2.7. HIV testing and ART enrollment

For children younger than 18 months of age, HIV testing was performed using DNA-PCR techniques. For children aged 18 months and above, HIV testing was performed using 2 HIV antibody rapid tests according to the Cameroon national guidelines. During the study period, the WHO test and treat policy was not yet effective at the site level. Thus, children who tested HIV positive were assessed for ART eligibility using WHO clinical staging and/or baseline biological analysis, including CD4 count. Eligible children were initiated on ART and monitored according to the Cameroon national guidelines.

2.8. Outcomes measurement

2.8.1. HIV case detection. Proportion of children/adolescents diagnosed for HIV infection.

2.8.2. ART enrollment. Proportion of HIV+ children/adolescents enrolled on ART.

2.9. Data management and analysis

Anonymous data from the questionnaires were entered into a database and analyzed using STATA 2013 (College Station, TX: StataCorp LP). For the purpose of this article, we excluded from the ASPA dataset, all parents as well as children 15 years or above. Only children <15 years were included in this analysis because pediatric HIV age ranges from 0 to 14 years. The study outcomes were determined by computing and comparing the proportions using Chi-square or Fisher exact tests at 5% significant level.

2.10. Ethical considerations

Participation in the study was voluntary for both parents and children. Only parents who consented were enrolled and assent was requested from children above 11 years of age. Consent from parents was obtained via signed written consent form. Likewise, assent for children over the age of 11 years was obtained through a signed written assent form. The ASPA study received ethical approval from the Cameroon National Ethics Committee, the Ludwig-Maximilians-Universität, Munich (Germany) and the Albert Einstein College of Medicine (NY, USA). The ASPA study was also registered at clinicaltrial.gov (NCT03024762).119 This study was also approved by the Cameroon Ministry of Public Health.

3. Results

A total of 4719 children and adolescents were enrolled in the ASPA study. Six hundred forty adolescents who were 15 years of age or above were excluded from the dataset. Among children/adolescents included in this analysis (n=4079), the age ranged from 6 weeks to 14 years. 51.8% were men, 49.6% had primary school level, 71.1% enrolled in the study through their mothers, and only 24.8% had previously tested for HIV (Table 1).

The age disaggregation shows that 19.7% and 80.3% of enrolled children/adolescents were <18 and ≥18 months of age, respectively. These proportions correspond to children/adolescents eligible for PCR HIV testing (<18 months) and HIV serology (≥18 months), respectively. Overall, 3156 (77.4%) children/adolescents were tested for HIV. Of those, 22.4% and 77.6% (P < .001) were <18 months and ≥18 months, respectively. Among children/adolescents who tested for HIV, 98.1% (3095) received their results. Among those who did not receive their HIV test result (n=61), 90.2% and 9.8% (P < .001) were children/adolescents <18 months and ≥18 months, respectively. Among children/adolescents who tested for HIV, 63 (2.0%) were found HIV+. Of those, 9 (14.3%) and 54 (85.7%) (P = .24) were children <18 and ≥18 months, respectively. Among children/adolescents who tested HIV+, 48 (76.1%) were enrolled on ART. Children/adolescents <18 months and those ≥18 months contributed by 8.3% and 91.7% (P = .02), respectively to the total number of children/adolescents who enrolled on ART (Table 2).

4. Discussion

According to WHO, “efforts in the global research agenda on pediatric HIV must be focused on generating targeted evidence that improves HIV programme implementation through a better understanding of what works for infants and children.”20 Our study is a response to this need.

We found that the HIV testing uptake was statistically higher among children ≥18 months compared with those <18 months (77.6% vs 22.4%, P < .001). Indeed, among children enrolled in the study, those ≥18 months of age were 3-fold more likely to be tested for HIV compared with those <18 months. This was due to the fact that during the study period, the HIV antibody rapid tests...
were readily available in the laboratory, unlike the DBS-PCR testing kits that were affected by the recurrent stock outs during that same period. The stock outs of PCR testing commodities had been previously reported as factors affecting the early infant diagnosis (EID) of HIV among neonates in sub-Saharan Africa countries.\cite{21,22} The use of point-of-care (PoC) HIV testing technology could address this gap and improves the turn-around time of EID results as well as the early ART initiation among HIV infected neonates.\cite{23-25}

Among children who tested for HIV, 98.1% received their results. Of those who did not receive results, the proportion of children <18 months were significantly higher compared with children ≥18 months (90.2% vs 9.8%, \(P < .001\)). This was mainly due to the long turned around time of EID PCR testing for children <18 months and the indeterminate HIV results for those ≥18 months. The use of PoC technology as indicated above and the availability of 3rd HIV rapid test (tie-breaker) in the laboratory should improve this outcome.

Among children who tested for HIV (\(n = 3156\)), 63 (2%) tested HIV+. Among those, children ≥18 months were predominantly represented compared with children <18 months (85.7% vs 14.3%, \(P = .24\)). Though this difference was not statistically significant, the HIV positivity among children ≥18 months was 6-fold higher compared with that of children <18 months. This suggests that a greater proportion of children living with HIV/AIDS are found among children ≥18 months. According to UNAIDS estimates, our finding is in line with the proportion of children <18 months living with HIV/AIDS in Cameroon and elsewhere.\cite{26} Therefore, adequate resources should be invested in testing children ≥18 months using HIV rapid tests. Among children tested HIV+ (\(n = 63\)), 76.1% were enrolled on ART. Of those, the proportion of children ≥18 months was significantly higher compared with those <18 months (91.7% vs 8.3%, \(P = .02\)). Actually, the contribution of children ≥18 months in the pediatric ART cohort was 11-fold higher compared with children <18 months. This further demonstrates the need of adequate investments in case finding strategies among the subpopulation of children ≥18 months in Cameroon and probably beyond.

### 4.1. Policy implications

Our results show that children ≥18 months have the highest contribution with regards to pediatric HIV case load and ART enrollment. This indicates the role the rapid HIV test could play in bridging the current gap in pediatric HIV care. That notwithstanding, HIV testing for older children is still limited in most high burden pediatric HIV countries. In these countries, the most common entry point for HIV testing of children ≥18 months is the outpatient pediatric consultations or inpatient pediatric wards using the WHO recommended PITC.\cite{13}

However, as indicated above PITC implementation had remained suboptimal across countries. As a result, only approximately 10% and 15% of HIV-infected young (15–24 years) men and women, respectively, in Sub-Saharan Africa are aware of their HIV status.\cite{27} This is a clear indication that many children ≥18 months are not adequately tested for HIV despite the higher HIV seropositivity among them as indicated by this study. This finding urges for a paradigm shift to give HIV testing for older children the adequate and proportionate attention and resources. This is a sine qua non condition to achieve the AIDS-free generation in high burden pediatric HIV countries.

We believe that in addition to all the barriers to the uptake of PITC reported to date\cite{28,29} the lack of adequate indicators to monitor the implementation of this strategy at facility level is a significant contributing factor to the current low uptake of PITC at facility level. Indeed, to monitor the performance of health facilities regarding HIV testing of their clients, it is important for these facilities to report both the number of outpatient consultations (denominator) and the number of patients tested for HIV (numerator). However, unlike the numerator, the denominator is most often not captured in the monthly summary HIV testing reports. This results to the lack of routine monitoring of health facilities performance with regards to HIV testing strategies, including PITC. On the other hand, other entry points for testing of older children (≥18 months) had not been adequately used by care providers for pediatric HIV case finding across countries in sub-Saharan Africa, including Cameroon. For example, though recommended by WHO since 2010\cite{10} the uptake of index case HIV testing, using HIV infected parents in

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**Table 2**

**Contribution of children’s age in HIV testing, case detection, and ART enrollment.**

| Outcome | Total n (col %) | Age <18 months n (row%) | Age ≥18 months n (row%) | \(P\) value* |
|---------|----------------|-------------------------|------------------------|----------|
| Children eligible for rapid antibody test | | | | |
| No | 805 (19.7) | 805 (19.7) | 0 | N/A |
| Yes | 3274 (80.3) | | 3274 (80.3) | |
| Children tested | | | | |
| No | 923 (22.6) | 99 (10.7) | 824 (89.3) | <.001 |
| Yes | 3156 (77.4) | 706 (22.4) | 2450 (77.6) | |
| Results received | | | | |
| No | 61 (1.9) | 55 (90.2) | 6 (9.8) | <.001 |
| Yes | 3095 (98.1) | 651 (21.0) | 2444 (79.0) | |
| Children’s HIV results | | | | |
| Negative | 3032 (98.0) | 642 (21.2) | 2390 (78.8) | .24 |
| Positive | 63 (2.0) | 9 (14.3) | 54 (85.7) | |
| Children enrolled on ART | | | | |
| No | 15 (23.8) | 5 (33.3) | 10 (66.7) | .02 |
| Yes | 48 (76.1) | 4 (8.3) | 44 (91.7) | |

\(\text{ART} =\) antiretroviral therapy, \(\text{HIV} =\) human immunodeficiency virus.

* \(P\) value = Chi-square or Fisher exact test.

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contact with health facilities (e.g., ART clinics) remains low in these countries. Recent studies from Kenya,[31] Malawi,[32] and Cameroon[19,33,34] have demonstrated that index case testing is a high-yield strategy for pediatric HIV case finding. Thus, in addition of monitoring the denominator of clients visiting the health facilities, the adequate implementation of index case testing among children ≥18 months should also contribute significantly to improving ART expansion among children.

4.2. Strengths and limitations
To the best of our knowledge, this is the first analysis from primary data examining the contribution of the PCR HIV testing technology against the rapid HIV antibody tests in pediatric HIV case detection and ART enrolment. Thus, our study has filled an important scientific gap in ART expansion among children. However, our study is limited by the fact that the sites were not randomly selected and therefore, our results may have been affected by selection bias. That notwithstanding, the external validity of our study is stronger as it was conducted in 3 hospitals from 3 different geographic locations covering urban, semi-urban, and rural populations in Cameroon. Hence, this study has provided new evidence that should guide the investments of the already scarce HIV funding in achieving greater impact in resource-limited settings.

4.3. Conclusions and recommendations
This study shows that 4 out of 5 children found HIV+ and over 90% of those enrolled on ART were children ≥18 months. Therefore, while rolling out the POC HIV testing technology for children below 18 months, committing adequate and proportionate resources in testing children above 18 months should be a smart investment. Translating this evidence into practice would require tweaking the data collection tools to ensure the routine monitoring of HIV testing performance of health facilities, but also expanding HIV testing to all high-yield entry points. These adjustments are necessary to reducing the current gap in pediatric HIV treatment, and to achieve the AIDS-free generation in Cameroon and beyond.

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