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Early antiretroviral treatment of infants to attain HIV remission

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ABSTRACT

Background: Studies in adults and children suggested that starting antiretroviral therapy (ART) soon after infection positively influences early events in HIV infection raising the possibility that remission may be achieved in some.

Methods: We designed an analytic treatment interruption (ATI) trial to test the hypothesis that a sizable minority of HIV-infected neonates who initiated ART <14 days of birth and maintained on ART would be able to maintain viral suppression when ART was withdrawn. To yield the target cohort for this trial, 73 HIV-infected neonates identified at one hospital in Johannesburg, South Africa, were initiated on ART <14 days of birth and maintained on ART tracking viral load (VL) decline and immune recovery (clinicaltrials.gov # NCT02431975).

Findings: Three HIV-infected infants (4.1%) died and nine (12.3%) were lost to follow-up before 48 weeks of age. Of those surviving on study, 52.5% attained and sustained VL <50 copies/ml and half of these sustained CD4+ T-cell percentage >30% which were the primary entry criteria for the ATI trial. Proportions achieving ATI eligibility criteria were similar in the 46 infants starting ART <48 h (19.6%) to 27 infants starting 2–14 days (25.9%) (p = 0.567).

Interpretation: Very early ART on its own, using regimens available when the trial was designed, is insufficient to attain minimum entry criteria needed to justify our trial of ART interruption. Decisions about how quickly to start ART should be based on optimizing standard clinical outcomes rather than with the expectation that remission can be attained.

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1. Introduction

Studies in adults given antiretroviral therapy (ART) soon after primary infection have observed that some individuals can achieve periods of post-treatment viral control when ART is withdrawn (remission) [1–6]. The report of the infant in Mississippi who started treatment within 30 h of birth and who was able to maintain viral suppression off treatment for over two years raised the
2. Methods

2.1. Study design

We designed an ATI trial to test the hypothesis that a sizable minority (>20%) of HIV-infected neonates started on ART within 14 days of birth and maintained on ART for at least two years would be able to sustain viral suppression after ART withdrawal. To yield the target population for the ATI trial, we established clinical protocols at Rahima Moosa Mother and Child Hospital (RMMCH), Johannesburg, South Africa, to provide HIV diagnostic testing at birth to identify HIV-infected neonates with a positive result within 48 h of birth and b) to initiate ART within 14 days of birth and maintain ART to at least two years of age (clinicaltrials.gov # NCT02431975). RMMCH is a centrally-located, large urban public tertiary hospital and one of the teaching facilities of the University of the Witwatersrand. Children who met virologic and immunologic endpoints would be eligible for entry into the ATI trial. The protocols were approved by the Institutional Review Boards (IRB) of the University of the Witwatersrand and Columbia University. Written informed consent was obtained from mothers for their own and their infants’ participation. In addition to IRB approval, the ATI trial protocol was developed in consultation with a Data Safety and Monitoring Board (DSMB) who met annually and monitored progress of the target cohorts’ potential to enter the trial.

2.2. Participants

When the study was initially proposed, we hypothesized that ART would need to be initiated within 48 h of birth to have the later beneficial outcomes. For practical purposes, in order to start ART <48 h of birth, a point-of-care neonatal diagnosis program was established that used Xpert HIV-1 Qual (Cepheid, Sunnyvale, CA) [10]. The site already had a routine birth testing program sending samples to the local laboratory for diagnosis (HIV-1 total nucleic acid (TNA) COBAS TaqMan HIV-1 Qualitative Test Version 2.0 Roche Molecular Systems, Inc., Branchburg, NJ). As part of the routine program, all HIV-exposed neonates have blood collected by venipuncture after birth prior to discharge. When staff capacity permitted, one of the vials from this blood collection was tested on-site using Xpert. Positive Xpert results were re-run with residual sample from the same blood collection. Neonates with two positive Xpert results were eligible for immediate on-site ART initiation. The on-site testing protocol was intended to yield a cohort enriched with infants initiating ART <48 h after birth although inclusion criteria for the ATI trial required only ART initiation <14 days. Two clinical protocols were in place to recruit infants intended for the trial. These protocols included Xpert co-tested neonates as well as infants with positive or indeterminate results from the routine diagnosis program not co-tested with Xpert. The latter neonates were recalled to the site for ART initiation once routine results were available.

2.3. Procedures

The initial ART regimen consisted of nevirapine, lamivudine and zidovudine. Nevirapine was changed to lopinavir-ritonavir no sooner than 42 weeks post-menstrual age taking into account patient readiness. Drugs were given in liquid form twice daily. Either stavudine or abacavir were given in the event of zidovudine toxicity and abacavir substituted for zidovudine once infants were ≥3 months of age. Cotrimoxazole was started at 4–6 weeks of age. Routine infant prophylaxis was one dose of nevirapine as soon as possible after birth and daily nevirapine for 6 weeks after discharge. Infants considered high risk had twice-daily zidovudine added. For infected neonates identified before discharge, prophylaxis was discontinued and the treatment regimen initiated. For the infected neonates recalled to the site, prophylaxis was discontinued once ART was initiated.

Our study was designed shortly after the report of the infant in Mississippi who started antiretroviral treatment (ART) within 30 h of birth and who was able to maintain viral suppression off treatment for over two years. This case report was consistent with studies of ART started soon after primary HIV infection in adults that had observed that some individuals can achieve periods of post-treatment viral control when ART is withdrawn (remission). Taken together with knowledge from developmental biology of the neonatal immune profile, we hypothesized that starting and maintaining effective treatment in early life would lead to protection of critical immune processes and establishment of smaller viral reservoirs, leading to remission in a sizable minority (>20%). Adequate viral suppression and immune competence while ART is maintained is generally taken as minimum criteria for ART interruption trials. We conducted a PubMed search, with no language restrictions, including the following search terms in combination: HIV, neonate, newborn, children, perinatal, antiretroviral therapy, cure, remission, treatment interruption, to identify articles pertinent to our study through May 6, 2019.

Added value of this study

We started ART within 14 days of birth among 73 HIV-infected neonates identified within 48 h of birth at one hospital in Johannesburg, South Africa. Of those surviving on study, 52.5% attained and sustained viral load <50 copies/ml and half of these sustained CD4+ T-cell percentage >30%. Proportions achieving these virological and immunological endpoints were similar in the 46 infants who started ART <48 h to 27 infants starting 2–14 days.

Implications of all available evidence

Very early ART on its own, using the regimens available at the time our study was designed, is unlikely to be sufficient to achieve remission in a sizable minority of infants. The rare cases of perinatally-infected children who have been able to achieve prolonged periods of remission also suggest that early treatment may interact with host and viral factors to achieve remission but ART on its own may not be sufficient. Early initiation of more potent antiretroviral regimens, long-acting formulations and/or alternative interventions, such as broadly-neutralizing antibodies, need to be investigated to enable more rapid and sustained viral control and immune recovery as a stepping stone to achieve remission in perinatally-infected infants.

tantilizing possibility that similar results might be observed in other early treated infants [7]. ART started close to birth is not only close in time to acquisition of infection but is also being given during a developmentally-critical time period when the immune system is transitioning to a more mature form and is at its most quiescent [8,9]. We hypothesized that starting and maintaining effective treatment in early life would lead to protection of critical immune parameters and establishment of smaller viral reservoirs, leading to remission in a sizable minority (>20%).

To test the hypothesis, we designed an analytic treatment interruption (ATI) trial. Interruption of treatment is currently the only known approach to test whether periods of remission can be achieved. Here we describe the cohort of HIV-infected neonates identified within 48 h of birth, started on ART within 14 days and maintained on ART, that we intended as the target population for this trial. We describe the proportions of these early-treated, HIV-infected children who meet virologic and immunologic eligibility criteria for entry into the ATI trial.
ART was initiated based on results of the first round of diagnostic testing. To confirm diagnosis, a second blood sample was collected prior to the first ART dose. For those co-tested with Xpert, confirmatory testing was usually done on the same day as initial testing. The qualitative HIV-1 TNA diagnostic PCR was repeated and viral load (VL) done (quantitative HIV-1 RNA COBAS AmpliPrep/COBAS TaqMan HIV-1 test, version 2.0, Roche Molecular Systems, Inc., Branchburg, NJ). If subsequent tests did not confirm HIV diagnosis, infants were excluded from eligibility for the ATI trial and were managed based on the profile of their results.

ART was continued for a minimum of 104 weeks. Infants were followed on one of two protocols. The preferred protocol repeated VL tests at 1, 2 and 4 weeks of age, every 4 weeks to 24 weeks and then every 8 weeks to 104 weeks. The alternative protocol, repeated VL at 4, 8, 12, 16, and 24 weeks then every 12 weeks to 104 weeks. Diagnostic HIV PCR tests were repeated at 24, 48, 72 and 104 weeks. CD4+ T-cell count and percentage (TruCount Method, BD Biosciences, Germany) was measured at enrolment, 24, 48, 72 and 104 weeks. Complete blood counts were done at 4, 12, 24, 48, 72 and 104 weeks. Alanine aminotransferase (ALT) was measured at 16 weeks. Results were graded using the Division of AIDS toxicity tables [11]. All grade 3 and 4 abnormalities were repeated. A maternal blood sample was collected at enrolment for VL, CD4+ T-cell count and complete blood count. Antenatal information and maternal HIV treatment history was collected. Children were assessed for clinical progression at every visit and anthropometric data were collected.

![Flow diagram of screening to identify potential participants, their disposition prior to enrollment and follow-up after enrollment.](image-url)
2.4. Sample size

Power calculations determined the need for 40 children to enter the ATI trial. Initially, it was anticipated that 60 early-treated children would be sufficient to obtain this number based on the assumption that ~80% would become eligible for the interruption trial and ~85% of these would remain in care. As outcome data accumulated, it was clear that the cohorts of early-treated infants would need to be larger and enrolment continued. A DSMB review in May 2018 concluded that the numbers of cohorts of early-treated infants would need to be larger and enrolment remain in care. As outcome data accumulated, it was clear that the would become eligible for the interruption trial and ~85% of these would be sufficient size to enter the ATI trial as originally designed. Further enrolment into the early-treatment protocols ended shortly thereafter.

2.5. Outcomes

Here we report the proportions of children in the early-treated cohorts who met the primary virologic and immunologic endpoints for consideration for entry into the ATI trial. The primary virologic endpoint was VL <400 copies/ml by 24 weeks after ART initiation and <50 copies/ml by 48 weeks of age and no confirmed VL (i.e. two consecutive measurements) >50 copies/ml after suppression was attained. The primary immunologic endpoint was a CD4+ T-cell percentage >30% by 24 weeks which was sustained through follow-up.

| Characteristic | Total (N = 73) | Initiated ART <48 h (N = 46) | Initiated ART 2–14 days (N = 27) | p-value |
|----------------|---------------|-----------------------------|---------------------------------|---------|
| Sex, N (%)     |               |                             |                                 |         |
| Male           | 39 (53.4)     | 26 (56.5)                   | 13 (48.2)                       | 0.628   |
| Female         | 34 (46.6)     | 20 (43.5)                   | 14 (51.9)                       |         |
| Birth Weight (grams), Range | 905–4150 | 1860–4150                   | 905–3890                        |         |
| Birth Weight (grams), Mean (SD) | 2820 (602) | 2954 (477) | 2592 (723) | 0.026 |
| Birth Weight (grams), N (%) | 2500 | 29 (39.7) | 20 (43.5) | 9 (33.3) |
| ≥3000          | 28 (38.4)     | 20 (43.5)                   | 8 (29.6)                        |         |
| Gestational age by Ballard (weeks), N (%) | 37 weeks (term) | 63 (86.3) | 44 (95.6) | 19 (70.4) |
| Mode of delivery, N (%) | 37 weeks (pre-term) | 10 (13.7) | 2 (4.4) | 8 (29.6) |
| Vaginal        | 55 (75.3)     | 38 (82.6)                   | 17 (63.0)                       | 0.091   |
| Cesarean       | 18 (24.7)     | 8 (17.4)                    | 10 (37.0)                       |         |
| Ever breastfed, N (%) | Yes | 57 (78.1) | 39 (84.8) | 18 (66.7) |
| No             | 16 (21.9)     | 7 (15.2)                    | 9 (33.3)                        |         |
| Infant prophylaxis, N (%) | 34 (46.6) | 20 (43.5) | 14 (51.9) | 1 (3.7) |
| Maternal CD4+ T-cell count closest to birth (cells/mm³), Median (IQR) | 3272 (1474–2372) | 3191 (1474–2442) | 1875 (1182–2372) | 0.926 |
| Pre-treatment Viral load (copies/ml), Median (IQR) | 31,445 (5355–290,807) | 35,071 (5370–267,000) | 12,335 (1124–454,790) | 0.780 |
| Age when pre-treatment Viral load measured (days), Median (IQR) | 4150 (1015–8107) | 7 (1–1) | 7 (4–8) | <0.001 |
| Pre-treatment CD4+ T-cell count (cells/mm³), Median (IQR) | 1823 (1474–2372) | 1819 (1474–2442) | 1875 (1182–2372) | 0.926 |
| Pre-treatment CD4+ T-cell percentage (%), Median (IQR) | 40.6 (32.2–49.8) | 39.9 (32.2–48.8) | 47.4 (30.9–51.8) | 0.354 |
| Pre-treatment Viral load (copies/ml), N (%) | 400 | 9 (13.0) | 5 (10.9) | 4 (17.4) |
| 400–1000       | 4 (5.8)       | 3 (6.5)                     | 1 (4.4)                         | 0.271   |
| 1000–10,000    | 10 (14.5)     | 6 (13.0)                    | 4 (17.4)                        |         |
| 10,000–100,000 | 20 (29.0)     | 14 (30.4)                   | 6 (26.1)                        |         |
| 100,000–1,000,000 | 18 (26.1) | 15 (32.6) | 3 (13.0) |         |
| >1,000,000     | 8 (11.6)      | 3 (6.5)                     | 5 (21.7)                        |         |
| Pre-treatment CD4+ T-cell count (cells/mm³), Median (IQR) | 1823 (1474–2372) | 1819 (1474–2442) | 1875 (1182–2372) | 0.926 |
| Pre-treatment CD4+ T-cell percentage (%), Median (IQR) | 40.6 (32.2–49.8) | 39.9 (32.2–48.8) | 47.4 (30.9–51.8) | 0.354 |
| Pre-treatment Viral load (copies/ml), N (%) | 25 (severe) | 6 (9.8) | 3 (7.1) | 3 (15.8) |
| 25–30 (advanced) | 6 (9.8) | 6 (14.3) | 0 (0.0) |         |
| 30–35 (mild) | 9 (14.8) | 6 (14.3) | 3 (15.8) |         |
| >35 (none or not significant) | 40 (65.6) | 27 (64.3) | 13 (68.4) |         |
| Mother age (years), Mean (SD) | 28.4 (6.0) | 28.4 (5.7) | 28.4 (6.7) | 0.976 |
| Maternal antiretroviral therapy (ART) category, N (%) | 12 (16.4) | 6 (13.0) | 6 (22.2) | 0.276 |
| ART started during pregnancy, ≥12 weeks | 20 (27.4) | 15 (32.6) | 5 (18.5) |         |
| ART started during pregnancy, <12 weeks | 25 (34.3) | 14 (30.4) | 11 (40.7) |         |
| ART started during pregnancy, unknown time | 1 (1.4) | 0 (0.0) | 1 (3.7) |         |
| No ART until delivery | 15 (20.6) | 11 (23.9) | 4 (14.8) |         |
| Maternal ART regimen at delivery, N (%) | 56 (76.7) | 35 (76.1) | 21 (77.8) | 0.134 |
| Elavir-based | 2 (2.7) | 0 (0.0) | 2 (7.4) |         |
| No ART before delivery | 15 (20.6) | 11 (23.9) | 4 (14.8) |         |
| Maternal Viral load closest to birth (copies/ml), Median (IQR) | 38,459 (1760–104,538) | 31,433 (1292–104,538) | 50,400 (1760–125,515) | 0.547 |
| Maternal CD4+ T-cell count closest to birth (cells/mm³), Median (IQR) | 327 (207–567) | 357 (231–603) | 259 (128–480) | 0.097 |
| Maternal CD4+ T-cell count closest to birth (cells/mm³), N (%) | 200 | 17 (23.3) | 8 (17.4) | 9 (33.3) |
| 200–349 | 23 (31.5) | 15 (32.6) | 8 (29.6) |         |
| 350–499 | 13 (17.8) | 9 (19.6) | 4 (14.8) |         |
| >500 | 20 (27.4) | 14 (30.4) | 6 (22.2) |         |

Denominators are as shown.
Abbreviations: Antiretroviral treatment (ART), Standard deviation (SD), Inter-quartile range (IQR).
2.6. Statistical analysis

The analysis included children with confirmed HIV infection (first sample indicating HIV-infection collected < 48 h) and ART initiated within 14 days born between March 1, 2015 (start of prospective enrolment into the earliest study protocol) and September 30, 2017 (to allow at least 48 weeks of follow-up data on all children). Follow-up data accrued through September 2018 or truncated at 104 weeks of age were utilized. The primary analysis describes proportions meeting eligibility criteria for the ATI trial. Binomial proportion confidence intervals were reported under 0.05 significance level. If cell counts for proportions were > 10, Wald-type confidence intervals were reported; for counts ≤ 10, Clopper-Pearson estimation method was used. Secondary analyses compared proportions stratified by whether ART was initiated before or after 48 h of birth. Proportions were compared with Fisher’s exact tests and continuous variables were compared with t-tests, if normally-distributed, in which case the means were shown, and with Wilcoxon signed-rank tests otherwise, in which case the medians were shown. Statistical analyses were conducted in SAS version 9.4 (Cary, NC).

3. Results

Between March 1, 2015 and September 30, 2017, 7217 HIV-exposed neonates were tested with the routine diagnostic HIV PCR among which 4233 (59%) were co-tested with Xpert yielding 115 neonates with an initial positive or indeterminate result. From these, 79 confirmed infected infants were enrolled and 73 of these started ART within 14 days of birth (Fig. 1).

Among 46 infants who started ART < 48 h of birth, the screening blood sample was collected at a median of 13.0 h (IQR 7.1–24.0; range 0.9–60.0 h) after birth; in 66.7% of cases after having received a dose of nevirapine (median time of blood draw 4.8 h after prophylaxis [IQR 2.8–6.3; range 2.0–7.8 h]). All infants, except one, received at least one dose of nevirapine prophylaxis and were prescribed daily nevirapine on discharge; four of these were prescribed zidovudine as well. Three infants were co-tested with Xpert but did not initiate ART on-site at the time and 24 infants had not been co-tested with Xpert on-site and had to be recalled. One of the 27 infants had an indeterminate result on initial screening (Fig. 1). The median baseline VL was 12,335 copies/ml (IQR 1124–454,790; range 102–863,000 copies/ml) (Table 1).

Among 46 neonates initiating ART < 48 h of birth, nevirapine, lamivudine and zidovudine was initiated at a median of 24 h after birth (IQR 18.1–30.5; range 8.1–48.0 h). ART initiation was a median of 6.4 h (IQR 5.6–22.2; range 4.9–34.8 h) after the blood draw to identify infection. Among 27 neonates initiating ART 2–14 days of birth, the median age at ART initiation was 6 days (IQR 4–8 days). All, except one, initiated treatment with nevirapine, lamivudine and zidovudine (Table 2).

Nevirapine was changed to lopinavir-ritonavir at a median of 30 days (IQR 18–36; range 14–71 days) in neonates who started ART < 48 h and 33 days (IQR 28–57; range 18–132 days) in neonates who received a dose of nevirapine as prophylaxis (median time of blood draw 3.3 h after prophylaxis [IQR 2.5–4.3; range 0.5–9.0 h]). The routine diagnostic PCR run on the same sample was positive in 41 and indeterminate in five (all turned positive on subsequent PCR tests). The second sample was drawn a median of 5.9 h (IQR 5.0–22.6) after the first sample a median of 22.6 h (IQR 18.3–30.7) after birth. The median baseline VL was 35,071 copies/ml (IQR 5370–267,000; range 60–2445,950 copies/ml) (Table 1).

Among 27 infants who started ART > 48 h and < 14 days of age (2–14 days), the screening blood sample was collected a median of 13.0 h (IQR 7.1–24.0; range 0.9–60.0 h) after birth; in 66.7% of cases after having received a dose of nevirapine (median time of blood draw 4.8 h after prophylaxis [IQR 2.8–6.3; range 2.0–7.8 h]). All infants, except one, received at least one dose of nevirapine prophylaxis and were prescribed daily nevirapine on discharge; four of these were prescribed zidovudine as well. Three infants were co-tested with Xpert but did not initiate ART on-site at the time and 24 infants had not been co-tested with Xpert on-site and had to be recalled. One of the 27 infants had an indeterminate result on initial screening (Fig. 1). The median baseline VL was 12,335 copies/ml (IQR 1124–454,790; range 102–863,000 copies/ml) (Table 1).

Between March 1, 2015 and September 30, 2017, 7217 HIV-exposed neonates were tested with the routine diagnostic HIV PCR among which 4233 (59%) were co-tested with Xpert yielding 115 neonates with an initial positive or indeterminate result. From these, 79 confirmed infected infants were enrolled and 73 of these started ART within 14 days of birth (Fig. 1).

Among 46 infants who started ART < 48 h of birth, the screening blood sample was collected at a median of 13.0 h (interquartile range [IQR] 9.0–15.8; range 2.0–31.1 h) after birth; all but one after having

### Table 2

| Characteristic | Total (N = 73) | Initiated ART < 48 h (N = 46) | Initiated ART 2–14 days (N = 27) | p-value |
|----------------|---------------|-------------------------------|---------------------------------|---------|
| Age at ART start in days of life | | | | |
| Range | 0–14 | 0–2 | 2–14 | |
| Mean (SD) | 3.0 (3.2) | 1.0 (0.5) | 6.4 (2.9) | |
| Median (IQR) | 1.0 (1.0–5.0) | 1.0 (1.0–1.0) | 6.0 (4.0–8.0) | |
| Age at ART start in hours since birth | | | | |
| Range | 8.1–48.0 | | | |
| Mean (SD) | 25.1 (10.5) | | | |
| Median (IQR) | 24.0 (18.1–30.5) | | | |
| Initial regimen, N [%] | | | | |
| Nevirapine/Lamivudine/Zidovudine<sup>1</sup> | 72 (98.6) | 46 (100.0) | 26 (96.3) | 0.370 |
| Lopinavir-ritonavir/Lamivudine/Zidovudine | 1 (1.4) | 0 (0.0) | 1 (3.7) | |
| Lopinavir-ritonavir status in 48 weeks of follow-up, N [%] | | | | |
| Lost before age to switch | 3 (4.1) | 3 (6.5) | 0 (0.0) | 0.410 |
| Initiated on lopinavir-ritonavir | 1 (1.4) | 0 (0.0) | 1 (3.7) | |
| Switched to lopinavir-ritonavir in first 28 days | 34 (46.6) | 21 (45.7) | 13 (48.2) | |
| Switched to lopinavir-ritonavir after 28 days | 35 (48.5) | 22 (47.8) | 13 (48.2) | |
| If switched in first 28 days, age switched (days) | | | | |
| Range | 14–32 | 14–30 | 18–32 | |
| Median (IQR) | 21 (17–29) | 18 (16–24) | 28 (21–30) | |
| If switched after 28 days, age switched (days) | | | | |
| Range | 29–132 | 29–71 | 34–132 | |
| Median (IQR) | 44 (33–61) | 35 (31–47) | 57 (45–80) | |
| Lopinavir-ritonavir switch in relation to estimated post-menstrual age (PMA), N [%] | | | | |
| No follow-up | 3 (4.1) | 3 (6.5) | 0 (0.0) | 0.491 |
| Initiated on lopinavir-ritonavir | 1 (1.4) | 0 (0.0) | 1 (3.7) | |
| Before 42 weeks PMA | 2 (2.7) | 1 (2.2) | 1 (3.7) | |
| 42 weeks PMA | 25 (34.3) | 10 (21.3) | 9 (33.3) | |
| After 42 weeks PMA | 42 (57.5) | 26 (56.5) | 16 (59.3) | |
| Zidovudine changed to abacavir in 48 weeks of follow-up, N | 33 | 21 | 12 | |
| If switched to abacavir, median age switched in days (IQR) | 169 (125–198) | 149 (125–186) | 182 (130–260) | |
| Zidovudine changed to stavudine in 48 weeks of follow-up, N | 4 | 2 | 2 | |

<sup>1</sup> Exact hours known in 41/45. In the other 4 known to be < 48 h but exact numbers of hours not recorded.

<sup>2</sup> One child received one dose of lopinavir-ritonavir in error at the start of treatment but then continued with nevirapine.

Abbreviations: Antiretroviral treatment (ART), Standard deviation (SD), Inter-quartile range (IQR), Post-menstrual age (PMA).
started ART 2–14 days of life. In two infants this was sooner than the recommended 42 weeks, 34.3% at 42 weeks, and 57.5% > 42 weeks (Table 2). Two children with Grade 3 neutropenia substituted stavudine for zidovudine at 14 and 74 days of age (both in the 2–14 days group). One child with Grade 3 low hemoglobin substituted stavudine for zidovudine at 66 days of age (2–14 days group). All other Grade 3 and 4 laboratory abnormalities resolved on repeat testing without a change in regimen (Supplementary Table 1).

Fig. 2. Viral load and other characteristics of the three study participants who died on study. P indicates the timing of positive diagnostic PCR tests.

A:
Maternal ART started < 12 weeks before birth
Cesarean delivery
Male
Birthweight 2905 grams
Started ART 28 hours after birth
No breastfeeding
Age at death 61 days
Cause of death: Unknown. History of diarrhea and sepsis

B:
No Maternal ART
Vaginal delivery
Male
Birthweight 3710 grams
Started ART <48 hours after birth
Breastfed
Age at death 43 days
Cause of death: Pneumonia/Hypoxic encephalopathy

C:
No Maternal ART
Cesarean delivery
Male
Birthweight 2405 grams
Started ART 44 hours after birth
Breastfed
Age at death 89 days
Cause of death: Unknown. Febrile illness
Table 3
Proportion of 73 HIV-infected infants starting antiretroviral therapy (ART) within 14 days of birth who met virologic and immunologic eligibility criteria for the analytic treatment interruption study by age at ART initiation.

| Characteristic                             | Total Initiated ART | Initiated ART <48 h | Initiated ART 2 to 14 days | p-value |
|--------------------------------------------|---------------------|---------------------|----------------------------|---------|
|                                            | N  | % of total (95% CI) | % of those in follow-up (95% CI) N = 61 | N  | % of total (95% CI) | % of those in follow-up (95% CI) N = 35 | N  | % of total (95% CI) | % of those in follow-up (95% CI) N = 26 |         |
| Deaths                                     | 3  | 4.1 (0.9 – 11.5)   |                           | 3  | 6.5 (1.4 – 17.9)   |                               | 0  | 0 (0 – 12.8)       |                                    | 0.291*  |
| Loss to Follow-up before 48 weeks          | 9  | 12.3 (5.8 – 22.1)  |                           | 8  | 17.4 (7.8 – 31.4)  |                               | 1  | 3.7 (0 – 19.0)     |                                    | 0.141*  |
| **Meets virologic criteria**               |     |                     |                           |     |                     |                               |     |                     |                                    |         |
| No: VL not <50 copies/ml by 48 weeks       | 15 | 20.6                |                           | 6  | 13.0                |                               | 9  | 33.3               |                                    |         |
| No: VL <50 copies/ml. achieved but confirmed rebound >50 copies/ml | 14 | 19.2                |                           | 11 | 23.9                |                               | 3  | 11.1               |                                    |         |
| Yes: VL achieved and sustained <50 copies/ml | 32 | 43.8 (32.5–55.2)   | 52.5 (40.0–65.0)         | 18 | 39.1 (25.0–53.2)   | 51.4 (34.9–68.0)             | 14 | 51.9 (33.0–70.7)   | 53.9 (34.7–73.0)                  | 0.335*  |
| **Meets immunologic criteria**             |     |                     |                           |     |                     |                               |     |                     |                                    |         |
| No                                         | 35 |                     |                           | 19 |                     |                               | 16 |                     |                                    |         |
| Yes: CD4+ T-cell% sustained >30%           | 26 | 42.6 (30.2–55.0)   |                           | 16 | 45.7 (29.2–62.2)   |                               | 10 | 38.5 (19.8–57.2)   |                                    | 0.610*  |
| **Virologic and immunologic criteria**     |     |                     |                           |     |                     |                               |     |                     |                                    |         |
| Meets both                                 | 16 | 21.9 (12.4 – 31.4) | 26.2 (15.2 – 37.3)        | 9  | 19.6 (9.4 – 33.9)  | 25.7 (12.5 – 43.3)            | 7  | 25.9 (11.1 – 46.3) | 26.9 (11.6 – 47.8)                | 1.000*  |
| Meets virologic but not immunologic        | 16 | 21.9                | 26.2                      | 9  | 19.6                | 25.7                          | 7  | 25.9                | 26.9                          | 0.567*  |
| Meets immunologic but not virologic        | 10 | 13.7                | 16.4                      | 7  | 15.2                | 20.0                          | 3  | 11.1                | 11.5                          |         |
| Meets neither                              | 19 | 26.0                | 31.2                      | 10 | 21.7                | 28.6                          | 9  | 33.3                | 34.6                          |         |

* Fisher Exact test results among all children (N = 73) comparing proportions between two ART initiation groups.
† Fisher Exact test results among surviving children (N = 61) comparing proportions between two ART initiation groups

Abbreviations: Antiretroviral treatment (ART), Confidence intervals (CI), Viral load (VL).
Three infants died at 43, 61 and 89 days of age. These three infants had started ART at <48 (exact time unknown), 28, and 44 h respectively. All three were male born at term, one was low birth weight, and two were cesarian section deliveries. All had high baseline VL and had transitioned to the lopinavir-ritonavir regimen, but none attained viral suppression before death (Fig. 2).

Nine infants (12.3%) were lost to follow-up before 48 weeks of age: 8/46 who started ART <48 h and 1/27 who started 2–14 days. Taking into account the three deaths, survival on study was lower in those who started <48 h (35/46) compared to 2–14 days (26/27) (p=0.026). Overall 63.0% (46/73) of children achieved <50 copies/ml by 48 weeks of age (45/46 achieved <400 copies/ml by 24 weeks) and 32/73 (43.8%) sustained VL <50 copies/ml without a confirmed measurement above this threshold to 48 weeks or last follow-up visit (32/61 [52.5%] of those remaining in follow-up). Most infants (56/61; 91.8%) had CD4+ T-cell percentage ≥30% by 48 weeks compared to 2–14 days (35/46) (p=0.007), which was also significantly higher in those who met virologic criteria (17.1% [6.6–33.7]) versus those who did not (14.7% [4.6–46.3]) (p=0.05).

Non-eligible children had started ART at 48 (exact time unknown), 28, and 44 h respectively. Lack of apparent benefit of very early ART on viral suppression may, at first glance, appear to contradict previous studies [13]. However, prior studies considered the timing of ART in much wider age bands than examined here [14]. Moreover, maternal ART was in place before birth in almost 80% of the participants, and all infants received nevirapine prophylaxis at birth regardless of the time of ART initiation. Drug resistance is unlikely to explain the findings as all were switched to a ritonavir-raltegravir-based regimen. While studies of acutely infected adults have almost 80% of the participants, and all infants received nevirapine prophylaxis at birth regardless of the time of ART initiation. Drug resistance is unlikely to explain the findings as all were switched to a ritonavir-raltegravir-based regimen. While studies of acutely infected adults have

### Table 4
Proportion of 61 HIV-infected infants starting antiretroviral therapy (ART) within 14 days of birth surviving in follow-up who met more stringent virologic endpoints by age at ART initiation.

| Characteristic                        | Total | Initiated ART <48 h | Initiated ART 2 to 14 days | p-value |
|---------------------------------------|-------|---------------------|---------------------------|---------|
|                                       | N     | % of 61 in follow-up(95% CI) | n/N (%) in sub-group | N     | % of 35 in follow-up(95% CI) | n/N (%) in sub-group | N     | % of 26 in follow-up(95% CI) | n/N (%) in sub-group |         |
| Did not achieve “Target not detected” (TND) by 48 weeks | 25 | 41.0 | 13 | 37.1 | 12 | 46.2 |         |
| Achieved TND then any rebound          | 26 | 42.6 | 16 | 45.7 | 10 | 38.5 |         |
| Sustained TND                          | 10 | 16.4 (8.2–28.1) | 6 | 17.1 (6.6–33.7) | 4 | 11.1 (4.4–34.9) | 1.00 |
| Ever PCR negative                      | 14 | 23.0 (13.2–35.5) | 10 | 28.6 (14.6–46.3) | 4 | 15.4 (4.4–34.9) | 0.357 |
| Non-transient PCR negative             | 9 | 14.8 (7.0–26.2) | 6 | 17.1 (6.6–33.7) | 3 | 11.5 (2.5–30.2) | 0.720 |
| Non-transient PCR negative in achieved & sustained <50 copies/ml group | 9 | 9/32 (28.1) | 6 | 6/18 (33.3) | 3 | 3/14 (21.4) |         |
| Non-transient PCR negative in achieved & sustained TND group | 5 | 5/10 (50.0) | 4 | 4/6 (66.7) | 1 | 1/4 (25.0) |         |

**Abbreviations:** Antiretroviral treatment (ART), Confidence intervals (CI), Target not detected (TND), polymerase chain reaction (PCR).
Fig. 3. Individual plots of viral load and CD4+ T-cell response to antiretroviral therapy of ten children in the study. P indicates the timing of positive, I indeterminate and N negative diagnostic PCR tests. 

A: Achieved and sustained target not detected and CD4% > 30% (3/61) 

B: Achieved and sustained target not detected and CD4% not > 30% (7/61) 

C: Achieved and sustained < 50 copies/ml and CD4% > 30% (13/61) 

D: Achieved and sustained < 50 copies/ml and CD4% not > 30% (9/61) 

E: Achieved < 50 copies/ml but not sustained and CD4% > 30% (6/61)
participants’ caregivers live in impoverished economic circumstances with complex social problems and experience a high degree of HIV-related stigma. This social profile is likely representative of transmitters in an era of high maternal ART coverage. Moreover, there are major practical difficulties of sustaining adherence with twice-daily, poorly-palatable liquids for infants.
Mortality was only observed in those initiating ART <48 h of birth, however sample size was small and attrition before ART initiation may counter-balance this apparent excess.

Almost half of the early-treated neonates did not attain the protocol-specified immunologic criteria required for entry into the ATI trial; even among those who attained the protocol-specified virologic criteria. Discordance in virologic and immunologic response to ART in children has been described [17–19]. Reported frequency of discordance in other cohorts is lower than we observed but differences in definitions makes comparisons difficult [17–19]. Unless liberal cut-offs on CD4+ T-cell count had been used, an immunologic endpoint based on count rather than percentage would not have been meaningfully different.

Two recent case reports of long-term remission are of interest, although, in both cases ART was not started in the neonatal period as in our study [20,21]. Given that critical host and viral factors very likely contributed to remission in these cases, they support the findings from our study that early treatment on its own may not be sufficient. Moreover, in the one case, where a denominator could be ascertained, remission was attained in only one child out of more than 200 children who initiated ART reasonably early and interrupted ART [21]. The apparent rarity of post-treatment control supports our conclusion that very early ART on its own, with routinely-available regimens, is unlikely to lead to remission in a sizable minority of early-treated infants.

The major limitation of our study is that early antiretroviral therapy for infants is entirely reliant on maternal adherence. Although we supported adherence through an experienced, multi-disciplinary team, the challenges of sustained adherence likely contribute to the results. A second limitation is that we advised use of antiretroviral regimens and formulations that we considered best for this age group available in this setting at the time the study was designed. Other alternatives, including raltegravir which was only approved for use in this age group when the study was almost complete [22], may be more effective. Although raltegravir addresses palatability issues associated with lopinavir-ritonavir, it adds a layer of complexity around preparation for neonates and young infants. Thus raltegravir offers only minor respite from the adherence challenges which likely drive the poor response to early ART. Further research on possible long-acting formulations is needed. The virological and immunological outcomes occurred less frequently than expected necessitating longer recruitment. However, there were no other deviations from the original protocols.

Decisions about how quickly to start ART should be based on expectations about the capacity of ART to achieve standard clinical outcomes i.e. sustained viral suppression, and reduced morbidity and mortality rather than with the expectation that remission might be attained. Early initiation of more potent antiretroviral regimens or long-acting formulations that can address adherence challenges and/or alternative interventions, such as broadly neutralizing antibodies, need to be investigated to enable more rapid and sustained viral control and immune recovery as a stepping stone to achieving remission in this population.

Declaration of Competing Interest

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Representatives from the NIH were involved in the design of the study, in oversight of its implementation and contributed to the writing of this manuscript. None of the other funders played any role in the writing of this manuscript or the decision to submit for publication.

Supplementary materials

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