Risk of triple-class virological failure in children with HIV: a retrospective cohort study

The Pursuing Later Treatment Options II (PLATO II) project team for the Collaboration of Observational HIV Epidemiological Research Europe (COHERE)*

Summary

Background In adults with HIV treated with antiretroviral drug regimens from within the three original drug classes (nucleoside or nucleotide reverse transcriptase inhibitors [NRTIs], non-NRTIs [NNRTIs], and protease inhibitors), virological failure occurs slowly, suggesting that long-term virological suppression can be achieved in most people, even in areas where access is restricted to drugs from these classes. It is unclear whether this is the case for children, the group who will need to maintain viral suppression for longest. We aimed to determine the rate and predictors of triple-class virological failure to the three original drugs classes in children.

Methods In the Collaboration of Observational HIV Epidemiological Research Europe, the rate of triple-class virological failure was studied in children infected perinatally with HIV who were aged less than 16 years, starting antiretroviral therapy (ART) with three or more drugs, between 1998 and 2008. We used Kaplan-Meier and Cox regression methods to investigate the risk and predictors of triple-class virological failure after ART initiation.

Findings Of 1007 children followed up for a median of 4.2 (IQR 2.4–6.5) years, 237 (24%) were triple-class exposed and 105 (10%) had triple-class virological failure, of whom 29 never had a viral-load measurement less than 500 copies per mL. Incidence of triple-class virological failure after ART initiation increased with time, and risk by 5 years after ART initiation was 12.0% (95% CI 9.4–14.6). In multivariate analysis, older age at ART initiation was associated with increased risk of failure (p=0.02). Of 686 children starting ART with NRTIs and either a NNRTI or ritonavir-boosted protease inhibitor, the rate of failure was higher than in adults with heterosexually transmitted HIV (hazard ratio 2.2 [95% CI 1.6–3.0, p<0.0001]).

Interpretation Findings highlight the challenges of attaining long-term viral suppression in children who will be taking life-long ART. Early identification of children not responding to ART, adherence support, particularly for children and adolescents aged 13 years or older starting ART, and ART simplification strategies are all needed to attain and sustain virological suppression.

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Introduction

Antiretroviral therapy (ART) has strikingly improved the prognosis for children with HIV, greatly reducing morbidity and mortality.2–4 However, a major challenge for treatment of these children is to minimise virological failure and development of drug resistance, so that treatment options continue to be available through adolescence and adulthood.

During the past decade, paediatric ART guidelines recommended an age-related CD4 percentage or count threshold for initiation of ART in infants and children.4 However, in view of results from the Children with HIV Early Antiretroviral Therapy (CHER) trial and other evidence,17 paediatric ART guidelines now unanimously advocate initiation of ART early in infancy because of the high risk of disease progression.17–20 Even if ART is not started early, vertically infected children face many more years of ART than do adults. A serious challenge for children with HIV, as for any chronic illness, is maintaining long-term adherence to treatment regimens, and thus virological suppression and prevention of emergence of resistance.15–19 However, experience with these children suggests that, with present treatment strategies, adherence rates are frequently less than optimum,20–26 which combined with the increased potential for inadequate drug dosing,27,28 contribute to appreciable risk of children acquiring multidrug-resistant HIV before adulthood.

As for adults, most paediatric ART regimens include drugs from one or more of the original three ART classes: nucleoside or nucleotide reverse transcriptase inhibitors (NRTI), non-NRTIs (NNRTI), and protease inhibitors. The availability of drugs from the new classes (integrase and entry inhibitors) remains limited by the lag in availability of appropriate formulations and pharmacokinetic data for children, and, in developing countries, high drug costs.3 Virological failure of the three original drug classes during childhood will severely limit future treatment options; therefore, the rate of triple-class virological failure should be monitored, to estimate the number of children transferring to adult care in probable need of treatment with new drugs.

We aimed to determine the rate and predictors of triple-class virological failure to the three original drug...
classes in children within the Collaboration of Observational HIV Epidemiological Research Europe (COHERE). The study forms part of the Pursuing Later Treatment Options II (PLATO II) project; an analysis of triple-class virological failure in adults from this project has been published.

**Methods**

**Study design**

14 cohorts with paediatric data in COHERE submitted data in a standardised format to one of two regional coordinating centres, where error checks were done before data were merged. Children appearing in more than one cohort were identified, and duplicate records removed. This analysis (of data merged in 2008) was restricted to children perinatally infected with HIV aged less than 16 years who started ART from 1998 onwards with an initial regimen of two or more NRTIs and either a NNRTI or a protease inhibitor (ritonavir-boosted or unboosted), or three NRTIs only (children exposed to ART for the prevention of mother-to-child transmission before starting one of the above regimens were included). Unlike in the adult analysis, children starting initial regimens with unboosted protease inhibitors, more than two NRTIs with a NNRTI or protease inhibitor, and three NRTIs only were included to mirror the regimens commonly prescribed to children in the past decade. A second analysis was done to mirror the adult analysis. Children were followed up from the start of ART to their last viral-load measurement. Because the definition of virological failure used in this study required 4 months of use of a drug, children were included only if they had at least 4 months (122 days) of follow-up. Time spent off ART (after ART had been started) was included as follow-up. Children were said to have interrupted ART if they stopped all antiretrovirals.

**Data extraction**

The main outcome in this study was triple-class virological failure, defined as virological failure to at least two NRTIs, one NNRTI, and one protease inhibitor. We defined virological failure of a drug as one viral-load measurement of greater than 500 copies per mL after at least 4 months of continuous use of that drug, regardless of concomitant use of other drugs in that period. We did sensitivity analyses using two variations of this definition: first, 6 months instead of 4 months of continuous drug use; second, viral load above 500 copies per mL after 4 months continuous use had to be confirmed by a second viral load value greater than 500 copies per mL while still on or followed by discontinuation of the drug.

To enable comparisons of the rate of triple-class virological failure in children with that of adults, children starting initial regimens of two or three NRTIs and either a NNRTI or a ritonavir-boosted protease inhibitor (ie, those for whom the initial ART regimen included an unboosted protease inhibitor, or three NRTIs, were excluded).

![Figure 1: Inclusion criteria for initial regimens and definition of triple-class virological failure in the main analysis and the comparison of children with adults.](http://www.cphiv.dk/COHERE/StudyDocuments/tabid/298/Default.aspx)

**Table:**

| Number of children (n=1007) | Median follow-up person-years |
|-----------------------------|------------------------------|
| **Sex**                      |                              |
| Boy                         | 510 (51%)                    | 3.9 |
| Girl                        | 497 (50%)                    | 3.6 |
| **Age at start of ART (years)**|                             |
| <2                          | 350 (35%)                    | 4.1 |
| 2-4                         | 202 (20%)                    | 4.3 |
| 5-9                         | 265 (26%)                    | 4.0 |
| 10-15                       | 190 (19%)                    | 2.9 |
| Median (IQR)                | 4.2 (0.9-8.5)                |     |
| **Year of ART initiation**  |                              |
| 1998-2000                   | 368 (37%)                    | 6.8 |
| 2001-2003                   | 363 (36%)                    | 3.9 |
| 2004-2007                   | 276 (27%)                    | 1.7 |
| **Previous ART exposure for prevention of mother-to-child transmission** |                      |
| No                          | 937 (93%)                    | 3.9 |
| Yes                         | 70 (7%)                      | 3.1 |
| **Initial regimen**         |                              |
| NNRTI + 2 NRTIs             | 467 (46%)                    | 3.0 |
| NNRTI + 3 NRTIs             | 93 (9%)                      | 3.9 |
| Ritonavir-boosted protease inhibitor + 2 or 3 NRTIs | 126 (13%) | 3.0 |
| Unboosted protease inhibitor + 2 or 3 NRTIs | 270 (27%) | 6.7 |
| 3 NRTIs                     | 51 (5%)                      | 4.8 |

(Continues on next page)
Comparisons were made with adults heterosexually infected with HIV in COHERE (ie, excluding homosexual men), because this is the adult group that children with HIV are probably most similar to in terms of socioeconomic and migration status, and health seeking behaviour. Figure 1 shows the two inclusion criteria for initial ART regimens and definitions of triple-class virological failure for the main analysis and the comparison of children with adults.

Statistical analysis

We used Kaplan-Meier and Cox regression methods to investigate the risk of triple-class virological failure after starting ART, and to compare the rate of failure in children with that in adults in the PLATO II project. Possible predictors of failure at the time of ART initiation were sex, age, year of ART initiation, initial drug regimen, previous ART exposure for prevention of mother-to-child transmission, AIDS diagnosis before ART, CD4 percentages and viral load (within 6 months before ART initiation). Analyses were done with SAS software (version 9.1).

For children with triple-class virological failure who had failed a ritonavir-boosted protease inhibitor, we examined data from resistance tests done as part of routine care after the start of ART and up to 4 months after triple-class virological failure for a subset of cohorts where data were available. Major mutations from the International AIDS Society USA, 2008, mutation list were used to define resistance. Thymidine-associated mutations in HIV reverse transcriptase were defined as M41L, D67N, K70R, L210W, T215Y, T215F, K219Q, and K219E.

Role of the funding source

The sponsors of the study had no role in the study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results

A total of 1007 children were identified in the 14 COHERE cohorts. The UK and Ireland, Spanish, Dutch, and French cohorts contributed more than 90% of children, with a smaller proportion from Denmark, Italy, Belgium, and the European Collaborative Study. Table 1 shows the characteristics of children at the time of ART initiation. Just over a third of children initiated ART in each of the calendar periods 1998–2000 and 2001–03, and a quarter in 2004–07. Fewer than 10% had been exposed to ART as part of prevention of mother-to-child transmission; 55 were exposed to zidovudine only, and 4 to NNRTIs. About 20% had a previous AIDS diagnosis.

560 (56%) children initiated ART on a NNRTI-based regimen—232 (41%) on efavirenz and 328 (59%) on nevirapine. Most children starting ART with a ritonavir-boosted protease inhibitor did so with lopinavir (94%; 118 of 126).

Children were followed up for a median of 4.2 (IQR 2.4–6.5) years and 932 (93%) of 1007 for at least 12 months, with a median of 90 (56–112) days between viral-load measurements. 258 (26%) children interrupted ART at some time, of whom 158 (61%) restarted ART; most follow-up (93%) was of children on ART. Half

ART=antiretroviral therapy. IQR=inter-quartile range. NRTI=nucleoside or nucleotide reverse transcriptase inhibitors. NNRTI=non-NRTI. *Antiretroviral therapy (ART) for prevention of mother-to-child transmission was defined as one or two antiretrovirals initiated within 7 days of birth and stopped up to 60 days after, or three or more antiretrovirals initiated within 7 days of birth and stopped up to 60 days after, and then no antiretrovirals for at least 90 days.

Table 1: Characteristics of children at the time of antiretroviral therapy initiation

| Characteristics | Number of children (n=1007) | Median follow-up person-years |
|-----------------|-----------------------------|-------------------------------|
| AIDS diagnosis before ART initiation | | |
| No | 793 (79%) | 3.7 |
| Yes | 214 (21%) | 4.1 |
| CD4 percentage at ART initiation | | |
| 0–9% | 193 (19%) | 3.8 |
| 10–19% | 254 (25%) | 3.2 |
| ≥20% | 236 (23%) | 4.3 |
| Unknown | 324 (32%) | 4.0 |
| Median (IQR) | 15.0 (9–24) | |
| Viral load at ART initiation (log10 copies per mL) | | |
| 0–4.49 | 181 (18%) | 3.3 |
| 4.50–5.49 | 334 (33%) | 3.4 |
| ≥5.50 | 258 (26%) | 4.2 |
| Unknown | 234 (23%) | 4.1 |
| Median (IQR) | 5.1 (4.6–5.7) | |

Figure 2: Incidence per 100 person-years (95% CI) of triple-class virological failure in children with HIV by duration of antiretroviral therapy

*At end of year 9.
(518 [51%] of 1007) had first-line virological failure, with about 58% (95% CI 54–62) failure by 5 years after initiation of ART. Most children (470 [91%] of 518) had not started a third drug class at first virological failure; 206 (44%) of 470 started a third drug class before the end of follow-up, with 55% (50–61) doing so by 5 years after first virological failure.

By the end of follow-up, 335 (33%) of 1007 children had started a third drug class, of whom 105 (10% overall and 44% of those with triple-class exposure) had developed triple-class virological failure. Nine (1%) died without such treatment failure. Incidence of triple-class virological failure during each year after ART initiation increased sharply in the first 2 years, from 0·5 per 100 person-years (0·2–1·2) in the first year after ART initiation to 2·6 per 100 person-years (1·5–3·6) in the second year, and then rose more gradually up to the sixth year (4·0 per 100 person-years [2·1–7·0]; figure 2). By 5 years after ART initiation the estimated cumulative proportion of children who had triple-class virological failure was 12·0% (9·4–14·6), and by 8 years it was 20·3% (15·9–24·7). Redefinition of virological failure to 6 months of continuous drug use rather than 4 months gave an estimated cumulative proportion of children with triple-class virological failure by 5 years of 9·4% (7·1–11·8%), and modification of the definition to require confirmation of virological failure gave a 5-year proportion of 9·5% (7·2–11·8).

Older age (10–15 years) at the time of ART initiation was associated with an increased risk of failure (table 2).

**Articles**

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| Univariate analyses | Multivariate analyses |
|---------------------|----------------------|
|                     | Hazard ratio | 95% CI | p   | Hazard ratio | 95% CI | p   |
| **Sex**             |              |       |    |              |       |    |
| Boy                 | 1.0          |       | 0.96| 1.0          |       | 0.92|
| Girl                | 1.0          | 0.7–1.5|    | 1.0          | 0.7–1.5|    |
| **Age at start of ART (years)** |              |       |    |              |       |    |
| <2                  | 1.2          | 0.7–2.0| 0.03| 1.2          | 0.7–2.0| 0.02|
| 2–4                 | 0.9          | 0.5–1.7|    | 0.9          | 0.5–1.7|    |
| 5–9                 | 1.0          |       |    | 1.0          |       |    |
| 10–15               | 2.1          | 1.2–3.7|    | 2.3          | 1.2–4.1|    |
| **Year of ART initiation** |              |       |    |              |       |    |
| 1998–2000           | 1.0          |       | 0.46| 1.0          |       | 0.83|
| 2001–2003           | 0.7          | 0.5–1.2|    | 0.9          | 0.5–1.5|    |
| 2004–2007           | 0.9          | 0.4–2.0|    | 1.0          | 0.4–2.4|    |
| **Initial regimen**  |              |       |    |              |       |    |
| NNRTI+2 NRTIs       | 1.0          |       | 0.45| 1.0          |       | 0.24|
| NNRTI+3 NRTIs       | 0.7          | 0.3–1.6|    | 0.9          | 0.3–1.6|    |
| Ritonavir-boosted protease inhibitor + 2 or 3 NRTIs | 0.5 | 0.2–1.4|    | 0.4 | 0.2–1.3|    |
| Unboosted protease inhibitor + 2 or 3 NRTIs | 1.1 | 0.7–1.7|    | 1.1 | 0.7–1.7|    |
| 3 NRTIs             | 0.8          | 0.3–2.1|    | 0.8          | 0.3–2.0|    |
| **Previous ART exposure for prevention of mother-to-child transmission** |              |       |    |              |       |    |
| No or unknown       | 1.0          |       | 0.91| 1.0          |       | 0.64|
| Yes                 | 1.0          | 0.4–2.3|    | 1.3          | 0.5–3.4|    |
| **AIDS diagnosis before ART initiation** |              |       |    |              |       |    |
| No                  | 1.0          |       | 0.18| 1.0          |       | 0.12|
| Yes                 | 1.3          | 0.9–2.1|    | 1.4          | 0.9–2.3|    |
| **CD4 percentage at ART initiation** |              |       |    |              |       |    |
| 0–9%                | 1.0          |       | 0.61| 1.0          |       | 0.73|
| 10–19%              | 0.9          | 0.5–1.7|    | 1.0          | 0.6–1.9|    |
| ≥20%                | 0.8          | 0.4–1.5|    | 0.9          | 0.5–1.8|    |
| Unknown             | 1.1          | 0.7–1.9|    | 1.3          | 0.7–2.4|    |
| **Viral load at ART initiation (log10 copies per mL)** |              |       |    |              |       |    |
| 0–4·49              | 1.65         | 0.3–1.2| 0.58| 0.5          | 0.3–1.0| 0.24|
| 4·50–5·49           | 1.0          |       | 1.0 |    | 1.0          |       |    |
| ≥5·50               | 1.0          | 0.6–1.6|    | 0.9          | 0.5–1.6|    |
| Unknown             | 1.0          | 0.6–1.6|    | 0.7          | 0.4–1.3|    |

ART=antiretroviral therapy. NRTI=nucleoside or nucleotide reverse transcriptase inhibitors. NNRTI=non-NRTI.

Table 2: Predictors of triple class virological failure
Children with a previous AIDS diagnosis had a nonsignificant trend towards an increased risk of triple-class virological failure (table 2). We recorded no significant difference in the risk of failure by sex, year of ART initiation, type of initial ART regimen, previous ART exposure for prevention of mother-to-child transmission, or CD4 percentage or viral load at ART initiation.

Of the children with triple-class virological failure, 29 of 105 never had a viral load measurement of less than 500 copies per mL. Compared with the 76 who had at least one viral load measurement less than 500 copies per mL before developing treatment failure, these children started ART at a younger age (median 1·8 [IQR 0·5–4·4] vs 6·1 [1·2–10·1] years; p=0·008) and were triple-class exposed more quickly after starting ART (2·1 [1·4–3·8] vs 3·3 [1·9–4·6] years; p=0·018).

Of the 686 children starting ART with an initial regimen of two or three NRTIs and either a NNRTI or a ritonavir-boosted protease inhibitor, 39 (6%) had triple-class virological failure (defined as virological failure of two NRTIs, one NNRTI, and ritonavir-boosted protease inhibitor). The trend in the incidence of triple-class virological failure during each year after the start of ART was similar to the main analysis (data not shown), and by 5 years after ART initiation the estimated cumulative proportion of children who had such failure according to this narrow definition was 8·2% (95% CI 5·1–11·2). This proportion was significantly higher than that of adults heterosexually infected with HIV in the PLATO II project—4·2% (3·8–4·6), HR 2·2 (1·6–3·0; p<0·0001); adjusted for AIDS diagnosis before ART initiation and year of ART initiation. To explore the extent to which this raised risk of triple-class virological failure in children compared with adults was driven by treatment failure in older children and adolescents aged 10–15 years at ART initiation, we repeated this analysis, adjusted for AIDS diagnosis before ART initiation and year of ART initiation, for children aged 9 years or younger compared with adults; the HR was 1·7 (1·1–2·5; p=0·011).

Mutation data from resistance tests were available for 40 (77%) children who had triple-class virological failure (81 tests). These tests detected mutations against NRTI in 26 of 36 children, mutations against NNRTIs in 28 of 29, and mutations against protease inhibitors in 0 of 15, while they took ART drugs of the corresponding class. M184V (17%), M41L (6%), T215Y (6%), and D67N (5%) were the most common NRTI mutations; nine children had at least one thymidine-associated mutation. K103N (14%), Y181C (13), and Y188L (6) were the most common NNRTI mutations.

Discussion

12% of children had triple-class virological failure after 5 years on ART, and about a fifth had such treatment failure by 8 years on ART. Although these risks are low, and highlight the great success of antiretroviral treatment in children, they raise concerns about the proportion of children starting ART who are likely to maintain viral suppression for life, despite the potential availability of newer drugs from other classes (panel). The rate of triple-class virological failure was highest in the first 2 years after ART initiation, rising gradually thereafter. After 6 years on ART, the number of children with failure was small and the confidence intervals were wide, but the apparent decline in the rate of triple-class virological failure could also mirror early dropout of children who adhered most poorly to drug regimens, which is in line with our finding that about a quarter of children with triple-class virological failure had never achieved virological suppression. This small group started ART at an earlier age and became triple-class exposed more quickly, raising the issue of adequate dosing in young children and presumably relating to very poor adherence by their caregivers.

Restriction of the analysis to children starting ART with two NRTIs and either a NNRTI or a boosted protease inhibitor and use of the same definition of treatment failure showed that the overall rate of failure in children was over two-times higher than in adults with heterosexually transmitted HIV in COHERE. This increased rate of triple-class virological failure could be explained by lower virological suppression rates in children than in adults, absence of alternative regimens, adherence issues related to taste and formulation, a tendency for switches in regimen for children being delayed because of the need to manage adherence problems before switching treatment and hence switching regimen at high viral loads, and social factors, especially those that affect adolescents.

As in the adult analysis, adolescents and young adults had the highest risk of triple-class virological failure. Drug adherence is a challenge for children and young people with any chronic disease. For those with HIV infection, there are additional factors, including coming to terms with disclosure of their HIV status, secrecy and guilt among adult family members, and dealing with HIV alongside their own sexual development. Fear of stigma increases their isolation and tendency towards denial, all of which might adversely affect drug adherence. Although there are several studies of adherence to ART in children and young adults with HIV, most are small, heterogeneous—including both caregiver and child adherence measurements—and nearly all have been cross-sectional. Murphy and colleagues reported a decline in adherence by teenagers over time associated with duration of ART, and Kekitiinna and colleagues described poor responses to ART in newly diagnosed young people aged 10 years or older in the UK and Ireland.

However, when adolescents aged 10–15 years were excluded from the comparison with adults, the HR declined from 2·2 to 1·7, but remained significant. This finding suggests that factors relating to young children, whose drugs are mainly given by caregivers, are also important and need to be considered. Few adherence
studies have differentiated child and caregiver adherence or studied changes in adherence longitudinally with age. In addition to adherence, other complexities in treatment of children with HIV include dosing and pharmacokinetics of drugs, and palatability and tolerability of formulations, all of which vary substantially with age. All these factors might adversely affect virological response and contribute to poorer responses in children than in adults.

We recorded no association between drug classes used in the initial regimen and triple-class virological failure, supporting the use of regimens containing either NNRTIs or protease inhibitors as first-line ART in accordance with the current guidelines. Although our study was a non-randomised comparison, the PENPACT-I trial of ART initiation with protease inhibitor versus NNRTI combination ART regimens, in children with HIV-1 infection who have not received ART, showed no significant virological, immunological, or clinical differences between these two randomised arms at 4 years after ART initiation; of note, PENPACT-I included both boosted and unboosted protease inhibitors, as in the present study. Our results also suggest that the rate of triple-class virological failure was higher in children who had been diagnosed with AIDS before starting ART, compared with those without an AIDS diagnosis at ART initiation, supporting the current recommendations of ART initiation at higher CD4 counts than previously used, and before clinical disease progression. However, virological failure was not associated with CD4 percentage or viral load at ART initiation, unlike in adults with HIV and could be due in part to low power, but also to the longer time taken to attain a virological set-point after primary infection in children (about 5 years) than in adults. No association between virological failure and use of ART for prevention of mother-to-child transmission was seen, although numbers were small.

Virological failure to a drug class does not necessarily mean that no drugs from that class can successfully suppress the virus in the future, especially if no resistance mutations occurred. For children with resistance test data, we found that most children on a NRTI or a NNRTI had resistance mutations, implying loss of future drug options from these classes. By contrast, protease-inhibitor resistance was not seen, which is consistent with results from the linked adult study and a randomised trial in children. Because boosted protease inhibitors are now the standard of care for children with HIV, resistance to them is probably unusual in children failing a boosted protease-inhibitor regimen.

Our results have implications for developing and developed countries. Increasingly, drugs from the original three classes are widely available, and our results provide encouragement that viral load suppression can be achieved and maintained for many years with use of these drugs. In adults, adherence to ART has proved to be at least as high in sub-Saharan Africa as it is in North America, supporting the fact that outcomes reported in developed countries are relevant for developing countries. However, regardless of setting, there will be a need for availability of newer drugs if children with HIV infection are to maintain viral suppression for sufficient durations to enable them to live as long as their uninfected peers. If options do become exhausted then it becomes important to understand the immunological and clinical consequences of long-term virological failure on ART in children, as has been studied in adults. Such analyses are beyond the scope of this study and, in fact, there have not been large enough numbers of children in the PLATO II project in this situation to allow reliable conclusions to be drawn.

The rate of virological failure of the three original drug classes seen in this study shows the challenge of maintaining lifelong viral suppression in children who start ART much earlier in life than do adults. Further detailed analysis is needed to compare rates of switching to second-line ART and viral suppression on second-line ART between adults and children, and to compare the development of resistance, and to repeat this analysis once further follow-up time has accrued; there is continued need for strategies to promote optimum drug adherence in children, caregivers, and young people to minimise the likelihood of triple-class virological failure, and for development of suitable new drugs and formulations to optimise the treatment of children with treatment failure. Fixed drug combinations and simplification of strategies could be important ways to maintain treatment options while children move through adolescence and reach adulthood.
Contributors
All members of the PLATO II analysis and writing group participated in discussions about the design of the study, the choice of statistical analyses and interpretation of the findings, and were involved in the preparation and review of the final report. Additionally, Rebecca Lodwick and Andrew Phillips were responsible for undertaking all analyses; Rebecca Lodwick acted as guarantor for the analyses and had full access to the data set. Hannah Castro, Ali Judd, Diana M Gibb, Karina Butler, Rebecca Lodwick, and Andrew Phillips were responsible for the study concept and design. Ali Judd, Di Gibb, Arv van Sijghem, Jose T Ramos, Josiane Waszarski, Claire Thorne, Antoni Noguera-Julian, Niels Obel, Dominique Costagliola, Pat A Tookey, and Genevieve Chene acquired data for the study. Hannah Castro, Ali Judd, Di Gibb, Karina Butler, Rebecca Lodwick, and Andrew Phillips analysed and interpreted the data and drafted the report. All authors critically reviewed the report. Céline Colin, Jesper Kjaer, and Jesper Grarup provided administrative, technical, and material support.

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Conflicts of interest
We declare that we have no conflicts of interest.

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