Cohort profile: the Netherlands Cohort Study on Acute HIV infection (NOVA), a prospective cohort study of people with acute or early HIV infection who immediately initiate HIV treatment

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ABSTRACT

Purpose Initiation of combination antiretroviral therapy (cART) during acute or early HIV infection (AEHI) limits the size of the viral reservoir and preserves immune function. This renders individuals who started cART during AEHI promising participants in HIV-cure trials. Therefore, we established a multicentre prospective cohort study in the Netherlands that enrolls people with AEHI. In anticipation of future cure trials, we will longitudinally investigate the properties of the viral reservoir size and HIV-specific immune responses among cohort participants.

Participants Participants immediately initiate intensified cART: dolutegravir, emtricitabine/tenofovir and darunavir/ritonavir (DRV/r). After 4 weeks, once baseline resistance data are available, DRV/r is discontinued. Three study groups are assembled based on the preparedness of individuals to participate in the extensiveness of sampling. Participants accepting immediate treatment and follow-up but declining additional sampling are included in study group 1 (‘standard’) and routine diagnostic procedures are performed. Participants willing to undergo blood, leukapheresis and semen sampling are included in study group 2 (‘less invasive’). In study group 3 (‘extended’), additional tissue (gut-associated lymphoid tissue, peripheral lymph node) and cerebrospinal fluid sampling are performed.

Findings to date Between 2015 and 2020, 140 individuals with AEHI have been enrolled at nine study sites. At enrolment, median age was 36 (IQR 28–47) years, and 134 (95.7%) participants were men. Distribution of Fiebig stages was as follows: Fiebig I, 3 (2.1%); II, 20 (14.3%); III, 7 (5.0%); IV, 49 (35.0%); V, 39 (27.9%); VI, 22 (15.7%). Median plasma HIV RNA was 5.9 (IQR 4.7–6.7) log10 copies/mL and CD4 count 510 (IQR 370–700) cells/μL. Median time from cART initiation to viral suppression was 8.0 (IQR 4.0–16.0) weeks.

Future plans The Netherlands Cohort Study on Acute HIV infection remains open for participant enrolment and for additional sites to join the network. This cohort provides a unique nationwide platform for conducting future in-depth virological, immunological, host genetic and interventional studies investigating HIV-cure strategies.

INTRODUCTION

Despite the availability of combination antiretroviral therapy (cART), worldwide approximately 1.7 million people newly acquired HIV in 2019, 690 000 people died of AIDS and one-third of people living with HIV (PLWH) still did not have access to cART. Further expansion of access to cART alone will not be sufficient to address these gaps, as universal cART has only led to a modest decrease in HIV incidence in certain high incidence regions and stigma remains a major barrier to HIV care, leading to reduced access and adherence to cART. Furthermore, PLWH on cART may continue to experience a substantial physical and psychosocial burden.
illustrate the need to develop therapeutic strategies that lead to HIV cure.\(^7\) HIV cure is defined as a therapeutic strategy resulting in prolonged viral suppression in the absence of cART. There are several essential elements involved in the design of potentially favourable cure strategies. First, insight in viral and host-immune factors that are associated with viral control in the absence of cART is needed. Important in this respect is the observation that people who start cART in the early phase of infection potentially have the greatest possibility to benefit from such an intervention.\(^6\) Second, there is increasing insight, which indicates that the properties of the viral reservoir and the concomitant HIV-specific immune response are important factors determining the clinical outcome of cure interventions. The establishment of cohorts that have been well characterised with respect to viral and immunological properties is therefore pivotal to study potential cure strategies.\(^9\)

cART suppresses plasma viraemia to below detection limits in the vast majority of PLWH,\(^10\) but is not able to clear the virus completely and an intracellular latent viral reservoir persists. Within the first hours following HIV infection, a viral reservoir is established,\(^11\) which is mainly localised in peripheral CD4\(^{+}\) T cells and lymphoid tissues.\(^12\)\(^–\)\(^16\) The presence of the viral reservoir is responsible for a rapid rebound of viraemia when cART is interrupted.\(^17\)\(^–\)\(^18\) However, when cART is initiated during acute or early HIV infection (AEHI; here defined as the first 6 months after infection), a select group of individuals is able to remain aviraemic after treatment interruption. This phenomenon is often referred to as post-treatment control or HIV remission. Studies suggest that this rare event of viral control related to treatment initiation during AEHI is associated with a significantly smaller viral reservoir.\(^8\)\(^–\)\(^25\) The relatively minor proportion of individuals that achieve post-treatment control together with the results from a seminal study that showed viral rebound in individuals that initiated cART in the earliest stage of infection,\(^26\) indicate that early containment of the viral reservoir by cART is only partially related to viral control.

**Figure 1** Overview of NOVA cohort study procedures for study groups 2 and 3. cART, combination antiretroviral therapy; GALT, gut-associated lymphoid tissue; PBMC, peripheral blood mononuclear cells.

**Figure 2** Disposition of The Netherlands Cohort Study on Acute HIV infection cohort study participants, August 2015 to July 2020. ‘Standard’ group 1 includes immediate combination antiretroviral therapy initiation and routine monitoring; ‘Less invasive’ group 2 includes routines monitoring, blood sampling for peripheral blood mononuclear cells and virological analyses, leukapheresis and semen sampling. ‘Extended’ group 3 includes group 2 sampling procedures and gut-associated lymphoid tissue, peripheral lymph node and cerebrospinal fluid sampling.
The HIV-specific T-cell response importantly contributes to the control of viral replication.\(^8\)\(^{26-28}\) The effect of early start of cART on the immune response can be discerned from the few studies in which participants were treated during AEHI and subsequently interrupted treatment. In the Viro-Immunological Sustained CONtrol after Treatment Interruption (VISCONTI) study, participants who were temporarily treated during early infection and then discontinued cART, post-treatment control occurred in 14 participants and was associated with a lower level of T-cell activation after discontinuation of cART.\(^8\) A recent study in which participants initiated cART during even earlier stages of infection (Fiebig stages I, II and III) showed that, before the start of cART, very early CD8\(^{\text{pos}}\) T cells (in Fiebig stages I and II) had a memory signature, whereas later CD8\(^{\text{pos}}\) T cells (in Fiebig stage III) were equipped with effector function.\(^27\) Furthermore, in participants that started cART very early (Fiebig I), CD8\(^{\text{pos}}\) T cells before the start of cART had less breadth and a low activation state,\(^26\) and treatment resulted in enhanced effector function and less exhaustion.\(^28\)

Taken together, these studies show that early cART reduces inflammation and improves T-cell function, but at least some viral exposure may be needed to drive the development of a potent CD8\(^{\text{pos}}\) T-cell response.

As mentioned, individuals who start cART early in the course of infection have a potentially good starting point to benefit from cure interventions given their relatively small reservoir size and potentially a potent HIV-specific immune response, both of which depend on the Fiebig stage during which they initiated cART. The Netherlands Cohort Study on Acute HIV infection (NOVA) was initiated in 2015. This is a national prospective cohort study among participants with AEHI who immediately start cART. NOVA provides a platform for gaining detailed insight into participants’ viral reservoir size and composition and their HIV-specific immune responses before and during treatment, in anticipation of future clinical trials aimed at HIV cure.

### TABLE 1 Baseline characteristics of first 140 The Netherlands Cohort Study on Acute HIV infection cohort study participants, August 2015 to July 2020

| Study setting | NOVA was initiated in August 2015 as an ongoing multi-centre prospective cohort study enrolling participants diagnosed with AEHI who immediately initiate cART on diagnosis. Participants currently are enrolled in nine HIV treatment centres across the Netherlands (Amsterdam University Medical Centers (Academic Medical Center site), Amsterdam; Erasmus University Medical Center, Rotterdam; Maasstad Hospital, Rotterdam; Onze Lieve Vrouwe Gasthuis, Amsterdam; DC Klinieken, Amsterdam; University Medical Center, Utrecht; Radboud University Medical Center, Nijmegen; Leiden University Medical Center, Leiden and Rijnstate Hospital, Arnhem).

### Study population

The NOVA cohort study enrolls individuals who are (1) 18 years or older; (2) diagnosed with AEHI (defined according to Fiebig staging at the time of diagnosis);\(^29\) and (3) willing to initiate cART within 24 hours of enrolment. Acute infection (Fiebig stages I–II) is defined as either plasma HIV RNA detectable by reverse transcription (RT)-PCR or HIV p24 antigen detectable by fourth generation ELISA without detectable anti-HIV antibodies.
infection (Fiebig stages III–VI) is defined as plasma HIV RNA detectable by RT-PCR and with anti-HIV antibodies detectable by fourth generation ELISA in the presence of a negative, indeterminate or positive western blot. Individuals with a positive western blot are only included if their western blot is p31 negative or if they have a documented negative HIV test, 6 months prior to their HIV diagnosis.

**Study design**

Three study groups are assembled based on the preparedness of individuals to participate in the extensiveness of sampling (figure 1). Participants that accept immediate treatment and follow-up but decline additional blood and tissue sampling are included in study group 1 (‘standard’) and only routine diagnostic and follow-up procedures are performed. Participants willing to undergo blood sampling for peripheral blood mononuclear cells (PBMC) and virologic analyses, leukapheresis and semen sampling are included in study group 2 (‘less invasive’). In study group 3 (‘extended’), additional tissue (gut-associated lymphoid tissue, peripheral lymph node) and cerebrospinal fluid sampling are performed. In participants who provide written informed consent to participate in NOVA, samples are obtained at the time of enrolment and at several subsequent time points to analyse the size and characteristics of the viral reservoir and the immune response. Study participants are followed for at least 10 years and are allowed to change between groups if they wish so at any time point.

**Participant selection and recruitment**

Individuals diagnosed with AEHI who are referred for cART initiation to one of the participating study sites are informed about the study by their treating physician. The screening takes place within 24 hours after the study team has been made aware of a potential AEHI case. AEHI diagnosis can take place at a sexually transmitted infection (STI) clinic, general practice or hospital. STI clinics in the Netherlands routinely offer fourth generation ELISA antigen/antibody testing free of charge to men who have sex with men (MSM) and other high-incidence populations. These groups are advised to test for HIV every 3 months, in case of AEHI/STI symptoms or after they have been notified for HIV by a sexual partner. HIV-RNA testing by RT-PCR is only conducted if AEHI is highly suspected. For example, at the Amsterdam STI clinic this is offered to MSM with a high score on the ‘Amsterdam score’, an AEHI screening score, which includes AEHI symptoms and behavioural factors. Among 106 participants with a CD4/CD8 T-cell ratio of <1 at enrolment (10 participants with FI–II, 41 participants with FIII–IV, 55 participants with FV–VI), **after 3 years of follow-up, less than 75% of participants achieved a CD4/CD8 T-cell ratio ≥1, therefore it is not possible to estimate the 75th percentile.**

cART, combination antiretroviral therapy; F, Fiebig stage.

### Table 2  Clinical and laboratory parameters of The Netherlands Cohort Study on Acute HIV infection cohort study participants stratified by Fiebig stage, August 2015 to July 2020

<table>
<thead>
<tr>
<th></th>
<th>All participants (n=140)</th>
<th>FI–II (n=23)</th>
<th>FIII–IV (n=56)</th>
<th>FV–VI (n=61)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Median</td>
<td>IQR</td>
<td>Median</td>
<td>IQR</td>
</tr>
<tr>
<td>Baseline plasma HIV RNA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(log10 copies/mL)</td>
<td>5.9*</td>
<td>4.7–6.7</td>
<td>6.4†</td>
<td>5.0–7.0</td>
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<tr>
<td>Baseline CD4&lt;sup&gt;pos&lt;/sup&gt; T-cell count (cells/mm&lt;sup&gt;3&lt;/sup&gt;)</td>
<td>510†</td>
<td>360–700</td>
<td>500†</td>
<td>330–580</td>
</tr>
<tr>
<td>Baseline CD8&lt;sup&gt;pos&lt;/sup&gt; T-cell count (cells/mm&lt;sup&gt;3&lt;/sup&gt;)</td>
<td>940†</td>
<td>540–1430</td>
<td>540†</td>
<td>300–790</td>
</tr>
<tr>
<td>Baseline CD4/CD8 T-cell ratio</td>
<td>0.55†</td>
<td>0.37–0.96</td>
<td>1.02†</td>
<td>0.53–1.76</td>
</tr>
<tr>
<td>Days from HIV diagnosis to cART initiation</td>
<td>1</td>
<td>0–7</td>
<td>1</td>
<td>0–1</td>
</tr>
<tr>
<td>Weeks from cART initiation to viral suppression‡§</td>
<td>8.0*</td>
<td>4.0–16.0</td>
<td>10.0†</td>
<td>5.3–20.0</td>
</tr>
<tr>
<td>Weeks from cART initiation to CD4/CD8 T-cell ratio ≥1‡¶</td>
<td>52.1</td>
<td>11.6–∞**</td>
<td>10.0</td>
<td>4.1–∞**</td>
</tr>
</tbody>
</table>

*2 missing values.
†Missing value.
‡Assessed with Kaplan-Meier estimates.
§Defined as the first documented plasma HIV RNA <40 copies/mL.
¶Among 106 participants with a CD4/CD8 T-cell ratio of <1 at enrolment (10 participants with FI–II, 41 participants with FIII–IV, 55 participants with FV–VI).
**After 3 years of follow-up, less than 75% of participants achieved a CD4/CD8 T-cell ratio ≥1, therefore it is not possible to estimate the 75th percentile.

cART, combination antiretroviral therapy; F, Fiebig stage.
testing if a person presents with exposure to HIV in the previous 3 months combined with AEHI symptoms. For these reasons, it may be more likely that AEHI is identified in a person with symptoms than in a person without symptoms. Furthermore, due to the high HIV incidence in MSM and routine screening offered to MSM and other high-incidence populations, it is more likely to identify AEHI in these groups than in low-incidence populations, such as heterosexual individuals.

**Fiebig staging**

According to the study protocol, participants are enrolled within 24 hours of diagnosis and samples are obtained on the day of enrolment. Subsequently, ART should be initiated within 24 hours of enrolment. However, in some cases, there may be a delay between diagnosis and enrolment or between enrolment and ART initiation due to logistical factors (eg, study staff not being available for enrolment during weekends) or participant-related factors (eg, a participant needing time to decide whether they wish to initiate ART immediately). Fiebig stage at ART initiation (rather than at diagnosis) is correlated with the size of the HIV reservoir and the Fiebig stage may evolve rapidly between diagnosis and start of ART treatment. Therefore, in these cases, we determine Fiebig stages on samples obtained on the day of ART initiation or, if such a sample is not available, within 3 days before or after ART initiation. For participants of whom no sample is available within this time frame, the Fiebig stage is estimated by extrapolation based on the estimated duration of each stage as described by Fiebig et al (Fiebig stage I: 5 days; II: 5 days; III: 3 days; IV: 6 days; V: 70 days; VI: open ended). For example, if a sample has been obtained 6 days before ART initiation and indicated Fiebig stage IV, we classify this participant as Fiebig stage V at ART initiation.

**Treatment**

Consenting participants start ART within 24 hours of enrolment with a regimen including emtricitabine/tenofovir disoproxil fumarate 200/245 mg once daily, dolutegravir (DTG) 50 mg once daily and darunavir/ritonavir (DRV/r) 800/100 mg once daily. DRV/r was added to standard triple-drug treatment to account for potential transmitted HIV drug resistance, while awaiting the results of genotypic resistance testing. DRV/r was selected because of the virtual lack of transmitted resistance to HIV protease inhibitors in the Netherlands, its high genetic barrier and good tolerability. In all participants, DRV/r is discontinued at week 4 depending on the results of baseline resistance testing. From 2015 through 2019, participants used DTG two times per day during these first 4 weeks of treatment, and plasma samples to evaluate DTG and DRV/r pharmacokinetics were collected, as DRV/r could potentially decrease plasma levels of DTG. Since 2020, the protocol was adjusted to DTG 50 mg once daily, as a dose adjustment for DTG is no longer recommended when used with DRV/r. We have ensured continued pharmacokinetic analysis of DTG before and after DTG dose adjustment. The HIV treating physicians and a specialised HIV nurse perform HIV counselling with respect to readiness to initiate ART, ART adherence, quality of life and sexual behaviour at enrolment and throughout the study period. Data are collected in collaboration with AIDS Therapy Evaluation in the Netherlands (ATHENA) National HIV cohort, which encompass data of 98% of all PLWH in care in the Netherlands. Clinical data in the ATHENA cohort are collected prospectively by trained data monitors using standardised case record forms.

**Patient and public involvement**

The NOVA cohort study has a strong engagement with PLWH as well as individuals behaviourally vulnerable for acquiring HIV. First, an essential element in the design of the NOVA cohort study has been to create awareness for AEHI among MSM, by launching two communication campaigns focusing on AEHI, its symptoms and the benefits of immediate treatment (www.hebikhiv.nl/en). These campaigns have been developed and delivered through cocreation with MSM living with and without HIV, through MSM focus groups, communication experts from the MSM community and the Dutch HIV Association. Second, the Dutch HIV Association of PLWH (Hiv Vereniging) is represented among the NOVA collaborators and provided with regular updates on progress of the study as well as scientific development on HIV cure. Finally, future clinical trials of novel interventions aiming at achieving post-treatment control will likely include analytic treatment interruption (ATI) and thereby can be expected to have emotional and physical
Impact on participants. Assessing the views of potential participants concerning ATI during cure trials is therefore essential and will provide indispensable information for trial design. We have recently explored knowledge and perception of HIV cure and willingness to participate in cure trials among NOVA participants by conducting 20 in-depth interviews. Furthermore, as part of this substudy, we explored barriers to undergo additional sampling in NOVA (such as in study groups 2 and 3).

**Time from diagnosis to viral suppression**

Starting cART during AEHI improves clinical outcomes. Furthermore, initiation of cART during AEHI results in rapid plasma viral suppression and limits the size of the viral reservoir measured as HIV-DNA in PBMC. However, diagnosis of AEHI can be challenging and referral to an HIV treatment centre may cause delay in cART initiation. To overcome these barriers, an AEHI search and treat-to-suppression strategy was implemented at the STI clinic in Amsterdam in August 2015. In this strategy, mobilisation for AEHI testing through an AEHI communication campaign was combined with point-of-care HIV-RNA testing by RT-PCR, same-day delivery of HIV diagnosis and immediate referral to an HIV treatment centre for immediate cART initiation. This strategy proved to be a feasible and effective approach in diagnosing AEHI and significantly reducing the time from diagnosis to viral suppression. The HIV treatment centres in Rotterdam work together to liaise with STI clinics and general practitioners and have direct access to point-of-care HIV-RNA testing by RT-PCR to facilitate referring people diagnosed with AEHI and start cART within 24 hours. To facilitate early diagnosis, we implemented education on AEHI and NOVA in regional meetings with general practitioners, STI clinics and first-line laboratories and made point-of-care HIV-RNA testing available for them. Recently, an online platform was launched for infectious diseases specialists, general practitioners and physicians working at STI clinics to enable direct communication and potentially facilitating faster referral of people diagnosed with AEHI in the future.

**Findings to date**

**Baseline characteristics**

The NOVA cohort study was initiated in August 2015 and since then (data update July 2020) 140 participants have been enrolled. Of these, 131 (93.6%) remain in active follow-up. Overall, 76 (54.3%) participants have been included in group 1; 55 (39.3%) in group 2; and 9 (6.4%) in group 3. Barriers for the decision to participate in study group 2 or 3 explored in 20 in-depth interviews included fear of possible risks related to sampling procedures (such as lumbar puncture or intestinal biopsy), needle phobia, being overwhelmed by the HIV diagnosis and practical concerns, such as the difficulty to combine study visits with work. The majority of participants are men and reported to be MSM (124, 92.5%). Further baseline characteristics are provided in table 1. On average, approximately 30 participants have been enrolled each year, with the exception of 2020, during which HIV testing, diagnosis and NOVA enrolment was strongly reduced due to the COVID-19 pandemic. Median duration of follow-up is 2.4 (IQR 1.4–3.1) years.

![Figure 4](https://example.com/figure4.png)

**Figure 4** Median CD4/CD8 T-cell ratio and IQR among The Netherlands Cohort Study on Acute HIV infection cohort study participants, August 2015 to July 2020. (A) Displays CD4/CD8 T-cell ratio median and IQR among participants who initiated cART during Fiebig stages I–II; (B) Fiebig stages III–IV; and (C) Fiebig stages V–VI. D, day; M, month; Y, year.
estimated by extrapolation based on the estimated duration of each stage. Participants in the NOVA cohort study initiated cART a median of 1 (IQR 0–7) day after HIV diagnosis. No resistance-associated mutations relevant to the cART regimen provided in NOVA were detected at baseline. Median plasma HIV RNA at baseline was 6.4 (IQR 5.0–7.0) log10 copies/mL among participants with Fiebig stages I–II, 6.4 (IQR 5.5–7.0) among those with stages III–IV and 5.0 (IQR 4.5–5.9) among those with stages V–VI (table 2). Median time from cART initiation to viral suppression was slightly longer among participants with Fiebig stages I–II (10.0 (IQR 5.3–20.0) weeks) than among participants treated during stages V–VI (7.7 (IQR 4.0–12.0) weeks) (figure 3).

Time to CD4/CD8 T-cell ratio of at least 1
Median CD4<sup>pos</sup> T-cell count at baseline was 500 (IQR 330–580) cells/mm<sup>3</sup> among Fiebig stages I–II participants, 480 (IQR 325–620) among stages III–IV participants and 560 (IQR 422–730) among stages V–VI participants. Pre-cART CD4/CD8 T-cell ratio <1 is associated with chronic immune activation among PLWH who initiated treatment during chronic stages of infection<sup>30, 31</sup> and is inversely correlated with the size of the viral reservoir among treated people with AEHI<sup>22</sup>. Therefore, we assessed the time to the first CD4/CD8 T-cell ratio of ≥1 in our cohort stratified by Fiebig stage, using Kaplan-Meier estimates.

Longitudinal CD4<sup>pos</sup> T-cell and CD8<sup>pos</sup> T-cell results were available from 139 (enrolment), 70 (month 1,±15 days), 55 (month 2,±15 days), 61 (month 3,±15 days), 88 (month 6,±30 days), 90 (year 1,±60 days), 51 (year 2,±60 days) and 39 (year 3,±60 days) participants. Median CD4/CD8 T-cell ratio at enrolment was 0.55 (IQR 0.37–0.96), with 52 (23.0%) participants having a ratio ≥1, including all 3 (9.4%) who had initiated cART during Fiebig stage I. Figure 4 displays median CD4/CD8 T-cell ratios during the first 3 years of follow-up. Among participants with a CD4/CD8 T-cell ratio <1 at enrolment, median time to the first CD4/CD8 T-cell ratio of ≥1 was 52.1 (IQR 11.6–∞) weeks. This was shorter among participants treated during Fiebig stages I–II (n=10; 10.0 weeks) than among those treated during stages III–IV (n=41; 25.9 weeks) or during stages V–VI (n=55; 54.0 weeks; figure 4). However, only 10 participants with Fiebig stages I–II had a CD4/CD8 T-cell ratio <1 at cART initiation, these findings should be interpreted with caution.

In 2015, we established the NOVA as an ongoing multicentre prospective cohort study of people with AEHI in the Netherlands. Since then, we have been enrolling an average of approximately 30 participants per year, with the exception of 2020 during which enrolment was strongly reduced due to the COVID-19 pandemic. The high retention of 94% and a median of 8 weeks to viral suppression suggest that starting cART during AEHI is acceptable and adherence to cART is high. The considerable amount of time spent on counselling by a specialised HIV nurse and treating physicians within the context of the NOVA could be a factor influencing these findings. Alternatively, PLWH willing to participate in the study may be more motivated compared with those who refused to take part. Unfortunately, we were not able to collect data on PLWH not willing to participate.

In the years since start, we have been able to enrol participants in the early Fiebig stages I and II. This is particularly challenging due to the short duration of these stages. The challenge to rapidly enrol participants diagnosed with Fiebig stages I and II has been described by others, and a delay between diagnosis and enrolment often results in participants no longer being in this phase of infection at enrolment. We will conduct sensitivity analyses excluding participants with an estimated Fiebig stage rather than a measured Fiebig stage for these participants. This strategy may have the potential to introduce error, as the duration of each stage can vary from person to person. In future studies within this cohort, we will conduct sensitivity analyses excluding participants with an estimated Fiebig stage. Furthermore, we once more emphasised to the study sites that future participants should be enrolled within 24 hours of diagnosis and samples are obtained on the day of enrolment, and we have planned regular investigator meetings to support this practice.


Figure 5 Kaplan-Meier survival curve of months from cART initiation to reaching a CD4/CD8 T-cell ratio ≥1 among The Netherlands Cohort Study on Acute HIV infection cohort study participants with a baseline CD4/CD8 T-cell ratio <1, stratified by Fiebig stage, August 2015 to July 2020. The black hashes represent censored participants. The following participants were excluded from this analysis: one participant with missing CD4/CD8 T-cell ratio values, one participant with only one value available, which was obtained before cART initiation and 32 participants with a CD4/CD8 T-cell ratio ≥1 at baseline. cART, combination antiretroviral therapy; F, Fiebig stage.
Together now almost half of the total cohort participates in the ‘extended’ groups 2 and 3. The number of participants enrolled in group 3 is limited, and we may not have sufficient power to perform analyses within this group stratified by Fiebig stage. However, the number and material will be sufficient and valuable for longitudinal studies within the same participant, for example, to study viral reservoir dynamics, including within host viral evolution. In addition, we will continue our efforts to enrol participants in group 3 in the coming years.

The blood and tissue samples allow us to longitudinally characterise the viral reservoir and the properties of the HIV-specific immune response in great detail. We plan to comprehensively characterise the viral reservoir with respect to proviral latency and residual replication competence; and characterisation of the cellular localisation of the viral reservoir with respect to type of cell and cellular phenotype (including activation status). The HIV-specific immune response will be characterised by assessing the frequency and breadth of HIV-specific CD8\(^+\) T cells, the phenotype and functional properties of the HIV-specific CD8\(^+\) T cells and the distribution of CD8\(^{pos}\) T cells in lymph nodes versus peripheral blood. Furthermore, in the coming years, we expect novel approaches to HIV cure to be developed, for example, the in vivo efficacy of various approaches to purge the latent viral reservoir and new ex vivo studies investigating the response of latency reversing agents and immune-based therapies in various cell types from PLWH who are on cART.\(^{53}\) NOVA will provide a valuable platform to conduct clinical trials evaluating such newly developed approaches, also in light of existing and future international collaborations.\(^{34}\)

The NOVA cohort study complements several other prospective cohorts on AEHI worldwide (among others those from US, African, South American and Thai sites NCT00296660 and NCT02855558; San Francisco NCT02656511; Gent NCT03449706; Zurich NCT00537966; Bangkok RV254/SEARCH010 studies\(^{21}\); East Africa and Thailand RV217\(^{35}\); and the South African FRESH cohort).\(^{26}\) The course of HIV infection (including viral load at AEHI, viral setpoint and viral reservoir) is highly variable between cohorts.\(^{56}-^{58}\) This variation is in part determined by age, sex, mode of HIV transmission and, importantly, geographical location and HIV subtype (B vs non-B).\(^{37}\) Differences in level of viremia during AEHI and viral setpoint have been shown to be influenced by host genetic and viral factors.\(^{39},^{59}\) An important host genetic factor is HLA type, which is differentially distributed across geographic (ethnic) regions.\(^{37}\) A viral factor that may determine the course of infection includes clade. A post-hoc analysis by Omondi et al indeed showed a difference in viral reservoir size between a Ugandan (black) and US (white) cohort infected with different viral clades.\(^{38}\) Because these determinants of HIV infection may impact formation of reservoir and immune response, we need multiple cohorts from around the world and different geographic locations to study this.

The nine Dutch HIV treatment centres that participate in NOVA cover the areas of the Netherlands known to have the highest HIV prevalence and serving 65% of all PLWH in care. The vast majority of new HIV diagnoses in the Netherlands occur among MSM.\(^{32},^{60}\) This may explain the over-representation of MSM (93%) in NOVA. Thus far, approximately 35% of MSM diagnosed with AEHI in the Netherlands have been enrolled in NOVA.\(^{32}\) We expect to increase this proportion in the coming years as participating study sites were added over a period of years and some of these sites have started to enrol participants only recently.

In conclusion, the NOVA cohort study is a well-characterised nationwide cohort of people who initiated cART during AEHI and provides a unique platform to conduct detailed analyses of the HIV reservoir and the host immune response and future interventional studies aimed at achieving durable HIV control in the absence of cART and cure.

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Collaborators The Netherlands Cohort Study on Acute HIV infection (NOVA) cohort study currently collaborates with several research groups in the Netherlands and enrols participants from nine clinical HIV treatment centres, and aims to actively engage additional Dutch study sites. The NOVA cohort study is overseen by a governing board composed of representatives of the two coordinating sites, Amsterdam UMC, Academic Medical Centre Centre site and Erasmus University Medical Centre. For further information visit the H-TEAM web page (https://h-team.nl/?lang=en).

Contributors GdB, CR, JMP, PR, CB and AV designed the study. MD and HP conducted the statistical analysis. MD and HP drafted the manuscript which was revised by JMP, PR, CB, AV, CR and GdB. GdB and CR supervised the project and are also responsible for its continued management. All authors reviewed, critically revised and approved the manuscript. CR contributed equally as last authors. GJB acts as the guarantor.

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The nine Dutch HIV treatment centres that participate in NOVA cover the areas of the Netherlands known to have the highest HIV prevalence and serving 65% of all PLWH in care. The vast majority of new HIV diagnoses in the Netherlands occur among MSM.\(^{32},^{60}\) This may explain the over-representation of MSM (93%) in NOVA. Thus far, approximately 35% of MSM diagnosed with AEHI in the Netherlands have been enrolled in NOVA.\(^{32}\) We expect to increase this proportion in the coming years as participating study sites were added over a period of years and some of these sites have started to enrol participants only recently.

In conclusion, the NOVA cohort study is a well-characterised nationwide cohort of people who initiated cART during AEHI and provides a unique platform to conduct detailed analyses of the HIV reservoir and the host immune response and future interventional studies aimed at achieving durable HIV control in the absence of cART and cure.

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Patient and public involvement Patients and/or the public were involved in the design, or conduct, or reporting, or dissemination plans of this research. Refer to the Cohort description section for further details.

Patient consent for publication Not applicable.

Ethics approval Study participants provide written informed consent. The study has been approved by the medical ethical committee of the Amsterdam Medical Centers (Academic Medical Center site) (NL51613.018.14).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. We endeavour to make the data used in any The Netherlands Cohort Study on Acute HIV infection (NOVA) manuscript publicly available, within the limits of the ethical governance under which the data were collected. To this end, we will share data directly with interested parties for two purposes: (1) verification and replication of an already published analysis derived from NOVA, (2) novel scientific research projects using NOVA data. To facilitate this, requests for data sharing can be made on a case-by-case basis following submission of a concept sheet. Once submitted the proposed research/analysis will undergo review by the NOVA team for evaluation of the scientific value, relevance to the study, design and feasibility, statistical power and overlap with existing projects. If the proposed analysis is for verification.replication, data will then be made available. If the proposed research is for novel science, upon completion of the review, feedback will be provided to the proposer(s). In some circumstances, a revision of the concept may be requested. If the concept is approved for implementation, a writing group will be established consisting of the proposers (up to three people that were centrally involved in the development of the concept) and members of the NOVA study group (or other appointed cohort representatives). All people involved in the process of reviewing these research concepts are bound by confidentiality. For more information about the procedure, data sharing or collaboration in general, please contact dr GdB: g.j.debree@amsterdamumc.nl.

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