Discordant Treatment Responses to Combination Antiretroviral Therapy in Rwanda: A Prospective Cohort Study

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters.

| Citation |
|----------|
| Kayigamba, Felix R., Molly F. Franke, Mirjam I. Bakker, Carly A. Rodriguez, Emmanuel Bagiruwigize, Ferdinand WNM Wit, Michael L. Rich, and Maarten F. Schim van der Loeff. 2016. "Discordant Treatment Responses to Combination Antiretroviral Therapy in Rwanda: A Prospective Cohort Study." PLoS ONE 11 (7): e0159446. doi:10.1371/journal.pone.0159446. http://dx.doi.org/10.1371/journal.pone.0159446. |

| Published Version |
|--------------------|
| doi:10.1371/journal.pone.0159446 |

| Citable link |
|--------------|
| http://nrs.harvard.edu/urn-3:HUL.InstRepos:29002550 |

| Terms of Use |
|--------------|
| This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA |
Discordant Treatment Responses to Combination Antiretroviral Therapy in Rwanda: A Prospective Cohort Study

Felix R. Kayigamba1*, Molly F. Franke2,3, Mirjam I. Bakker4, Carly A. Rodriguez2, Emmanuel Bagiruwigize1, Ferdinand WNM Wit5,6, Michael L. Rich3,7, Maarten F. Schim van der Loeff5,6,8

1 INTERACT, CPCD, PO Box 2181, Kigali, Rwanda, 2 Department of Global Health and Social Medicine, Harvard Medical School, Boston, MA, United States of America, 3 Partners In Health/Inshuti Mu Buzima, Rwinkwavu, Rwanda, 4 Royal Tropical Institute, KIT Biomedical Research, Amsterdam, the Netherlands, 5 Amsterdam Institute for Global Health and Development (AIGHD), Amsterdam, the Netherlands, 6 Center for Infection and Immunity Amsterdam (CINIMA), Academic Medical Center (AMC), Amsterdam, the Netherlands, 7 Division of Global Health Equity, Brigham and Women’s Hospital, Boston, MA, United States of America, 8 Public Health Service of Amsterdam (GGD), Amsterdam, the Netherlands

* fkai-gamba@gmail.com

Abstract

Introduction

Some antiretroviral therapy naïve patients starting combination antiretroviral therapy (cART) experience a limited CD4 count rise despite virological suppression, or vice versa. We assessed the prevalence and determinants of discordant treatment responses in a Rwandan cohort.

Methods

A discordant immunological cART response was defined as an increase of <100 CD4 cells/mm³ at 12 months compared to baseline despite virological suppression (viral load [VL] <40 copies/mL). A discordant virological cART response was defined as detectable VL at 12 months with an increase in CD4 count ≥100 cells/mm³. The prevalence of, and independent predictors for these two types of discordant responses were analysed in two cohorts nested in a 12-month prospective study of cART-naïve HIV patients treated at nine rural health facilities in two regions in Rwanda.

Results

Among 382 patients with an undetectable VL at 12 months, 112 (29%) had a CD4 rise of <100 cells/mm³. Age ≥35 years and longer travel to the clinic were independent determinants of an immunological discordant response, but sex, baseline CD4 count, body mass index and WHO HIV clinical stage were not. Among 326 patients with a CD4 rise of ≥100 cells/mm³, 56 (17%) had a detectable viral load at 12 months. Male sex was associated...
with a virological discordant treatment response \((P = 0.05)\), but age, baseline CD4 count, BMI, WHO HIV clinical stage, and travel time to the clinic were not.

**Conclusions**

Discordant treatment responses were common in cART-naïve HIV patients in Rwanda. Small CD4 increases could be misinterpreted as a (virological) treatment failure and lead to unnecessary treatment changes.

**Introduction**

The aim of combination antiretroviral therapy (cART) is to suppress plasma human immunodeficiency virus (HIV) viral load (VL) to undetectable levels. The usual median time to achieve full viral suppression is about 100 days \([1,2]\). Most HIV patients, both in high-income and in resource-poor countries, also display an immunological response to treatment, measured as an increase in CD4 count.\([3–5]\) In 14–25% of patients CD4 count does not rise substantially despite successful viral suppression.\([1,6–9]\) This phenomenon has been referred to as an immunological discordant treatment response.

Studies have reported an increased incidence of AIDS events or death among those with immunological discordant responses.\([1,6,8–11]\) The mortality risk among immunological discordant responders is between that of complete responders and that of complete non-responders, \([6,8]\) thus, discordant treatment responses are regarded as suboptimal treatment outcomes.

Older age and lower baseline VL have consistently been shown to be associated with discordant response.\([1,6,7,10,12–14]\) Low adherence and lamivudine or zidovudine containing regimens were also found to be associated with a discordant response.\([6]\) Studies examining the relationship between baseline CD4 cell count and discordant response show conflicting results, with some reporting a positive association between low CD4 count and a discordant treatment response,\([1,6,15]\) and others the reverse.\([7,10]\) Most studies on discordant responses have been done in cohorts from high-income countries.

Another type of discordant treatment response is a positive immunological response despite incomplete suppression of viral replication. This type of response is was found to be associated with a history of injecting drug use, high baseline HIV VL, and poor adherence.\([6]\) Those with a discordant virological response have a higher mortality risk,\([6,8,9,11]\) and like discordant immunological responses, is regarded as a suboptimal treatment response.

In practice, routine viral load monitoring is recommended to detect treatment failure earlier and accurately;\([16]\) however, in resource-limited settings where routine virological monitoring is not available, immunological and clinical criteria are often used. As a result, patients presenting with a negative immunological response may be misclassified as having failed treatment, and unnecessarily switched to costly second-line regimens. For this reason, understanding discordant treatment responses, and the factors that influence them, is critical to optimizing cART use. We studied the frequency of discordant treatment responses in a cohort of cART-naïve HIV patients starting cART in Rwanda, and assessed determinants of discordant responses in this setting.

**Methods**

Rwanda is one of only three countries in sub-Saharan Africa with a generalised HIV epidemic where over 90% of ART eligible HIV patients are on cART.\([17]\) A dense network of clinics and
hospitals provide HIV care and treatment, free of cost. We conducted a prospective study of 610 ART-naïve HIV infected patients starting cART at nine health facilities in Rwanda. We identified two nested cohorts of patients for analyses of discordant immunological and virological response.

A detailed description of the study methods and of treatment outcomes of the full cohort has been published. In brief, patient enrollment started in June 2007 and ended in August 2008. Inclusion criteria were: (1) documented HIV infection; (2) starting cART at one of the nine selected Ministry of Health (MOH) centers in the two study regions; (3) residence in one of the study regions for at least the past one year. Patients were excluded if CD4 count was above 350 cells/mm³ at the time of cART initiation, if they were aged less than 21 years or if they had previously initiated cART (except for women who had received short-term antiretrovirals during pregnancy).

**Standard of care for cART**

cART was provided free-of-charge to all individuals who met eligibility criteria based on Rwandan Ministry of Health guidelines. In short, eligibility criteria for cART at the time of the study included: confirmed HIV seropositivity; WHO clinical stage 4 regardless of CD4 count or WHO clinical stage 1, 2, or 3 with a CD4 count of <350/mm³; and fulfillment of mandatory social conditions. Current guidelines call for consideration of regimen switches when there is suspicion of clinical, immunological or virological treatment failure, and only after careful assessment of adherence and repeat CD4 cell count and/or viral load testing. The first-line cART regimen for HIV-infected individuals consisted of either stavudine or zidovudine, plus lamivudine and nevirapine. Efavirenz replaced nevirapine in individuals who were receiving tuberculosis (TB) treatment. Co-trimoxazole was routinely prescribed to individuals with CD4 cell counts <350 cells/mm³ or World Health Organization (WHO) HIV clinical stage 3 or 4.

Data collection

A baseline clinical exam was done before cART initiation for all patients enrolled into the study. A standardized intake form was completed regarding age, sex, marital status, literacy, study site, patient travel time to the clinic, weight and height, CD4 cell count, WHO clinical stage, cART start date, antiretroviral regimen, and whether the patient was being treated for TB at the time of initiation of cART. CD4 cell count measurements were done at baseline and after 12 months, using the FACS Count system (Becton Dickinson TM, La Pont de Claix, France). Plasma VL measurement was done only after 12 months of cART using the Cobas TaqMan 48 Analyzer (Roche, Geneva, Switzerland); the threshold level for detection of VL was 40 copies/mL. During monthly clinic visits patients were examined and any diagnosed opportunistic infections were treated. No study participant was prescribed a second-line cART regimen during their first year on cART (i.e., their study follow up period). Adherence was assessed 3 months and 12 months after the start of cART, using the validated Center for Adherence Support Evaluation (CASE) adherence index. This index is a simple composite measure of self-reported ART adherence, based on three questions. The theoretical score range is from 3 to 16, however pilot data suggested that participants had difficulty distinguishing between...
two response categories: missing a dose an average of “zero times per week” and “less than once a week”. We therefore combined these response categories for a total maximum score of 15. A CASE index score of ≤15 indicates suboptimal adherence.[22]

Selection of Nested Cohorts

Of the 610 patients included in the full cohort, 35 (6%) died by the end of the 12-month observation period, 13 (2%) defaulted, 15 (3%) were transferred to other clinics, and 547 patients were retained in care at 12 months (Fig 1). From 17 (3%) of these 547 patients, no VL measurement was available; the baseline CD4 count was done > 7 days after start of cART in 3 patients (1%); from 22 (4%) no CD4 count was available at the 12-month time point; and from 39 patients (7%) the dates of the end-of-observation period CD4 count and VL measurement were >60 days apart. Thus, essential measurements were available for 466 patients still in care at 12 months.

**Nested Cohort 1** consisted of patients who had undetectable VL at 12 months (n = 382). Nested Cohort 1 allowed us to study what is referred to as “immunological cART discordance”. This was defined as an increase of <100 CD4 cells/mm$^3$ at 12 months compared to baseline in spite of full virological suppression (VL<40 copies/mL).

**Nested Cohort 2** consisted of patients enrolled in the prospective study who had experienced an increase in CD4 count of ≥100 cells/mm$^3$ at 12 months compared to baseline (n = 326). Nested Cohort 2 allowed us to examine “virological cART discordance”, defined as an increase of CD4 count of ≥100 cells/mm$^3$ in the first 12 months and incomplete viral suppression at month 12.

Of note, the definitions used to define suboptimal immunological (increase of <100 CD4 cells/mm$^3$) and virological responses (VL≥40 copies/mL) at twelve months were intentionally broad, relative to WHO definitions of immunological and virological treatment failure,[23] in order to encompass the majority of patients at risk of either type of treatment failure.

Statistical analysis

Bivariate logistic regression analysis was done to identify factors that were associated with discordant treatment responses. Subsequently, multivariable logistic regression was performed to identify independent determinants of a discordant treatment response. The variables sex, age, CD4 count at baseline and region were retained in the model irrespective of P values. Other variables were included into a starting model if they were associated with the outcome at P<0.20 in bivariate analysis. These variables were dropped one by one, based on a criterion of P<0.05, using the likelihood ratio test, until a parsimonious model was obtained.

In a secondary analysis, we examined whether adherence to cART, assessed 3 and 12-months (+/- 1 month) after the start of cART, was predictive of discordant treatment responses.

All reported P values were two-sided. P values <0.05 were considered statistically significant. All analyses were done using Stata 11 (StataCorp, College Station, Texas, USA).

Ethical approval

The study protocol was approved by the Rwanda National Ethics Committee Kigali, Rwanda and the Partners Human Research Committee, Boston, USA. All individual patients recruited into the study signed consent forms before they were enrolled. Data were double entered into an electronic medical record system (OpenMRS) that was password-protected; patient identification codes instead of names were used during analysis to ensure the confidentiality.
Results

Of the 466 patients who were retained in care after 12 months and who had a 12-month VL and CD4 count measurement done, 140 had a CD4 rise <100 cells/mm³ (Table 1). Of these, 28 had a detectable VL while 112 had full viral suppression (i.e., a discordant immunological

Table 1. Virological and immunological treatment responses among 466 patients retained in care at 12 months and with full data, Rwanda, 2007–2008.

| CD4 increase | Viral Load | Nested Cohort 1 | Nested Cohort 2 | All |
|--------------|------------|-----------------|-----------------|-----|
| <100 cells/mm³ | <40 copies/mL | 112 | 28 | 140 |
| ≥100 cells/mm³ | ≥40 copies/mL | 270 | 56 | 326 |
| All          | All        | 382 | 84 | 466 |

doi:10.1371/journal.pone.0159446.t001

doi:10.1371/journal.pone.0159446.g001
response). Only seven of those 28 had a VL of 1,000 copies/mL or above. Thus, of patients identified with a limited CD4 rise after 12 months, only 5% (7/140; 95%CI 2–10%) had virological failure (here defined as a single VL ≥1,000 copies/mL). With regard to virological discordance, 84 of 466 patients had a viral load ≥ 40 copies/mL at month 12, of whom 56 experienced a CD4 count increase of ≥100 cells/mm³.

Analysis of immunological discordant treatment responses: Nested Cohort 1

Table 2 shows the baseline characteristics of Nested Cohort 1, alongside the characteristics of the full cohort. The median age (interquartile range [IQR]) was 40 (34–47) years and 64% of patients were female. The median (IQR) CD4 count at baseline was 240 (150–295) cells/mm³ and 46% (177/382) of patients were in WHO stage 3 or 4. The median time between baseline CD4 count and start of cART was 22 (9–41) days.

Table 3 shows the follow-up data of Nested Cohort 1, next to the data of the full cohort for comparison. The median (IQR) CD4 count at 12 months was 390 (179–502) cells/mm³ and the median increase between the baseline and the 12-month CD4 count was 163 (87–259) cells/mm³.

Of the 382 patients in Nested Cohort 1, 112 (29%) had a rise in CD4 cell count <100 cells/mm³ between baseline and 12-month measurements. In bivariate analysis, only older age was significantly associated with an immunological discordant response (Table 4). In multivariable analysis, older age (P = 0.002), and having a longer travel time to the clinic (P = 0.01) were significantly associated with an immunological discordant response. Those from the Kayonza/Kirehe region appeared to be less likely to have a discordant treatment response; however, this difference was not statistically significant (P = 0.09). There was no association between baseline CD4 count and an immunological discordant response (P = 0.9), and none of the other demographic or health factors were associated (Table 4).

Of the 382 patients included in the analysis of an immunological discordant treatment response, 326 (85%) had a 3-month adherence measurement (assessed at a median of 85 days (IQR 82–98) after start of cART). In bivariate analysis good adherence at 3 months was not significantly associated with a lower risk for a discordant treatment response at 12 months (OR = 0.5, 95%CI 0.3–1.2, P = 0.1). When adherence was added to the multivariable model obtained in the primary analysis, good adherence was also not significantly associated (aOR = 0.5, 95%CI 0.2–1.1, P = 0.08). The effect size of all other variables in the model only changed marginally (data not shown). A similar analysis was done for adherence at 12 months; 12-months adherence was not associated with a discordant treatment response (P = 0.7), nor did it change the effect of the other variables in the model.

Analysis of virological discordant treatment responses: Nested Cohort 2

Table 2 shows the baseline characteristics of Nested Cohort 2, alongside the characteristics of the full cohort and those of Nested Cohort 1. The median age (interquartile range [IQR]) was 39 (33–46) years and 63% of patients were female. The median (IQR) CD4 count at baseline was 239 (155–294) cells/mm³. Forty-six percent (149/326) of patients were in WHO stage 3 or 4 at the start of the treatment. The median time between baseline CD4 count and the date of start of cART was 23 (9–42) days.

Table 3 shows the follow-up data of Nested Cohort 2, next to the data of the full cohort and of Nested Cohort 1 for comparison. The median (IQR) CD4 count at 12 months was 454 (362–541) cells/mm³ and the median increase between the baseline CD4 count and the 12-month
## Table 2. Baseline characteristics of Nested Cohorts 1 and 2 and of the full cohort of 610 HIV infected patients starting cART, Rwanda, 2007–2008.

|                  | Full Cohort N = 610 | Nested Cohort 1 (for the analysis of immunological discordant cART response) N = 382 | Nested Cohort 2 (for the analysis of virological discordant cART response) N = 326 |
|------------------|---------------------|------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|
| **Age**          |                     |                                                                                         |                                                                                         |
| Median age (IQR) in years | 40 (34–47)          | 40 (34–47)                                                                              | 39 (33–46)                                                                              |
| 20–34 years      | 159 (26.1%)         | 98 (25.7%)                                                                              | 101 (31.0%)                                                                             |
| 35–44 years      | 256 (42.0%)         | 159 (41.6%)                                                                              | 130 (39.9%)                                                                             |
| ≥45 years        | 195 (32.0%)         | 125 (32.7%)                                                                              | 95 (29.1%)                                                                              |
| **Sex**          |                     |                                                                                         |                                                                                         |
| Male             | 234 (38.4%)         | 139 (36.4%)                                                                              | 122 (37.4%)                                                                             |
| Female           | 376 (61.6%)         | 243 (63.6%)                                                                              | 204 (62.6%)                                                                             |
| **Literacy**     |                     |                                                                                         |                                                                                         |
| Unable to read    | 203 (33.3%)         | 131 (34.3%)                                                                              | 113 (34.7%)                                                                             |
| Able to read     | 407 (66.7%)         | 251 (65.7%)                                                                              | 213 (65.3%)                                                                             |
| **Marital status** |                   |                                                                                         |                                                                                         |
| Single           | 28 (4.6%)           | 16 (4.2%)                                                                                | 16 (4.9%)                                                                               |
| Married/cohabiting | 351 (57.8%)       | 221 (58.9%)                                                                              | 193 (59.2%)                                                                             |
| Divorced/separated | 60 (9.9%)         | 38 (10.0%)                                                                               | 33 (10.1%)                                                                              |
| Widowed          | 168 (27.7%)         | 107 (28.0%)                                                                              | 84 (25.8%)                                                                              |
| Missing          | 3                   | -                                                                                        | -                                                                                        |
| **CD4 count at baseline** |     |                                                                                         |                                                                                         |
| Median CD4 (IQR) cells/mm³ | 231 (148–289)     | 240 (150–295)                                                                             | 239 (155–294)                                                                           |
| <100 cells/mm³   | 91 (14.9%)          | 53 (13.9%)                                                                               | 43 (13.2%)                                                                             |
| 100–199 cells/mm³ | 149 (24.4%)         | 82 (21.5%)                                                                               | 78 (23.9%)                                                                             |
| 200–350 cells/mm³ | 370 (60.7%)         | 247 (64.7%)                                                                              | 205 (62.9%)                                                                             |
| Median time (IQR) in days CD4 count—start cART | 22 (9–41) | 22 (9–41) | 23 (9–42) |
| WHO Stage at baseline |                  |                                                                                         |                                                                                         |
| 1                | 117 (19.2%)         | 73 (19.2%)                                                                               | 61 (18.8%)                                                                             |
| 2                | 196 (32.2%)         | 131 (34.4%)                                                                              | 115 (35.4%)                                                                             |
| 3                | 281 (46.1%)         | 168 (44.1%)                                                                              | 139 (42.8%)                                                                             |
| 4                | 15 (2.5%)           | 9 (2.4%)                                                                                 | 10 (3.1%)                                                                              |
| Missing          | 1                   | 1                                                                                        | 1                                                                                        |
| **BMI at baseline in kg/m²** |          |                                                                                         |                                                                                         |
| Median BMI       | 20.7 (18.6–22.7)    | 20.7 (18.7–22.7)                                                                         | 20.8 (18.8–22.9)                                                                        |
| <18.5            | 151 (25.0%)         | 87 (22.9%)                                                                               | 69 (21.3%)                                                                             |
| 18.5–24.9        | 398 (65.8%)         | 256 (67.2%)                                                                              | 223 (68.4%)                                                                             |
| ≥25              | 56 (9.3%)           | 37 (9.7%)                                                                                | 32 (9.8%)                                                                              |
| Missing          | 5                   | 2                                                                                        | 2                                                                                        |
| **On TB treatment at start of cART** |          |                                                                                         |                                                                                         |
| No               | 585 (96.2%)         | 366 (96.1%)                                                                              | 313 (96.3%)                                                                             |
| Yes              | 23 (3.8%)           | 15 (3.9%)                                                                                | 12 (3.7%)                                                                              |
| Missing          | 2                   | 1                                                                                        | 1                                                                                        |
| **Regimen with d4T** |                  |                                                                                         |                                                                                         |
| No               | 191 (31.3%)         | 117 (30.9%)                                                                              | 95 (29.1%)                                                                             |
| Yes              | 416 (68.5%)         | 264 (69.3%)                                                                              | 229 (70.7%)                                                                             |
| Missing          | 3                   | 1                                                                                        | 2                                                                                        |
CD4 count was 206 (156–297) cells/mm$^3$; by definition, all had an increase in CD4 count of at least 100 cells/mm$^3$.

Of the 326 patients in Nested Cohort 2, 56 (17%) did not have full virological suppression. In bivariate analysis women had a lower risk for a virological discordant response (OR = 0.5, 95%CI 0.3–1.0; $P = 0.03$), but none of the other variables were associated with a virological discordant response (Table 4). In multivariable analysis, including (a priori) age, sex, baseline CD4 count and region, none of the variables were significantly associated with a discordant virological response (Table 4), although female sex was of borderline significance ($P = 0.05$).

Sixteen of the 326 patients in Nested Cohort 2 (5%) had a VL of 1,000 copies/mL or above. We also assessed determinants of a discordant treatment response defined in this less strict way. No significant determinants of such a discordant treatment response were identified, but power was limited.

An adherence measurement at 3 months was available for 278 (85%). In bivariate analysis good adherence at 3 months was not associated with a virological discordant treatment response (OR = 0.7, 95%CI 0.3–2.1; $P = 0.5$). When we added adherence to the previously obtained multivariable model, good adherence at 3 months was not a significant determinant of this type of discordant treatment response (aOR = 0.7, 95%CI 0.2–2.2; $P = 0.5$); the effect of the other variables did not change substantially (data not shown). A similar analysis was done for adherence at 12 months; 12-months good adherence was associated with a non-significantly decreased likelihood for a discordant treatment response (OR = 0.6, 95%CI 0.3–1.3, $P = 0.2$). When the 12-month adherence was added to the multivariable model, it did not change the effect of the other variables in the model substantially, but good adherence was associated with lower odds of a discordant treatment response (aOR = 0.5, 95%CI 0.2–1.2, $P = 0.1$).
Among patients with a CD4 rise less than 100 cells/mm$^3$, 20% had a detectable VL, but only 5% had virological failure defined as VL $>1,000$ copies/mL. Conversely, 17% of patients with a CD4 rise of 100 cells/mm$^3$ or more did not have full virological suppression. Together, these results indicate the poor predictive value of a rise in CD4 count for virological suppression and underscore the importance of viral load testing for treatment monitoring. Discordant treatment responses (i.e., the absence of an adequate immunological response despite an undetectable VL or vice versa) were observed in 36% of cART-naïve HIV patients in Rwanda, 12 months after the start of cART. Older age and long travel distance to the clinic were associated with an immunological discordant treatment response.

**Table 3. Follow-up data of 610 HIV patients on cART, Rwanda, 2007–2008.**

| CD4 count at 12 months | Full cohort N = 610 | Nested Cohort 1 N = 382 | Nested Cohort 2 N = 326 |
|------------------------|---------------------|-------------------------|-------------------------|
| Median CD4 (IQR) cells/mm$^3$ | 392 (275–507) | 390 (179–502) | 454 (362–541) |
| <100 cells/mm$^3$ | 11 (1.8%) | 7 (1.8%) | - |
| 100–199 cells/mm$^3$ | 47 (9.0%) | 32 (8.4%) | 12 (3.7%) |
| 200–349 cells/mm$^3$ | 145 (27.7%) | 110 (28.8%) | 57 (17.5%) |
| 350–499 cells/mm$^3$ | 182 (34.8%) | 134 (35.1%) | 141 (43.3%) |
| ≥500 cells/mm$^3$ | 137 (26.2%) | 99 (25.9%) | 116 (35.6%) |
| Missing | 88 | - | - |

**Difference in CD4 count between baseline and 12 months**

| Median difference (IQR) cells/mm$^3$ | Full cohort N = 610 | Nested Cohort 1 N = 382 | Nested Cohort 2 N = 326 |
|--------------------------------------|---------------------|-------------------------|-------------------------|
| <0 cells/mm$^3$ | 29 (4.8%) | 20 (5.2%) | - |
| 0–99 cells/mm$^3$ | 127 (24.3%) | 92 (24.1%) | - |
| 100–199 cells/mm$^3$ | 167 (31.9%) | 126 (33.0%) | 155 (47.6%) |
| 200–299 cells/mm$^3$ | 102 (19.5%) | 74 (19.4%) | 90 (27.6%) |
| ≥300 cells/mm$^3$ | 97 (18.6%) | 70 (18.3%) | 81 (24.9%) |
| Missing | 88 | - | - |

**Viral load after 12 months of treatment**

| Median VL (IQR) copies/mL | Full cohort N = 610 | Nested Cohort 1 N = 382 | Nested Cohort 2 N = 326 |
|---------------------------|---------------------|-------------------------|-------------------------|
| <40 copies/mL | 39.9 (39.9–39.9) | 39.9 (39.9–39.9) | 39.9 (39.9–39.9) |
| 40–999 copies/mL | 430 (81.1%) | 382 (100.0%) | 270 (82.8%) |
| 1,000–9,999 copies/mL | 71 (13.4%) | - | 40 (12.3%) |
| ≥10,000 copies/mL | 16 (3.0%) | - | 10 (3.1%) |
| Missing viral load | 80 | - | - |

**Outcome at 12 months follow-up**

| Died | 35 (5.7%) | - | - |
| Defaulted | 13 (2.1%) | - | - |
| Transferred out | 15 (2.5%) | - | - |
| Retained in care, undetectable viral load | 430 (70.5%) | 382 (100.0%) | 270 (82.8%) |
| Retained in care, detectable viral load | 100 (16.4%) | - | 56 (17.2%) |
| Retained in care but no viral load measured done at 12 mo | 17 (2.8%) | - | - |
| Median time (IQR) between baseline & 12 mo CD4 count in days | 386 (372–413) | 386 (372–412) | 391 (372–411) |
| Median time (IQR) between start of cART & 12 mo VL in days | 366 (361–374) | 366 (361–374) | 366 (361–374) |
| Median time (IQR) between start of cART & 12 mo CD4 count in days | 364 (356–374) | 364 (357–373) | 364 (356–373) |

Numbers in **Table 3** are N (%), unless mentioned otherwise. HIV Human immunodeficiency virus; cART combination antiretroviral treatment; IQR Interquartile range; N Number; mo month; VL Viral load.

doi:10.1371/journal.pone.0159446.t003

**Discussion**

Among patients with a CD4 rise less than 100 cells/mm$^3$, 20% had a detectable VL, but only 5% had virological failure defined as VL $>1,000$ copies/mL. Conversely, 17% of patients with a CD4 rise of 100 cells/mm$^3$ or more did not have full virological suppression. Together, these results indicate the poor predictive value of a rise in CD4 count for virological suppression and underscore the importance of viral load testing for treatment monitoring. Discordant treatment responses (i.e., the absence of an adequate immunological response despite an undetectable VL or vice versa) were observed in 36% of cART-naïve HIV patients in Rwanda, 12 months after the start of cART. Older age and long travel distance to the clinic were associated with an immunological discordant treatment response.
Table 4. Analysis of determinants of immunological discordant treatment responses (CD4 count rise <100 cells/mm³ despite complete virological suppression) in Nested Cohort 1, and of virological discordant treatment responses (CD4 count rise ≥100 cells/mm³ but incomplete virological suppression, i.e. VL≥40 copies/mL) in Nested Cohort 2, Rwanda, 2007–2008.

|                          | Immunological discordant response (Nested Cohort 1) | Virological discordant response (Nested Cohort 2) |
|--------------------------|------------------------------------------------------|--------------------------------------------------|
|                          | Bivariate analysis | Multivariable analysis                        | Bivariate analysis | Multivariable analysis |
|                          | n/N (%) | OR (95% CI) | P | aOR (95% CI) | P | n/N (%) | OR (95% CI) | P | aOR (95% CI) | P |
| Age-group                |          |            |    |              |    |          |            |    |              |    |
| 20–34 years              | 112/382 (29.3%) | 1          | 1  | 1            | 1  | 56/326 (17.2%) | 1  | 0.36 | 0.34 |
| 35–44 years              | 47/159 (29.6%) | 2.3 (1.2–4.4) | 2.4 (1.2–4.6) | 18/130 (13.9%) | 0.7 (0.4–1.5) | 0.6 (0.3–1.3) |
| ≥45 years                | 50/125 (40.0%) | 3.7 (1.9–7.1) | 3.9 (2.0–7.8) | 20/95 (21.1%) | 1.2 (0.6–2.5) | 1.0 (0.5–2.2) |
| Sex                      |          |            |    |              |    |          |            |    |              |    |
| Male                     | 45/139 (32.4%) | 1          | 1  | 28/122 (23.0%) | 1  | 1          |            |    |              |    |
| Female                   | 67/243 (27.6%) | 0.8 (0.5–1.3) | 0.8 (0.5–1.3) | 28/204 (13.7%) | 0.5 (0.3–1.0) | 0.5 (0.3–1.0) |
| Marital status           |          |            |    |              |    |          |            |    |              |    |
| Single                   | 3/16 (18.8%) | 0.6 (0.2–2.1) | 3/16 (18.8%) | 1.1 (0.3–4.0) |
| Married/cohabiting       | 62/221 (28.1%) | 1          | 1  | 34/193 (17.6%) | 1  | 1          |            |    |              |    |
| Divorced/separated       | 11/38 (29.0%) | 1.0 (0.5–2.2) | 6/33 (18.2%) | 1.0 (0.4–2.7) |
| Widowed                  | 36/107 (33.6%) | 1.3 (0.8–2.1) | 13/84 (15.5%) | 0.9 (0.4–1.7) |
| Literacy                 |          |            |    |              |    |          |            |    |              |    |
| Unable to read           | 41/131 (31.3%) | 1          | 23/113 (20.4%) | 1  |
| Able to read             | 71/251 (28.3%) | 0.9 (0.5–1.4) | 33/213 (15.5%) | 0.7 (0.4–1.3) |
| CD4 count at baseline    |          |            |    |              |    |          |            |    |              |    |
| <100 cells/mm³           | 16/53 (30.2%) | 1          | 1  | 6/43 (14.0%) | 1  | 1          |            |    |              |    |
| 100–199 cells/mm³        | 22/82 (26.8%) | 0.8 (0.4–1.8) | 1.1 (0.5–2.1) | 18/78 (23.1%) | 1.9 (0.7–5.1) | 2.1 (0.7–5.8) |
| 200–350 cells/mm³        | 74/247 (30.0%) | 1.0 (0.5–1.9) | 1.1 (0.6–2.2) | 32/205 (15.6%) | 1.1 (0.4–2.9) | 1.3 (0.5–3.3) |
| WHO Stage at baseline    |          |            |    |              |    |          |            |    |              |    |
| 1 and 2                  | 58/204 (28.4%) | 1          | 30/176 (17.1%) | 1  |
| 3 and 4                  | 54/177 (30.5%) | 1.1 (0.7–1.7) | 26/149 (17.5%) | 1.0 (0.6–1.8) |
| BMI at baseline          |          |            |    |              |    |          |            |    |              |    |
| <18.5 kg/m²              | 30/87 (34.5%) | 1.4 (0.8–2.3) | 12/69 (17.4%) | 1.0 (0.5–2.1) |
| 18.5–24.9 kg/m²          | 71/256 (27.7%) | 1          | 38/223 (17.0%) | 1  |
| ≥25 kg/m²                | 11/37 (29.7%) | 1.1 (0.5–2.3) | 6/32 (18.8%) | 1.1 (0.4–3.0) |
| On TB treatment at start cART |          |            |    |              |    |          |            |    |              |    |
| No                       | 109/366 (29.8%) | 1          | 56/313 (17.9%) | N.A. |
| Yes                      | 3/15 (20.0%) | 0.6 (0.2–2.1) | 0/12 (0.0%) |
| Regimen with d4T         |          |            |    |              |    |          |            |    |              |    |
| No                       | 39/117 (33.3%) | 1          | 17/95 (17.9%) | 1  |
| Yes                      | 73/264 (27.7%) | 0.8 (0.5–1.2) | 38/229 (16.6%) | 0.9 (0.5–1.7) |
| Regimen with NVP         |          |            |    |              |    |          |            |    |              |    |
| No                       | 9/34 (26.5%) | 1          | 5/30 (16.7%) | 1  |
| Yes                      | 103/347 (29.7%) | 1.2 (0.5–2.6) | 50/294 (17.0%) | 1.0 (0.4–2.8) |
| Travel time to clinic    |          |            |    |              |    |          |            |    |              |    |
| <30 minutes              | 23/78 (29.5%) | 1          | 7/62 (11.3%) | 1  |
| Between 30 & 60 min.     | 27/93 (29.0%) | 1.0 (0.5–1.9) | 1.0 (0.5–2.0) | 17/83 (20.5%) | 2.0 (0.8–5.2) |
| Between 1 & 2 hrs        | 28/124 (22.6%) | 0.7 (0.4–1.3) | 0.8 (0.4–1.5) | 17/113 (15.0%) | 1.4 (0.5–3.6) |
| > 2 hrs                  | 34/86 (39.5%) | 1.6 (0.8–3.0) | 2.4 (1.1–5.5) | 15/67 (22.4%) | 2.3 (0.9–6.0) |
| Region                   |          |            |    |              |    |          |            |    |              |    |
| Ruhengeri                | 60/197 (30.5%) | 1          | 26/163 (16.0%) | 1  |

(Continued)
The frequency of immunological discordant treatment response in this study population was higher than that observed in several other studies, where the proportion ranged between 9% and 28%.\(^{[1,6–8,10,24–27]}\) Immunological discordant treatment response remains an unsolved challenge to physicians involved in long-term HIV care and treatment. In agreement with most other studies we found that older age was a significant risk factor for immunological discordant treatment responses.\(^{[1,6,7,9,10,15]}\) The degree of immune restoration is dependent on the thymic function; as thymic function decreases with age,\(^{[9,24,25]}\) this may explain why older people are at higher risk of incomplete immune restoration. All our patients were on a regimen including a non-nucleoside reverse transcriptase inhibitor (NNRTI). NNRTIs are known to be associated with a lower CD4 increase after the start of cART,\(^{[26]}\) with particular evidence of this effect for zidovudine.\(^{[28,29]}\) In our study, neither type of NNRTI (zidovudine or stavudine) was associated with either type of discordant treatment response.

Most studies found that low VL at start of cART was associated with immunological discordant treatment responses,\(^{[6,7,10,15]}\) but as we did not have baseline VL available, we could not assess this in our cohort. Regarding the effect of baseline or nadir CD4 count, studies have provided contradictory results. Some found that a low baseline or nadir CD4 count was predictive of immunological discordant treatment responses,\(^{[1,6,15]}\) while others found the reverse,\(^{[7,10]}\) and some studies found no association between baseline CD4 count and immunological discordant treatment response.\(^{[9,11]}\) The remarkable finding that a higher baseline CD4 count is a significant predictor of a discordant treatment response is probably due to regression to the mean. Due to random variation in the measurement of CD4 counts and due to individual variation of CD4 counts, those with the highest CD4 counts are bound to have lower CD4 counts upon subsequent measurements even in the absence of any intervention.\(^{[8,27]}\) In our cohort we did not observe any effect of baseline CD4 count on discordant treatment responses. Evolving definitions of discordant treatment response,\(^{[23,30,31]}\) and variation in the definitions used across studies may in part explain discrepancies in findings as well as variability in the frequency of this condition.\(^{[1,7,11]}\)

We also found that an immunological discordant treatment response was more common in patients who had to travel more than two hours to the clinic. This was not explained by a lower adherence among them and could be a chance finding. In a previously published analysis of this cohort,\(^{[32]}\) time to clinic was not associated with retention with VL suppression. Future investigations could further explore the association between travel time to health clinics, and treatment outcomes and discordant responses. We did not find that TB treatment at the start of cART predicted an immunological discordant response though only a small number of patients had TB.

Discordant virological treatment response—incomplete suppression of viral replication but a good immunological response—was observed in 17% of patients with a CD4 rise of 100 cells/
mm³ or more. We could not identify any independent determinants of such discordant treatment response, although female sex was associated with lower odds of discordance at $P = 0.03$. The interpretation of these data is not straightforward, as the VL might have become detectable just because of a brief interruption of therapy. This is supported by the analysis in which adherence at 12-months was included; poor adherence at that time point was significantly associated with a discordant response. Ongoing screening for suboptimal adherence and targeted, intensive counselling may improve virological responses and prevent acquired resistance for individuals with inconsistent adherence. Repeat CD4 count and viral load assessments in this group will be critical to distinguishing between incomplete viral suppression due to suboptimal adherence and that due to drug resistance. Most of the patients with detectable VL (71%; 40/56) had VL <1,000 copies/ml. We only had a single VL measurement, so we could not distinguish between transient viremia, also referred to as “viral blips,” or full-blown prolonged viremia associated with poor adherence or resistance. Nevertheless, these data underscore that clinicians should be equally careful not to switch cART too early because of transient viremia, especially when viral blips do not exceed 400 copies/mL. In this case, adherence counselling should be considered, as viral blips are often associated with short-term lapses in adherence rather than resistance.[33,34]

Because this was a secondary analysis of a prospective study of the first year of cART, and viral load was conducted one time at the end of study follow-up, we lacked information regarding physician’s decision-making patterns in the absence of viral load monitoring. Although Rwanda has since implemented routine viral load monitoring, many settings still lack access to this important tool. In such programs limited CD4 count rises might be erroneously interpreted as treatment failure and patients may subsequently be switched to second line regimens. In our study only 5% of patients with a limited CD4 rise after 12 months had virological failure (VL ≥1,000 copies/mL). This suggests that in most such cases the (costly) treatment change is unnecessary. VL tests are needed to avoid this; in public health clinics such tests could differentiate between ART failure and a (less threatening) discordant treatment response. Nevertheless, both discordant treatment responses are associated with poorer outcomes,[6,8,9,11] and should, if diagnosed, be regarded as a danger sign to the clinician.

Limitations

Our study has several limitations. We did not have baseline VL measurements and for the definition of viral success we relied on a single VL determination approximately 12 months after the start of cART. If that single VL determination was not representative of the VL over time in a particular patient, misclassification might have occurred, regarding some patients as virological responders who in fact were not having a complete virological response to therapy and vice versa. Furthermore, a single VL measurement of more than 1,000 copies/mL does not infallibly identify the development of viral resistance to the antiretroviral therapy used. A high VL can be caused by temporary treatment interruptions, which is usually quickly and fully re-suppressed after re-initiation of cART. Also, we relied on only one CD4 count taken at approximately 12 months. Daily fluctuations in CD4 counts and fluctuations induced by opportunistic infections are common, so this may have led to another misclassification, in two directions: regarding some persons unfairly as immunological non-responders and others unfairly as immunological responders. So long as the misclassification was not associated with the predictors under study, we would expect it to lead to underestimations of the true associations. This misclassification might have reduced power to detect significant associations with baseline variables.
The size of the cohort and the small number of events (especially that of virological discordance) meant that the power of this study to identify significant determinants was limited, and this may be the reason we could not confirm some of the associations that earlier studies had found. Lastly, there may be other factors that were not measured in this cohort (i.e., drug substitutions, smoking) that predict a discordant response.

**Conclusion**

Discordant treatment response are relatively common and signal a patient who is vulnerable to negative health outcomes and may benefit from intensified adherence counseling and repeat laboratory monitoring over the short term. When VL monitoring is not done (as is the case in many ART programs in sub-Saharan African countries), a limited increase of the CD4 count after 12 months could be misinterpreted as a (virological) treatment failure and lead to unnecessary changes to more expensive second-line ART regimens. Development and implementation of low-cost point of care VL tests, coupled with ongoing adherence outcomes and a strong drug supply chain, will likely help optimize cART outcomes. Future research should be conducted to further understand the long-term consequences of discordant immunological treatment responses.

**Acknowledgments**

We acknowledge and thank the Doris Duke Charitable Foundation for financial support of this research. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. Technical support was provided by the Infectious Disease Network for Treatment and Research in Africa (INTERACT), funded by the Netherlands Organization for Scientific Research/Netherlands Foundation for the Advancement of Tropical Research (NWO/WOTRO) and the European Union (SANTE/2006/105-316). We are grateful to the study staff: Adrienne Socci, Massudi Hakizamungu, Wellars Ndayambaje, Eline Uwitonze, Albertine Mukeshimana, Ernest Nyirinkindi, Jean Damascene Uwamuhoro, Carine Dusenge, Claire Dusabe, and Jean Claude Nyiramana. We also thank Cheryl Amoroso, Benjamin Akimana, Christian Allen, Darius Jazayeri, Ellen Ball, the Rwanda-based PIH-EMR team, and Laboratory Management of the Rwanda National Reference Laboratory. We also thank Frank Cobelens for providing feedback on a draft of this manuscript.

**Author Contributions**

Conceived and designed the experiments: FRK MSVDL MIB MFF MLR. Analyzed the data: FRK MSVDL MIB CAR. Wrote the paper: FRK MSVDL MIB MFF MLR EB CAR FWNMW. Recruitment and follow up: FRK MFF EB. Data management: MFF EB. Data interpretation: FRK MFF MIB FWNMW MSVDL. Wrote first draft: FRK MSVDL.

**References**

1. Zoufaly A, van der Heiden M, Kollan C, Bogner JR, Fätkenheuer G, Wasmuth JC et al. Clinical outcome of HIV-infected patients with discordant virological and immunological response to antiretroviral therapy. J Infect Dis 2011; 203: 364–71. doi:10.1093/jinfdis/jiq055 PMID: 21208929
2. Coffey S, Volberding PA. Chapter 11: Overview of antiretroviral therapy. In: Volberding PA, Greene WC, Lange JMA, Gallant JE, Sewankambo N (eds). Sande’s HIV/AIDS Medicine. 2nd Ed. StLouis: Elsevier, 2012.
3. Mutevedzi PC, Lessells RJ, Rodger AJ, Newell ML. Association of age with mortality and virological and immunological response to antiretroviral therapy in rural South African adults. PLoS One 2011; 6: 21795.
4. Mocroft A, Phillips AN, Gatell J, Ledergerber B, Fisher M, Clumeck N, et al. Normalization of CD4 counts in patients with HIV-1 infection and maximum virological suppression who are taking combination antiretroviral therapy: an observational cohort study. Lancet 2007; 360: 407–10. PMID: 17659333

5. Bartlett JA, Fath MJ, Demasi R, Hermes A, Quinn J, Mondou E, et al. An updated systematic overview of triple combination therapy in antiretroviral-naive HIV-infected adults. AIDS 2006; 20: 2051–64. PMID: 17053351

6. Moore DM, Hogg RS, Yip B, Wood E, Tyndall M, Braithstein P, et al. Discordant immunologic and virologic responses to highly active antiretroviral therapy are associated with increased mortality and poor adherence to therapy. J Acquir Immune Defic Syndr 2005; 40: 288–93. PMID: 16249702

7. Tuboi SH, Brinkhof MW, Egger M, Stone RA, Braithstein P, Nash D, et al. Discordant responses to potent antiretroviral treatment in previously naive HIV-1-infected adults initiating treatment in resource-constrained countries: the antiretroviral therapy in low-income countries (ART-LINC) collaboration. J Acquir Immune Defic Syndr 2007; 45: 52–9. PMID: 17460471

8. Tuboi SH, Pacheco AG, Harrison LH, Stone RA, May M, Brinkhof MW, et al. Mortality associated with discordant responses to antiretroviral therapy in resource-constrained settings. J Acquir Immune Defic Syndr 2010; 53: 70–7. doi: 10.1097/QAI.0b013e3181c22d19 PMID: 20035163

9. Piketty C, Weiss L, Mohamed AS, Belec L, Kazatchkine MD. Long-term clinical outcome of human immunodeficiency virus-infected patients with discordant immunologic and virologic responses to a protease inhibitor-containing regimen. J Infect Dis 2001; 183: 1328–35. PMID: 11294663

10. Gilson RJ, Man SL, Copas A, Rider A, Forsyth S, Hill T, et al. Discordant responses on starting highly active antiretroviral therapy: suboptimal CD4 increases despite early viral suppression in the UK Collaborative HIV Cohort (UK CHIC) Study. HIV Med 2010; 11: 152–60. doi: 10.1111/j.1468-1293.2009.00755.x PMID: 19732175

11. Tan R, Westfall AO, Willig JH, Mugavero MJ, Saag MS, Kaslow RA, et al. Clinical outcome of HIV-infected antiretroviral-naive patients with discordant immunologic and virologic responses to highly active antiretroviral therapy. J Acquir Immune Defic Syndr 2008; 47: 553–6. doi: 10.1097/QAI.0b013e31819168565 PMID: 18285713

12. Julg B, Poole D, Ghebremichael M, Castilla C, Altford M, et al. (2012) Factors predicting discordant virological and immunological responses to antiretroviral therapy in HIV-1 clade C infected Zulu/Xhosa in South Africa. PLoS One 7: e31161. doi: 10.1371/journal.pone.0031161 PMID: 22348047

13. Prabhakar B, Banu A, Pavithra HB, Chandrashekhara P, Sasthri S (2011) Immunological failure despite antiretroviral therapy: suboptimal CD4 lymphocyte count response to HAART despite full viral suppression in the EuroSIDA study. HIV Med 2003; 4: 255–62. PMID: 12859325

14. Florence E, Lundgren J, Dreezen C, Fisher M, Kirk O, Blaxhult A, et al. Factors associated with a reduced CD4 lymphocyte count response to HAART despite full viral suppression in the EuroSIDA study. HIV Med 2003; 4: 255–62. PMID: 12859325

16. Consolidated Guidelines on the use of antiretroviral drugs for treating and preventing HIV Infection. Geneva: World Health Organization; 2013. Available: www.who.int/hiv/pub/guidelines/arv2013.

17. Joint United Nations Programme on HIV/AIDS (UNAIDS). Global AIDS epidemic 2013. Geneva: UNAIDS; 2013.

18. Nsanzimana S, Ruton H, Lowrance DW, Cishahayo S, Nyemazi JP, Muhayimpundu R, et al. Cell Phone- and Internet-based Monitoring and Evaluation of the National Antiretroviral Treatment Program during Rapid Scale-up in Rwanda: TRACnet, 2004–2010. J Acquir Immune Defic Syndr 2012; 59: e17–23. doi: 10.1097/QAI.0b013e31823e2278 PMID: 22067668

19. Franke MF, Kaigamba F, Socci AR, Hakizamunzu M, Patel A, Bagiruwigize E, et al. Improved retention associated with community-based accompaniment for antiretroviral therapy delivery in rural Rwanda. Clin Infect Dis 2013; 56: 1319–26. doi: 10.1093/cid/cis1193 PMID: 23249611

20. TRAC Plus, Ministry of Health, Rwanda: Guidelines for the provision of comprehensive care to persons infected HIV in Rwanda 2009. Kigali, Rwanda: Ministry of Health; 2009.

21. World Health Organization: Rapid advice: antiretroviral therapy for HIV infection in adults and adolescents: Geneva: World Health Organization; 2009.

22. Mannheimer SB, Mukherjee R, Hirschhorn LR, Dougherty J, Celano SA, Ciccarone D, et al. The CASE adherence index: A novel method for measuring adherence to antiretroviral therapy. AIDS Care 2006; 18: 853–61. PMID: 16971298

23. World Health Organization: Antiretroviral therapy for HIV infection in adults and adolescents: Recommendations for a public health approach. Geneva: World Health Organization; 2006.
24. Casotti JA, Passos LN, Oliveira FJ, Cerutti C Jr. Prevalence of discordant immunologic and virologic responses in patients with AIDS under antiretroviral therapy in a specialized care center in Brazil. Rev Inst Med Trop Sao Paulo 2011; 53: 301–7. PMID: 22183451

25. Piketty C, Castel P, Belec L, Batisse D, Si Mohamed A, Gilquin J, et al. Discrepant responses to triple combination antiretroviral therapy in advanced HIV disease. AIDS 1998; 12: 745–50. PMID: 9619806

26. Dronda F, Moreno S, Moreno A, José LC, Pérez-Elias MJ, A. Long-Term Outcomes among Antiretroviral-Naive Human Immunodeficiency Virus–Infected Patients with Small Increases in CD4+ Cell Counts after Successful Virologic Suppression. Clin Infect Dis 2002; 35: 1005–1009. PMID: 12355389

27. Falster K, Petoumenos K, Chuah J, Mijch A, Mulhall B, Kelly M, et al. Poor baseline immune function predicts an incomplete immune response to combination antiretroviral therapy despite sustained viral suppression. J Acquir Immune Defic Syndr 2009; 50: 307–13. doi:10.1097/QAI.0b013e3181945ed4 PMID: 19194311

28. Wandeler G, Gsponer T, Mulenga L, Garone D, Wood R, et al. (2013) AZT Impairs Immunological Recovery on First-line ART: Collaborative analysis of cohort studies in Southern Africa. AIDS (London, England) 27: doi: 10.1097/QAD.1090b1013e328362d328887

29. Wandeler G, Gsponer T, Mulenga L, Garone D, Wood R, et al. (2013) Zidovudine impairs immunological recovery on first-line antiretroviral therapy: collaborative analysis of cohort studies in southern Africa. Aids 27: 2225–2232. doi: 10.1097/QAD.0b013e328362d887 PMID: 23660577

30. World Health Organization: Antiretroviral therapy for HIV infection in adults and adolescents: recommendations for a public health approach. Geneva: World Health Organization; 2010.

31. World Health Organization: Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection: Recommendations for a public health approach. Geneva: World Health Organization; 2013.

32. Gupta N, Muyarurinca C, Mutagomma M, Nyigena JW, Kayigamba F, et al. (2016) Community-Based Accompaniment Mitigates Predictors of Negative Outcomes for Adults on Antiretroviral Therapy in Rural Rwanda. AIDS Behav 20: 1009–1016. doi: 10.1007/s10461-015-1185-9 PMID: 26346334

33. Podsadecki TJ, Vrijens BC, Tousset EP, Rode RA, Hanna GJ (2007) Decreased adherence to antiretroviral therapy observed prior to transient human immunodeficiency virus type 1 viremia. J Infect Dis 196: 1773–1778. doi: 10.1086/523704 PMID: 18190257

34. Nettles RE, Kieffer TL, Kwon P, et al. (2005) Intermittent HIV-1 viremia (blips) and drug resistance in patients receiving haart. JAMA 293: 817–829. PMID: 15713771