

1 **Lifetime changes in CD4 count, viral load suppression and adherence among**
2 **adolescents living with HIV in urban Peru**

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25 **Keywords:** Peru; adolescent health; HIV; adherence; viral load suppression; youth

26

27 **ABSTRACT**

28 Introduction

29 Viral load suppression and adherence to combined antiretroviral therapy (cART) have been
30 shown to be lower in adolescents than in other age groups; however, this relationship has not
31 been documented longitudinally from childhood to adolescence and has rarely been examined
32 outside of high-resource settings and sub-Saharan Africa. To address this knowledge gap, we
33 quantified longitudinal changes in CD4 cell count, viral load suppression, and cART adherence
34 in adolescents living with HIV in urban, Peru.

35

36 Methods

37 We conducted a retrospective chart review among adolescents ages 10-18 years on cART and
38 receiving care at a large, public sector pediatric hospital as of December 2015. We abstracted
39 clinical notes indicating nonadherence and viral load and CD4 counts from childhood to
40 adolescence. We modeled the association between age and each outcome with restricted cubic
41 splines accounting for multiple observations per person, and graphed study outcomes by age.

42

43 Results

44 A median of 7.7 years (25th percentile=4.9, 75th percentile=10.2) of follow up were observed for
45 128 adolescents. Nearly 70% of patients had at least one nonadherence note and the
46 proportion with nonadherence increased log-linearly with age ($p < 0.0001$). The peak proportion
47 with viral load suppression was 84% (95% CI: 79, 88) at age 13, which dropped to 67% (95%
48 CI: 47, 83) by age 18. Mean CD4 count decreased at age 13, dropping from 723 cells/mm³
49 (95% CI: 666, 784) to 429 cells/mm³ (95% CI: 356, 517) by age 18.

50

51 Conclusion

52 This is the first report from Latin America to examine longitudinal changes in HIV outcomes from
53 childhood into adolescence. Consistent with the limited evidence from other settings, decreases
54 in viral load suppression and mean CD4 count occurred in early adolescence in tandem with
55 increases in nonadherence. Adolescent-friendly cART adherence support interventions to target
56 this critical period are urgently needed.

57

58

59 **Introduction**

60 Combined antiretroviral therapy (cART) has greatly improved survival for infants and children
61 perinatally infected with HIV. As a result, adolescents make up a growing portion of the global
62 HIV burden with an estimated 1.8 million adolescents living with HIV (ALHIV) between ages 10
63 and 19 years in 2017.[1] Lower viral load suppression rates have been observed in children and
64 adolescents under 15 years than in adults[2] and uptake and adherence to cART is reported to
65 be lower in adolescents than other age groups.[3–6] Furthermore, HIV-related deaths among
66 adolescents have tripled over the last two decades, occurring primarily in those perinatally
67 infected.[7]

68 Little is known about longitudinal changes in HIV outcomes across the lifespan, particularly
69 during childhood and adolescence—a period of dramatic physical growth and cognitive
70 development. Available studies on the long-term outcomes of perinatally infected adolescents
71 have short follow-up periods and are largely from high resource settings or Sub Saharan Africa,
72 [8–11] with limited data from cohorts in Latin America.[12,13] The objective of this study was to
73 examine changes in absolute CD4 count, viral load suppression, and adherence from childhood
74 to adolescence among patients on cART in Lima, Peru.

75

76 **Methods**

77 **Study design**

78 We conducted a retrospective chart review of ALHIV ages 10 to 18 years receiving care at the
79 Instituto Nacional de Salud del Niño (National Institute for Child Health, henceforth INSN) in
80 Lima, Peru. INSN is a national public sector referral hospital for pediatric care and hosts the
81 largest HIV treatment clinic for children and adolescents in the country. In 2004, cART became
82 widely available in Peru due to the expansion of free, universal access to HIV care and
83 treatment.[14] The earliest guidance for treatment of HIV in children and adolescents in 2003

84 recommended two nucleoside reverse transcriptase inhibitors (NRTIs) (zidovudine and
85 lamivudine) and one protease inhibitor (nelfinavir).[15] This guidance was updated in 2013 to
86 recommend two NRTIs and one non-nucleoside reverse transcriptase inhibitor (nevirapine or
87 efaviranez).[16] Between June 2015 and April 2016, trained study personnel abstracted
88 demographic data and cART treatment history, including longitudinal CD4 measures, viral load
89 counts, and clinical notes describing nonadherence after cART initiation from paper clinical
90 charts for 132 adolescents. All adolescents were alive and on cART at the time of the chart
91 review.

92

93 **Study outcomes**

94 Absolute CD4⁺ lymphocyte monitoring was conducted every three months and viral load
95 monitoring was conducted every six months per national guidelines.[15,16] CD4 count and viral
96 load and measurements <6 months after cART initiation were excluded to allow values to
97 stabilize post-treatment initiation. This led to the exclusion of four adolescents from the analysis
98 who were on cART for <6 months at the time of the chart review. CD4 counts were additionally
99 restricted to ages 5 to 18 years because of the tendency for CD4 counts to be higher at young
100 ages due to age-related immune development.[17] Adherence and viral load were restricted to
101 the same age-years for comparability. Due to varying viral load lower limits of detection over the
102 follow-up period, we defined viral load suppression as <400 copies/ml, the least sensitive
103 threshold. Providers routinely assessed nonadherence using clinical judgement during medical
104 encounters; therefore, the absence of clinician-documented nonadherence at a given encounter
105 was presumed to indicate adequate adherence at that time.

106

107 **Statistical analysis**

108 We conducted longitudinal analyses of the three study outcomes using generalized estimating
109 equations and graphed the mean CD4 count and predicted probabilities of viral load
110 suppression and nonadherence by age-month.

111 In generalized estimating equations for CD4 count we used a normal response and log link; for
112 viral load suppression and nonadherence we used a binary response and logit link. We applied
113 an autoregressive correlation matrix to account for multiple observations per patient.[18,19] The
114 autoregressive correlation matrix assumes measurements closest in time are most correlated
115 and that this correlation decreases exponentially as measurements become more distant. To
116 examine the possible non-linear association between age and each outcome nonparametrically,
117 we used restricted cubic splines. We fit multiple models with varying numbers of knots for each
118 outcome and chose the best model by assessing the quasi-likelihood under the Independence
119 Criterion (QIC), where smaller values indicate better model fit.[20] For the outcomes of CD4
120 count and viral load suppression, modeling age with seven and four knots, respectively, fit the
121 data best. For the outcome of nonadherence, modeling age as linear fit the data best. We tested
122 for a relationship between age and each outcome using the likelihood ratio test. To examine
123 whether the overall association between age and viral suppression was driven by an increasing
124 probability of viral suppression at younger ages, versus a declining probability of suppression
125 during adolescence, we tested for an association between older age (16-18 years) versus
126 younger age (10-12 years) in the subset of measurements taken from 10 to 18 years of age.
127 Analyses were conducted in SAS version 9.4 (Cary, NC). This study was reviewed and
128 approved by research ethics committees at INSN, Lima, Peru and Harvard Medical School,
129 Boston, USA. Research ethics committees deemed that patient consent was not required for the
130 retrospective chart review.

131

132 **Results**

133 **Cohort characteristics**

134 Of 132 adolescents, 128 (97.0%) were on cART for ≥ 6 months at the time of chart review and
135 were included in the analysis. The median age at the time of the chart review was 14.6 years
136 (25th percentile=12.1, 75th percentile=16.6) with a median follow-up time after six months of
137 cART of 7.5 years (25th percentile=4.8, 75th percentile=10.0). The median age of cART initiation
138 was 5.7 years (25th percentile=3.8, 75th percentile=9.4) (Table 1). Because data were restricted
139 to the period that patients were age five or greater and on cART, the earliest CD4 counts, viral
140 load measurements, and nonadherence notes were recorded in 2004 (i.e. the year in which the
141 oldest patients included in the chart review were age five and had access to cART). The latest
142 CD4 counts, viral load measurements, and nonadherence notes were recorded in 2015 (i.e. the
143 year of the chart review).

144

145 **CD4 count**

146 A total of 2,449 CD4 counts after ≥ 6 months of cART were observed in 128 patients. Patients
147 had a median of 17 CD4 counts (25th percentile=11, 75th percentile=27) over the follow up
148 period and the median interval between measurements was 3.0 months (25th percentile=3.0
149 months, 75th percentile=4.1 months). We observed a statistically significant non-linear
150 relationship between age and CD4 count ($p=0.02$, Figure 1). The mean CD4 count trended
151 downward during childhood and decreased at a faster rate after approximately 12.8 years of
152 age (154 months) from a predicted mean CD4 count of 723 cells/mm³ (95% CI: 666, 784) to 429
153 cells/mm³ (95% CI: 356, 517) at 18 years of age (216 months).

154

155 **Viral load suppression**

156 Over the follow up period, patients had a median of 12 viral load measurements after having
157 been on cART for ≥ 6 months (25th percentile=9, 75th percentile=16.5). The median interval
158 between assessments was 6.0 months (25th percentile=5.8 months, 75th percentile=6.7 months).
159 Of 1,531 viral loads among 128 patients, 1,115 (73%) were suppressed (<400 copies/ml). At
160 least one suppressed viral load was observed in 123 (96%) patients. We observed a non-linear
161 relationship between age and viral load suppression ($p < 0.0001$) in which the predicted
162 proportion of patients with viral suppression increased during childhood and early adolescence
163 and declined thereafter (Figure 1). The predicted proportion of patients with viral load
164 suppression peaked at age 12.7 years (152 months), with 84% (95% CI: 79, 88) virally
165 suppressed. By 18 years of age (216 months), suppression rates decreased to 67% (95% CI:
166 47, 83). When we examined the association between age and viral load suppression during
167 adolescence, we found a significantly lower predicted probability ($p = 0.02$) of viral load
168 suppression in older adolescence (i.e., 16 to 18 years of age) than younger adolescence (i.e.,
169 10 to 12 years of age).

170

171 **Nonadherence**

172 A total of 328 nonadherence notes were recorded over 10,788 follow-up months. Eighty-six
173 (67%) patients had at least one nonadherence note recorded in their chart. The median number
174 of nonadherence notes per patient was one (25th percentile=0, 75th percentile=4). The predicted
175 proportion with a nonadherence note increased with age in a log-linear fashion ($p < 0.0001$,
176 Figure 1). At five years of age (60 months), the predicted proportion of patients with a
177 nonadherence note was 0.8% (95% CI: 0.5, 1.3) which increased to 11% (95% CI: 7, 15) by 18
178 years of age (216 months).

179

180 **Discussion**

181 We report lifetime changes of CD4 count, viral load suppression and nonadherence from
182 childhood into adolescence in a cohort of adolescent patients on cART at an urban pediatric
183 hospital in Lima, Peru. We found that dramatic declines in CD4 count and viral load suppression
184 were observed after approximately age 13 and that nonadherence increased with age. These
185 findings indicate that targeted interventions to improve clinical outcomes and support cART
186 adherence are needed early in adolescence in this population.

187 Adherence to cART is critical to maintaining viral load suppression.[21,22] In our study,
188 decreases in viral load suppression and mean absolute CD4 count with age are likely due to
189 increases in nonadherence, which changed in tandem with these outcomes. Other work in this
190 population supports that adherence suffers during adolescence and that the mechanisms
191 through which nonadherence occur are amenable to intervention. Through a health behaviors
192 survey implemented in this study population, we found self-reported nonadherence was greatest
193 in the 13 to 15 year age group, with 82% of adolescents missing ≥ 3 doses in a 30 day
194 period.[23] In psychosocial support groups, adolescents described barriers to adherence at the
195 individual- and family/caregiver-levels, providing the ideal opportunity to deliver support or
196 education interventions.[24] Over the last decade, many adherence interventions leveraging
197 technology have been studied.[25,26] Some of this work has been conducted exclusively in
198 adolescents[27,28] and may be particularly acceptable in populations with ready access and
199 experience using mobile devices, like adolescents in Lima.[29] As adolescents mature and
200 responsibility for their care is transferred from caregiver to the adolescent, health education on
201 living with HIV becomes increasingly important. Education through non-traditional mediums
202 such as social media and music have been explored in young persons,[30–32] however further
203 work from a variety of settings is needed. At 18, most adolescents must transition from pediatric
204 to adult HIV care, a period associated with poor outcomes.[33] In our study, we observed

205 declines in viral load suppression and adherence even before this transition. In anticipation of
206 this change, education and skills building on how to stay healthy into adulthood should be
207 provided early on in adolescence.

208 Adolescents continue to be underrepresented in HIV research and policy, despite calls to
209 prioritize this group.[34,35] Systematic reviews assessing interventions for ALHIV have found
210 that most studies are conducted in adults or in high-resource settings.[36–38] When studies
211 include ALHIV from low-resource settings, they are primarily from Sub Saharan Africa,[38]
212 limiting the generalizability of findings outside settings with generalized HIV epidemics.[38] A
213 2015 systematic review of interventions to improve linkage and retention in care among ALHIV
214 did not identify any studies from Latin America,[27] and only 10% of studies in a review
215 investigating adherence among adolescents were conducted in Latin America—all of which
216 were from Brazil.[38] In Peru, adolescents ages 10 to 19 years are largely perinatally infected,
217 while new cases of HIV in youth ages 15 to 24 years are concentrated in men who have sex
218 with men and transgender women.[39] These key differences in risk populations demand
219 tailored interventions.

220 Limitations include that our study population is a survivor cohort. A retrospective chart review of
221 all children <18 years receiving HIV care at INSN from 2003 to 2012 reported mortality rates
222 were under 9%, indicating survivor bias in our study is likely small.[40] Additionally, data on
223 nonadherence was based on the presence of a clinician note documented in the chart. In
224 modeling these data, we assumed that the absence of a nonadherence note signified
225 adherence, thus our estimate of the predicted proportion of patients with a nonadherence note
226 may be of greater magnitude. Despite these limitations, the nonadherence results triangulate
227 with those of CD4 count and viral load, as would be expected. Together, these findings provide
228 evidence that adolescence is an important period for interventions aimed at improving clinical
229 outcomes, of which adherence support is one strategy.

230

231 **Conclusions**

232 Studies assessing lifetime HIV outcomes from childhood to adolescence are limited, especially
233 in Latin America. Decreases in mean CD4 count and viral load suppression were observed
234 during early adolescence and occurred in accordance with provider-reported nonadherence.
235 Research on effective, tailored interventions aimed at improving clinical outcomes and
236 adherence during adolescence are needed in this population.

237

238 **Competing interests**

239 The authors report no conflicts of interest.

240

241 **Authors' contributions**

242 LK, MFF and MM conceived of the study. MFF designed the study. AR and MW collected the

243 data and MM supervised data collection. CAR conducted the analysis and wrote the first draft.

244 AR, CAR, KP, LK, MFF and MW interpreted the results. All authors critically reviewed the

245 manuscript and approved the manuscript for submission.

246

247 **Acknowledgements**

248 This study was support by funding from the William F. Milton Fund of Harvard University.

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Table 1. Clinical characteristics among adolescents living with HIV in urban Peru, (N=128)

	Median (25th, 75th percentiles)^a
Male, N (%)	68 (53.1)
Age at chart review (years)	14.6 (12.1, 16.6)
Age at cART initiation (years)	5.7 (3.8, 9.4)
Follow-up time on ≥6 months of cART (years)	7.5 (4.8, 10.0)
Time on cART (years)	8.2 (5.4, 10.7)
Viral loads measurements	
Number of viral loads measurements per adolescent after ≥6 months of cART	12 (9, 16.5)
Interval between viral load measurements (days)	182 (175, 203)
CD4 count measurements	
Number of CD4 count measurements per adolescent after ≥6 months of cART	17 (11, 27)
Interval between CD4 counts measurements (days)	91 (91, 126)
Nonadherence	
Adolescents with a nonadherence note, N (%)	86 (67.2)
Number of nonadherence notes per adolescent	1 (0, 4)

^a Unless specified otherwise

Figure 1. Lifetime changes in CD4 count (A), viral load suppression (B), and nonadherence (C) from age 5 (60 months) to 18 years (216 months) among adolescents living with HIV on cART in Lima, Peru

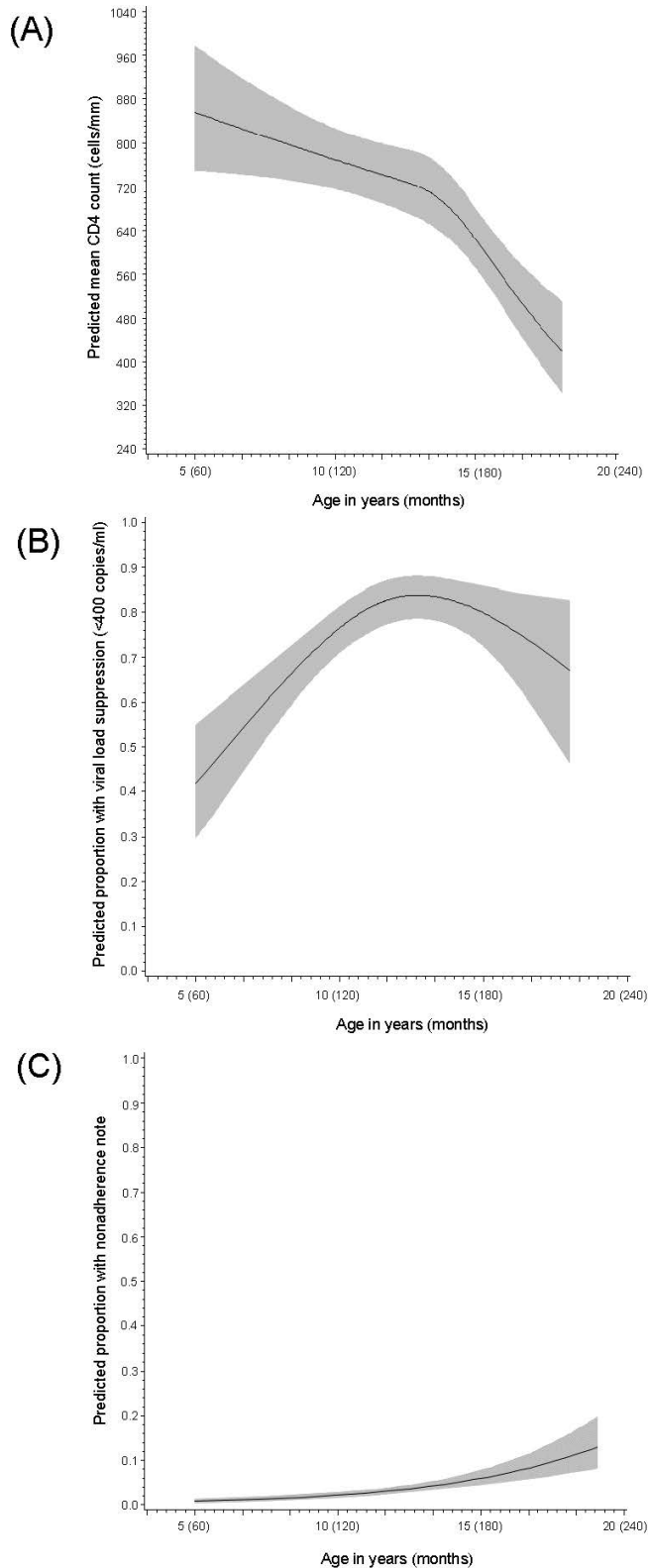


FIGURE CAPTIONS

Figure 1. (A) Mean CD4 count (cell/mm³), by age in years and months; (B) Proportion of adolescents with viral load suppression (<400 copies/ml), by age in years and months; (C) Proportion of adolescents with a nonadherence note, by age in years and months