CD4 Deficit and Tuberculosis Risk Persist With Delayed Antiretroviral Therapy: 5-Year Data From CIPRA HT-001

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CD4 deficit and tuberculosis risk persist with delayed antiretroviral therapy: 5-year data from CIPRA HT-001


**SETTING:** Port-au-Prince, Haiti.

**OBJECTIVE:** To determine long-term effects of early vs. delayed initiation of antiretroviral therapy (ART) on immune recovery and tuberculosis (TB) risk in human immunodeficiency virus (HIV) infected individuals.

**DESIGN:** Open-label randomized controlled trial of immediate ART in HIV-infected adults with CD4 counts between 200 and 350 cells/mm³ vs. deferring ART until the CD4 count was <200 cells/mm³. The primary comparisons were CD4 counts over time and risk for incident TB, with 5 years of follow-up.

**RESULTS:** A total of 816 participants were enrolled, with 408 in each treatment arm. The early treatment group started ART within 2 weeks, while the deferred treatment group started ART a median of 1.3 years after enrollment. After 5 years, the mean CD4 count in the early treatment group was significantly higher than in the deferred treatment group (496 cells/mm³, 95% confidence interval [CI] 477–515 vs. 373 cells/mm³, 95%CI 357–389; *P* < 0.0001). TB risk was higher in the deferred treatment group (unadjusted HR 2.41, 95%CI 1.56–3.74; *P* < 0.0001) and strongly correlated with lower CD4 counts in time-dependent multivariate analysis.

**CONCLUSION:** Delays in ART initiation for HIV-infected adults with CD4 counts of 200–350 cells/mm³ can result in long-term immune dysfunction and persistent increased risk for TB.

**KEY WORDS:** human immunodeficiency virus; TB incidence; CD4 lymphocyte count; HIV-TB; ART

FOR HUMAN IMMUNODEFICIENCY VIRUS (HIV) infected patients with CD4 counts <350 cells/mm³, early initiation of antiretroviral therapy (ART) improves survival, reduces the risk of HIV transmission, and prevents opportunistic infections.¹⁻⁴ The randomized Comprehensive Program for Research in AIDS (CIPRA) HT-001 trial in Haiti demonstrated that starting ART with a CD4 count between 200 and 350 cells/mm³ reduced mortality by 75% compared to delaying ART until the CD4 count was <200 cells/mm³ or an acquired immune-deficiency syndrome (AIDS) defining illness occurred, with the majority of deaths occurring before initiation of ART.⁵ Based on accumulated evidence from this trial and others, the World Health Organization (WHO) revised their guidelines in 2010, raising the CD4 threshold for ART initiation to ≤350 cells/mm³, and again in 2013 to ≤500 cells/mm³.³,⁶,⁷

It is not known if and for how long after ART initiation patients who delay therapy have an immune deficit and increased risk of tuberculosis (TB). Observational studies suggest that individuals with low CD4 counts at ART initiation have impaired immunologic recovery on therapy, and low CD4 counts are the strongest mediator of TB risk.⁸⁻¹² To determine the effect of early ART on CD4 recovery and TB risk, we conducted a study of the randomized controlled CIPRA HT-001 cohort of HIV-infected adults who received early vs. deferred ART. The primary comparisons were CD4 count and risk of incident TB over 5 years of follow-up.

**STUDY POPULATION AND METHODS**

**Design and setting**

CIPRA HT-001 was an open-label randomized controlled trial conducted at the Center of the Haitian Group for the Study of Kaposi’s Sarcoma and
Opportunistic Infections (Gheskio) in Port-au-Prince, Haiti, and was approved by the institutional review boards at Gheskio and Weill Cornell Medical College, New York, NY, USA. All participants provided written informed consent.

Between August 2005 and July 2008, HIV-infected participants aged ≥18 years, with a CD4 count between 200 and 350 cells/mm³, and no history of an AIDS-defining illness (WHO Stage 4), and no prior ART were enrolled. Pulmonary TB was not an exclusion criterion. Participants were randomly assigned to initiate ART within 2 weeks of enrollment (early treatment group), when their CD4 count was <200 cells/mm³, or when they developed an AIDS-defining illness (deferred treatment group). In June 2009, a scheduled interim analysis showed a survival benefit for early ART, and the Data Safety and Monitoring Board (DSMB) recommended that all participants start ART with continued follow-up. The 2009 analysis and details on study design have been reported previously.

In the present study, we extended follow-up until 1 January 2012 and examined the effect of early ART on CD4 counts and incident TB for 5 years after randomization.

Study intervention
Participants in both groups were seen monthly by a clinician and received an identical package of services provided to all HIV-infected patients at Gheskio, including prophylaxis with trimethoprim-sulfamethoxazole, nutritional support and adherence counseling. Enrollees were screened for latent tuberculous infection (LTBI) and active TB using a purified protein derivative (PPD) skin test, chest X-ray (CXR) and three sputum samples for acid-fast bacilli (AFB) and mycobacterial culture. Participants with symptoms suggestive of TB were screened using CXR and three sputum samples for AFB smear and mycobacterial culture. For suspecting extra-pulmonary TB, a biopsy with AFB smear, mycobacterial culture, and histopathology was performed at the discretion of the study physician.

We used the TB case definition of the American Thoracic Society as described in previous reports. For diagnosis, the required criteria were symptoms of active TB, including fever, night sweats, weight loss, cough, dyspnea, hemoptysis, or lymphadenopathy and AFB on sputum smear, a positive culture for M. tuberculosis, or histopathological findings consistent with mycobacterial disease. For cases with negative AFB smear, culture and histopathology, we required a CXR highly suggestive of active TB and clinical response to anti-tuberculosis treatment. Cases were defined as microbiologically confirmed if AFB smear or M. tuberculosis culture was positive.

Patients with active TB were treated using directly observed therapy with daily rifampin (R), isoniazid (H), ethambutol (E), and pyrazinamide (Z) for 2 months, followed by daily RH for 4 months. Patients who were retreated due to recurrence, treatment failure, or default with drug-susceptible M. tuberculosis received streptomycin plus RHEZ. Patients with drug-resistant TB were treated according to WHO guidelines.

Statistical analysis
Clinical and laboratory information were entered electronically in Haiti, managed by Frontier Science and Technology Research Foundation and exported into SAS (Statistical Analysis System, Cary, NC, USA) for analysis. Analyses were based on intention to treat.

Mean CD4 counts and 95% confidence intervals (CIs) were plotted over 24-week intervals for each treatment group, starting from randomization and separately from initiation of ART. If a participant had more than one CD4 measurement over a 24-week period, the lower value was incorporated. Multiple imputations were used for missing data. The most conservative models using weighted averages are reported. Generalized estimating equations (GEE) were used to compare mean CD4 counts and the rate of change in CD4 counts of the two treatment groups. For analysis of time to occurrence of incident TB, we excluded participants with active TB at enrollment. Primary data analysis was conducted from the time of randomization and a secondary analysis from
the time of ART initiation. The outcome of interest was TB diagnosis. Participants were censored on the date they were lost to follow-up, died, or on 1 January 2012. The probability of TB-free survival was calculated using Kaplan-Meier survival methods, and differences were evaluated using the log-rank test. Incidence rates were compared using Fisher’s exact test.

We examined previously reported predictors of TB, including time-independent variables (age at randomization, sex, education, income, PPD status, history of TB) and time-dependent variables (CD4 count, body mass index [BMI]) using univariate Cox proportional hazards regression models to estimate the risk of TB as hazard ratios (HRs) with 95%CI from the time of enrollment. Variables with \( P \leq 0.05 \) in univariate analysis were retained in multivariate models. We hypothesize that the CD4 count lies on the causal pathway between ART and TB risk; we therefore analyzed models with and without treatment group as a predictor variable. Two-sided tests were adopted for all statistical inferences.

RESULTS

Study participants

Of the 1066 subjects screened, 816 were enrolled between August 2005 and July 2008. The median age was 40 years; 470 (58%) were women, and the median CD4 count was 281 cells/mm\(^3\). Forty-three had TB at randomization, 28 (7%) in the early treatment group and 15 (4%) in the deferred treatment group (\( P = 0.042 \)). The proportion of participants with a history of TB and those with a positive PPD were balanced between the groups (Table 1).

Status at time of analysis

Data were collected until 1 January 2012. The median follow-up was 4.4 years (interquartile range [IQR] 3.5–5.0, 1531 person-years [py]) in the early treatment arm and 3.9 years (IQR 2.8–4.7, 1412 py) in the deferred treatment arm.

At the time of analysis, of 408 participants in the early treatment arm, 23 (6%) had died, 30 (7%) were lost to follow-up, and 355 (87%) remained in care. Of the 408 participants in the deferred treatment arm, 41 (10%) had died, 39 (10%) were lost to follow-up, 4 (1%) declined ART but remained in care, and 325 (80%) remained in care and on ART (Figure 1).

In the deferred treatment arm, 351 (86%) initiated ART a median of 1.3 years (IQR 1–2.1) after randomization. The median follow-up on ART was 2.5 years (IQR 2.3–3.3). After starting ART, 18 of 351 (5%) died, 9 (3%) were lost, and 324 (93%) remained in care.

### CD4 count recovery

Using GEE analysis, the mean CD4 count of the early treatment group was significantly higher over 5 years than that of the deferred treatment group (\( P < 0.0001 \)). At the time of randomization, the mean CD4 counts were the same (early treatment group, 264 cells/mm\(^3\), 95%CI 258–266; deferred treatment group, 264 cells/mm\(^3\), 95%CI 259–267). At 5 years, the mean CD4 count of the early treatment group reached 496 cells/mm\(^3\) (95%CI 476–515), while that of the deferred treatment group reached 373 cells/mm\(^3\) (95%CI 357–389) (Figure 2). Multiple imputations for missing data did not significantly alter the results of this or subsequent analyses.

Using GEE analysis, the mean CD4 count of the early treatment group was significantly higher from the time of ART initiation through 3 years of follow-up on ART (\( P < 0.0001 \)). In the early treatment group, the mean CD4 count rose from a baseline value of 262 cells/mm\(^3\) to 483 cells/mm\(^3\) (95%CI 464–503) after 3 years of ART (Figure 3). In the 351 deferred treatment group participants who started ART, the mean CD4 count rose from 186 cells/mm\(^3\) (95%CI 179–193) to
393 cells/mm$^3$ (95% CI 374–411) after 3 years of ART. There was no difference in the rate of change of CD4 counts between groups ($P = 0.47$).

**Incident tuberculosis**

Forty-three (5%) participants had active pulmonary TB at enrollment and were excluded from the TB incidence analysis. Among the 773 subjects who were TB-free at enrollment, 96 (12%) developed TB during the study, 30 in the early treatment and 66 in the deferred treatment arms ($P < 0.0001$).

Among 30 early treatment group participants with TB, 29 (97%) had pulmonary TB, one had extrapulmonary TB, and 14 (47%) were microbiologically confirmed. Of the 66 participants in the deferred treatment group with incident TB, 33 (50%) were diagnosed before starting ART and 33 (50%) after starting ART. Of these, 63 (95%) developed pulmonary TB, 3 (5%) had extrapulmonary TB, 45 (68%) were microbiologically confirmed and one had multidrug-resistant TB.

In a Kaplan-Meier survival analysis from the time of randomization (Figure 4), the probability of developing TB within 5 years was 9.5% in the early treatment group compared to 20% in the deferred treatment group ($P < 0.0001$ using log-rank test). Deferred ART carried an increased TB risk in the 5 years after enrollment; the unadjusted HR was 2.41 (95% CI 1.56–3.74).

In a Kaplan-Meier survival analysis from the time of ART initiation (Figure 5), the probability of developing active TB within 3 years was 6.2% in the early treatment group compared to 11.5% in the deferred treatment group ($P = 0.026$ using log-rank test). The unadjusted HR for deferred ART was 1.69 (95% CI 1.01–2.78).

The TB incidence rate was 2.0 cases/100 py (95% CI 1.3–2.8) in the early treatment arm vs. 5.0 cases/100 py (95% CI 3.9–6.2) in the deferred treatment arm ($P = 0.001$). In the deferred treatment group, the TB incidence rate was 6.0/100 py (95% CI 4.3–8.3) in the pre-ART and 4.2/100 py (95% CI 3–5.8) in the post-ART period. All participants with TB started anti-tuberculosis treatment. Two participants (7%) in the early treatment arm and 11 (17%) in the deferred treatment arm died while on anti-tuberculosis treatment.

**Tuberculosis risk factors**

On univariate analysis of the characteristics associated with incident TB, significant associations were found with treatment group, history of TB prior to enrollment, WHO HIV clinical stage at enrollment, CD4 count at the time of diagnosis and BMI at the time of diagnosis. In the multivariate model, WHO
HIV clinical stage at enrollment, CD4 counts at diagnosis, and BMI at diagnosis were independently associated with incident TB (Table 2, Model 1). When treatment group was included in the multivariate model, both CD4 count and treatment group were significant predictors of incident TB (Table 2, Model 2). Including both variables reduced the HR for treatment group, suggesting that differences in CD4 counts mediate some, but not all, of the differential TB risk between randomization groups.

PPD status at enrollment was not significantly associated with incident TB (HR 1.27, 95%CI 0.83–1.95, P = 0.28). Of note, patients with a positive PPD received INH prophylaxis, and PPD status at enrollment was not a further effect modifier of the association between randomization group and TB risk.

**DISCUSSION**

In this randomized controlled trial, HIV-infected adults with a CD4 count of 200–350 cells/mm³ who...
Figure 4  Kaplan-Meier estimates of the probability of TB-free survival from the time of study randomization. TB tuberculosis; ART antiretroviral therapy.

Figure 5  Kaplan-Meier estimates of the probability of TB-free survival from the time of ART initiation. TB tuberculosis; ART antiretroviral therapy.
delayed ART initiation for an average of 1.3 years had a lower mean CD4 count over 5 years of follow-up than participants who started ART immediately. The difference between the mean CD4 counts of the treatment groups was 123 cells/mm³ after 5 years. The TB incidence was 2.5 times higher in the deferred treatment group and correlated with lower CD4 counts. The study demonstrates that short delays in ART initiation can result in long-term immune dysfunction and increased risk of TB.

Our findings are consistent with observational cohorts in which the CD4 count at ART initiation predicts the degree of immune recovery. More than 75% of participants in our early treatment group achieved a CD4 count >500 cells/mm³ by 5 years. When CD4 counts are maintained above 500 cells/mm³, mortality rates in HIV-infected individuals can approach those of the general population. Relatively short delays in ART initiation for HIV-infected adults with a CD4 count of 200–350 cells/mm³ can result in long-term immune dysfunction and increased risk of TB.

Early initiation of ART is cost-effective in developing countries. Our previous cost-effectiveness analysis of the CIPRA HT-001 trial did not account for the long-term impact of delayed ART on TB risk. The benefit of early ART will therefore likely be greater than previously estimated. In our multivariate analysis, lower CD4 counts were an independent risk factor for TB, suggesting a causal relationship consistent with data from South Africa. These studies did not report impaired immune recovery in patients with lower CD4 counts, a finding we believe underlies the elevated TB risk in our deferred treatment participants.

PPD status was not an independent risk factor for incident TB and did not modify the effect of early ART on incident TB. Participants were enrolled when their CD4 counts were <350 cells/mm³, and as a consequence already had some immune deficit. Partial anergy may explain why only 25% of the cohort was PPD-positive, when data from Haiti suggest that 60% of the population has LTBI. Furthermore, all the participants who tested PPD-positive received INH prophylaxis, which can reduce the incidence of TB by 75%,. The net effect was that PPD status was a poor predictor of incident TB in our cohort.

**CONCLUSION**

Relatively short delays in ART initiation for HIV-infected adults with a CD4 count of 200–350 cells/mm³ can result in long-term immune dysfunction and increased risk of TB.

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Conflicts of interest: none declared.
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