Early Antiretroviral Therapy is Associated with Lower HIV DNA Molecular Diversity and Lower Inflammation in Cerebrospinal Fluid but Does Not Prevent the Establishment of Compartmentalized HIV DNA Populations

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Abstract

Even when antiretroviral therapy (ART) is started early after infection, HIV DNA might persist in the central nervous system (CNS), possibly contributing to inflammation, brain damage and neurocognitive impairment. Paired blood and cerebrospinal fluid (CSF) were collected from 16 HIV-infected individuals on suppressive ART: 9 participants started ART <4 months of the estimated date of infection (EDI) (“early ART”), and 7 participants started ART >14 months after EDI (“late ART”). For each participant, neurocognitive functioning was measured by Global Deficit Score (GDS). HIV DNA levels were measured in peripheral blood mononuclear cells (PBMCs) and CSF cell pellets by droplet digital (dd)PCR. Soluble markers of inflammation (sCD163, IL-6, MCP-1, TNF-α) and neuronal damage (neurofilament light [NFL]) were measured in blood and CSF supernatant by immunoassays. HIV-1 partial C2V3 env deep sequencing data (Roche 454) were obtained for 8 paired PBMC and CSF specimens and used for phylogenetic and compartmentalization analysis. Median exposure to ART at the time of sampling was 2.6 years (IQR: 2.2–3.7) and did not differ between groups. We observed that early ART was significantly associated with lower molecular diversity of HIV DNA in CSF (p<0.05), and lower IL-6 levels in CSF (p = 0.02), but no difference for GDS, NFL, or HIV DNA detectability compared to late ART. Compartmentalization of HIV DNA populations between CSF and blood was detected in 6 out of 8 participants with available paired HIV DNA sequences (2 from early and 4 from late ART group). Phylogenetic analysis confirmed the presence of monophyletic HIV DNA populations within the CSF in 6 participants, and the same population was repeatedly sampled over a 5 months period in one participant with longitudinal sampling. Such compartmentalized provirus in the CNS needs to be considered for the design of future eradication strategies and might contribute to the neuropathogenesis of HIV.
Introduction

Human Immunodeficiency Virus (HIV) invades the central nervous system (CNS) early during the course of infection [1,2] providing the foundations for neurocognitive impairment (NCI) and potentially establishing a latent reservoir [3,4]. Newly infected individuals typically have homogeneous HIV populations in blood [5,6] that evolve during untreated infection to generate diverse viral variants [2,7,8]. Compartment-specific selective pressures can subsequently lead to the emergence of unique HIV populations in different anatomical sites during the course of infection, including the CNS [2,7,9–11], the genital tract [12], and other tissues [13,14]. HIV RNA variants can be sequestered from blood into the CNS early after infection (within 2–6 months) and give rise to a separate HIV RNA population in the cerebrospinal fluid (CSF) [2,8], which remains genetically distinct from blood throughout the course of infection. Overall, these observations suggest that the CNS can be permissive for HIV replication from a very early period after HIV infection. The presence of compartmentalized HIV variants within the CNS has important implications: (1) compartmentalization of HIV RNA in CNS has been associated with greater inflammation and worse neurocognitive outcomes [15–17] and, (2) independent replication of HIV within the CNS might hinder HIV eradication efforts by providing a distinct reservoir of HIV persistence different from that found in peripheral CD4+ T cells. This has been suggested by previous observations reporting differential emergence of drug resistance mutations between CSF and blood during antiretroviral therapy (ART) failure [18–20].

Combination ART has markedly reduced the incidence of HIV-associated dementia [21,22]. However, the true impact of early ART initiation on HIV-associated neurocognitive impairment is still under investigation [23]. While the viral replication and evolution of HIV RNA in the CNS has been extensively studied even during early HIV infection [2,8,24,25], little is known about the HIV DNA populations persisting in this anatomic compartment during the earliest phase of HIV infection, and especially during suppressive ART. Similar to blood [26,27], initiation of ART during early HIV infection might limit the diversification of HIV DNA within the CNS, affecting the size and molecular diversity of the HIV reservoir, preventing inflammation, and limiting brain damage. But these features have not been evaluated yet for the CNS. Our study used a unique set of samples from a well-characterized cohort of HIV-
infected individuals followed longitudinally from early HIV infection to investigate the effects of early ART initiation on the size and molecular and phylogenetic characteristics of the HIV DNA populations while on long-term suppressive ART. Additionally, since chronic inflammation has been associated with HIV persistence [28], we evaluated the effects of early ART on selected inflammatory markers in blood and CSF supernatant.

Results

Study population and Samples

Study participants (n = 16) were all HIV-infected males with a median age of 41 years (Inter Quartile Range [IQR]: 32.5–52.5) selected among participants of the San Diego Primary Infection Resource Consortium (SD PIRC). At baseline (pre ART), the median plasma viral load was 176,000 HIV RNA copies/μl (IQR: 40,287–515,900). Participants achieved viral suppression after a median of 76 days (IQR: 47–256) from the estimated date of infection [EDI] and remained undetectable during the entire follow-up (median of 3.5 viral load measurements per participant, median of 168 days between visits, median % of time-points with suppressed HIV RNA during follow-up 100%). Participants received ART for a median duration of 2.6 years (IQR: 2.2–3.7) and had suppressed levels of HIV RNA in blood plasma (<50 copies/ml) and in the CSF supernatant (at single copy level) at the time of sample collection. Six out of sixteen participants were on a protease inhibitor (PI)-based ART regimen, 6/16 were on a non-nucleoside reverse transcriptase inhibitor (NNRTI)-based regimen and 4/16 on an Integrase Strand Transfer Inhibitor (INSTI)-based regimen, all in combination with two nucleoside reverse transcriptase inhibitors (NRTI).

While we recruited participants with early and late ART initiation according to study design, the exact categorization (<4 months or >14 months) was performed retrospectively to participant enrollment, but a priori to any molecular data generation or interpretation. The “early ART group” (n = 9) started ART within a median of 1.8 months from EDI (range: 1.5–3) while the “late ART group” (n = 7) started ART within a median of 17.2 months from EDI (range: 14.8–30.9). Detailed demographic and clinical characteristics of the study population are summarized in Table 1. No significant differences between the early and late ART groups were observed for any recorded demographic or clinical characteristics (p > 0.2). Paired CSF and blood samples were obtained at baseline from all 16 participants. Two participants (both belonging to the late ART group) agreed to donate CSF and blood at a second (T0338 and T0366) and a third (T0366) longitudinal visit. These additional samples were obtained 5 and 3 months from the first evaluation and 2 months from the second evaluation, respectively.

HIV DNA levels and detectability in CSF cells and PBMC

Overall, this study comprised 16 participants with baseline samples (9 early ART and 7 late ART) and 3 extra time points from 2 participants (both belonging to the late ART group). Among the 16 baseline samples, we detected HIV DNA from 6 CSF cell pellet samples (31%) by ddPCR and amplified the HIV partial env gene (C2V3, HXB2 coordinates 6,928–7,344) in 8 CSF cellular samples (50%) by nested PCR (Summarized in Supplementary S1 Table). For the purpose of our study, we considered as “positive” any CSF sample with detectable HIV DNA by either ddPCR or nested PCR (or both). This resulted in 10 HIV DNA positive CSF samples at baseline (62.5%, 5 in the early ART and 5 in the late ART group) and 6 undetectable (negative for both ddPCR and nested PCR). Of the 3 extra time point samples (longitudinal), we detected HIV DNA from one CSF cellular sample by ddPCR (T0338 TP2) but we were able to amplify C2V3 env in all 3 CSF cellular samples (T0338 TP2 and T0366 TP2 and TP3).

Of note, only 5 samples (out of the 13 with detectable HIV DNA) had consistent detection of HIV DNA by ddPCR and nested PCR across both aliquots. This inconsistency across
aliquots is not surprising because of the low number of infected cells which increases the proportional impact of unequal cell numbers across the two separate aliquots during processing. Also, the dilution of lysates before the ddPCR droplet generation may have significantly reduced the sensitivity of the ddPCR assay.

When comparing the two groups, HIV DNA was detected in 5 out of 9 CSF cell pellet samples as part of the early ART group and in 5 of 7 in the late ART group, but this difference was not statistically significant (55% versus 71%, relative risk 0.78, p = 0.63); HIV DNA was detected in all but one (93.8%) of the 16 PBMC samples.

**HIV DNA molecular characteristics in blood and CSF cells**

To further characterize the HIV DNA population, we sequenced *env* from CSF cell pellets (*n* = 8) and PBMCs (*n* = 14) at baseline. For two participants, we also obtained *env* sequences

### Table 1. Baseline demographic and clinical characteristics, HIV DNA and inflammatory markers levels between different ART initiation groups.

| Parameters | All subjects (*n* = 16) | Early ART (*N* = 9) | Late ART (*N* = 7) | p-value (Mann-Whitney) |
|------------|-------------------------|---------------------|-------------------|-------------------------|
| **Characteristics** | | | | |
| Age | 41 [32.5, 55] | 44 [40, 57] | 35 [29.5, 47.5] | 0.22 |
| Education (years) | 14 [12, 16] | 13.5 [12, 16] | 15 [13.5, 16.5] | 0.60 |
| Male, % | 93.3 | 88.9 | 100 | 1.00 |
| Caucasian, % | 86.6 | 77.8 | 100 | 0.51 |
| Current CD4 | 726 [603, 1051] | 786 [583, 1292] | 726 [603, 816] | 0.59 |
| Nadir CD4 | 374 [287, 505] | 473 [300, 592] | 373 [300, 414] | 0.52 |
| Estimated Duration of infection (years) | 3.4 [2.3, 5.1] | 2.7 [2.1, 3.4] | 5.1 [4.4, 5.7] | 0.52 |
| Exposure LT ARVs (years) | 2.6 [2.2, 3.7] | 2.8 [2.1, 3.7] | 2.6 [2.4, 3.2] | 0.81 |
| CNS Penetration Effectiveness | 7 [4, 7] | 6.5 [4, 7] | 7 [4, 7] | 0.93 |
| GDS | 0.2 [0.1, 0.9] | 0.6 [0.1, 1.1] | 0.2 [0.1, 0.2] | 0.38 |
| **CSF** | | | | |
| HIV DNA (cps/million cells) | 2,701.5 [1118.9, 4526.2] | 10,559.7 (N/A) | 1,554 [578.9, 3078.7] | 0.34 |
| Diversity (%) | 1.1 | 0.9 | 2.5 | 0.11 |
| **Cytokines** | | | | |
| sCD163 (ng/mL) | 45.8 [37.5, 63] | 39.7 [35.6, 62.6] | 49.4 [40.4, 63.2] | 0.60 |
| IL-6 (pg/mL) | 1.0 [0.9, 1.2] | 0.9 [0.6, 1.1] | 1.2 [0.9, 1.4] | 0.03 |
| MCP-1 (pg/mL) | 376.9 [339, 437.5] | 414.9 [343.8, 447.3] | 344.8 [330, 377] | 0.17 |
| TNF-α (pg/mL) | 0.24 [0.1, 0.3] | 0.2 [0.02, 0.2] | 0.3 [0.3, 0.5] | 0.02 |
| NFL (ng/L) | 168.1 [109.7, 270.5] | 173.8 [148.5, 269.8] | 138.7 [109.2, 245.5] | 0.53 |
| **Blood** | | | | |
| HIV DNA (cps/million cells) | 18.6 [9, 44.2] | 14.1 [10.1, 37] | 39.1 [7.1, 62.7] | 0.62 |
| Diversity (%) | 2.5 | 2.1 | 2.5 | 0.26 |
| **Cytokines** | | | | |
| sCD163 (ng/mL) | 648.6 [479.2, 826.4] | 790.5 [560.9, 870.5] | 508.2 [478.9, 570.1] | 0.20 |
| IL-6 (pg/mL) | 0.5 [0.2, 0.7] | 0.4 [0.2, 0.6] | 0.7 [0.5, 0.7] | 0.68 |
| TNF-α (pg/mL) | 1.7 [1.3, 2.3] | 1.7 [1.3, 2.1] | 1.8 [1.4, 3.2] | 0.80 |
| MCP-1 (pg/mL) | 118.7 [103.5, 145.8] | 126.1 [106.1, 149.1] | 113.9 [95.4, 131.9] | 0.52 |

GDS = Global Deficit Score; ART = antiretroviral therapy; LT = lifetime.

aData shown as median [interquartile range].

bData shown as median [interquartile range] among HIV DNA detectable samples. Fisher test p value is shown.

^cNot computed because only two values were available for this group.

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Early ART reduces CSF inflammation and HIV DNA diversity but does not prevent compartmentalization.
from one (T0338) or two additional time-points (T0366). Detailed characteristics of the viral populations are provided in supplementary S2 Table (for PBMC) and S3 (for CSF cell pellets).

Overall, participants in the early ART group presented a lower molecular diversity of the CSF HIV DNA population, as compared to the late ART group (Fig 1; Median: 0.9% versus 2.5%, p = 0.11). In contrast, no difference in molecular diversity was observed in the PBMC HIV DNA population between the two ART groups (Fig 1, Median: 2.1% versus 2.5%, p = 0.26). The CSF/PBMC diversity ratio was 0.58% (range: 0.31–0.69) for the early ART group and 0.84 (range: 0.33–1.06) late ART group (p = 0.12).

Next, we used a mixed-effects model where baseline viral diversity was predicted by log-transformed time to ART from EDI as a continuous variable to evaluate its association with percentage of diversity (Fig 2). We observed a higher percentage of diversity among participants with the longer time to ART from EDI, collapsed across blood and CSF (b = 0.36, p = 0.04, $\eta^2_p = 0.28$). When evaluating the compartments separately, this association was significant in CSF (p = 0.05, $\eta^2_p = 0.22$), but not in blood (p = 0.08, $\eta^2_p = 0.19$). Diversity was significantly higher in PBMC than in CSF by 0.8% (p = 0.02, $\eta^2_p = 0.31$), regardless of time to ART. We also included five covariates (age, peak viral load, CD4, CD8, and CD4/CD8 ratio) separately in the model to examine their potential effects on diversity and the association between time to ART and diversity. None of the covariates was significantly associated with

![Time to Start ART: Early vs Late](image)

Fig 1. Comparison of molecular diversity for HIV DNA (env gene) in CSF cells and PBMC between early ART versus late ART groups. Mann Whitney comparison between early ART versus late ART groups for rank-transformed percentage of molecular diversity for HIV-1 env in CSF cells and PBMC. p values for both comparisons were >0.10.

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diversity (all p-values >0.1, all η^2_p <0.05) while the association between time to ART start and diversity remained consistently significant (p-values <0.05).

The average number of input HIV DNA templates from CSF cells into the first round PCR reaction was estimated using the number of HIV DNA and RPP30 copies (based on our ddPCR data). The median HIV DNA copies per million cells among HIV-positive CSF cell samples was 2,701 copies/million cells (IQR: 1,119–4,526). The median number of CSF cells for each ddPCR reaction (estimated by RPP30) was 2,340 (IQR: 1,527–3,123 cells). After adjusting for the different volumes (5 μl for ddPCR and 10 μl for nested PCR) and the dilution factor, we estimate that the average calculated HIV template input was 22 copies of HIV DNA (range: 4–64). It should be noted that these levels are likely an under-estimate, given the inherent dilution with the ddPCR methods, as described in the method section and above.

To further evaluate if the low HIV DNA input for the sequencing reaction influenced our measures of molecular diversity, we performed additional sensitivity analyses based on our

Fig 2. Regression analysis between time to ART start and HIV DNA molecular diversity in blood and CSF. Linear mixed-effects regression models revealing the relationship between molecular diversity of HIV DNA env with log-transformed time to ART start. *p = 0.05 for the correlation within CSF.

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baseline model described above. We first assessed the potential impact of HIV DNA copies on diversity measures by including log-transformed HIV DNA levels (measured in blood and CSF when available) into our model; we found no statistical evidence that the number of HIV DNA copies was associated with any bias in molecular diversity ($p = 0.21, \eta^2_p = 0.10$). Second, to take into account the lack of consistency across aliquots, we compared diversity measures between cases with consistent versus inconsistent detectability across aliquots (assuming that cases with ddPCR+/nested PCR+ will have higher HIV DNA levels compared to cases with ddPCR-/nested PCR+) and we did not find a significant difference ($p = 0.46, \eta^2_p = 0.04$). While the ability to detect a significant effect in our sensitivity analysis was surely limited by the small sample size, this analysis suggests that the effect size of our primary predictor (time from EDI to ART, $\eta^2_p = 0.28$) on molecular diversity of partial env was greater than the effect sizes of each covariate, including the number of template HIV DNA copies ($\eta^2_p = 0.10$) and the number of positive aliquots ($\eta^2_p = 0.04$). Finally, to test the consistency of the diversity measures across blood and CSF, we performed a correlation analysis, and found that molecular diversity in CSF pellets was significantly associated with molecular diversity in PBMC (Pearson $r = 0.78, p = 0.02$), strongly supporting the validity of our conclusion and measurements within the context of all the aforementioned limitations.

Compartmentalization analysis of HIV DNA between CSF cells and PBMC

Paired HIV DNA sequences (env) from CSF cell pellets and PBMCs were obtained for 8 participants, 3 from the early ART group and 5 from the late ART group. Two individuals (both from the late ART group) had additional HIV DNA sequences from a second time-point available (obtained 3 and 5 months from the first evaluation, respectively). One individual had a third time-point (2 months apart). Compartmentalization was assessed using three distinct methods: distance-based $F_{ST}$ test with and without collapsed haplotypes and tree-based Slatkin-Maddison (SM) test. Applying our conservative definition (i.e., significant compartmentalization for all three methods), we observed a significant genetic compartmentalization between the HIV DNA populations sampled from CSF cells and PBMCs in 6 of 8 participants, including 2 individuals in the early ART group (T0104 and T0430) (Table 2). Of note, the $F_{ST}$

| Subject | TP | ART Group | Location | $F_{ST}$ test* | $p$ value | Slatkin Maddison $p$ value | Compartamentalization |
|---------|----|------------|----------|--------------|-----------|--------------------------|----------------------|
| T0430   | Early | PBMC | CSF | 0.39 | <0.01 | 0.03 | Yes |
| T0073   | Early | PBMC | CSF | -0.03 | 1 | 0.79 | No |
| T0104   | Early | PBMC | CSF | 0.2 | <0.01 | <0.01 | Yes |
| T0020   | Late | PBMC | CSF | 0.08 | 0.09 | 0.04 | Yes |
| T0133   | Late | PBMC | CSF | 0.33* | <0.01 | 0.03 | Yes |
| T0156   | Late | PBMC | CSF | 0.37 | <0.01 | <0.01 | Yes |
| T0338   | TP1 | Late | PBMC | 0.33 | <0.01 | <0.01 | Yes |
| T0366   | TP2 | Late | PBMC | 0.16 | <0.01 | 0.01 | Yes |
| TP3     | Late | PBMC | CSF | 0.24 | <0.01 | <0.01 | Yes |

Compartmentalization is called conservatively when all three tests indicate compartmentalization [see text].

*Indicates tests which become significant (or not significant) if copy numbers are ignored during $F_{ST}$ calculations. [see text]. Statistical significance was derived via a 1,000-fold population-structure randomization/ permutation test. CSF: Cerebrospinal Fluid; PBMC: Peripheral Blood Mononuclear Cells.

While $F_{ST}$ could assume negative values, none of the samples with negative $F_{ST}$ could be called compartmentalized.

Table 2. Compartmentalization Analysis for partial env HIV DNA in PBMC and CSF.

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estimates were congruent between both distance-based approaches, with and without collapsed haplotypes (Kendall τ test p < 0.01).

Phylogenetic structure of the HIV DNA populations

Maximum likelihood (ML) phylogenetic trees were created to evaluate the structure of the HIV DNA populations for participants with paired env sequences from CSF cells and PBMCs (Fig 3 and Fig 4).

Tree topologies revealed the presence of monophyletic HIV DNA populations in CSF for 6 participants (Figs 3 and 4, indicated with an asterisk). Two (T0104 and T0430) of the six individuals with evidence of well-segregated viral populations in the CSF were part of the early ART group. The same monophyletic CSF virus population was sampled from longitudinal CSF pellets over a period of 5 months for the one individual with a second time-point (T0338; Fig 4, see asterisk).

Soluble markers of inflammation and neuronal damage

Next, we investigated the effect of early ART on inflammatory markers and a marker of neuronal damage. In our cross-sectional analysis (including baseline samples), participants from the early ART group had lower levels of interleukin (IL)-6 (Fig 5 and Table 1, p = 0.03) and tumor necrosis factor (TNF)-α (Fig 5 and Table 1, p = 0.02) in CSF compared to participants from the late ART group. ART groups did not differ for any of the other soluble inflammatory markers in CSF (sCD163 and MCP-1) or blood (sCD163, IL-6, TNF-α and MCP-1) or for neurofilament light (NFL) chains in CSF (p > 0.1; Table 1).

We also used the time to ART as a continuous variable to evaluate its association with the levels of the four cytokines. We observed higher IL-6 levels among participants with the longest time to ART from EDI, collapsed across blood and CSF (b = 0.19, p = 0.02, η²_p = 0.16). When evaluated separately, this association was significant in CSF (p = 0.02, η²_p = 0.16), but not in blood (p = 0.54, η²_p = 0.01). Again the five covariates were included in the model to control for their potential effects. The CD4/CD8 ratio was significantly negatively correlated with IL-6 levels (b = -0.37, p = 0.05, η²_p = 0.12), while the other four were not correlated (all p-values > 0.1, all η²_p < 0.07). Regardless of the covariate included in the model, the association between time to ART and IL-6 remained consistently significant (p-values < 0.05).

Since IL-6 levels and HIV DNA diversity showed a similar, positive association with time to ART, we performed an additional mediation analysis to test the hypothesis that time to ART might have influenced diversity through its effect on IL-6 levels (Fig 6). While the direct effect of time to ART on diversity was still significant (p = 0.02), its indirect effect through IL-6 levels was not (p = 0.52), suggesting that IL-6 is unlikely the main mechanism connecting shorter timing of ART initiation to lower HIV DNA diversity.

Discussion

To cure HIV, all forms of viral persistence should be considered, including viral reservoirs in different tissues and anatomical compartments [2,17,29–31]. Strong evidence supports that HIV can independently replicate in the CNS during untreated infection [2,11,32] and that the virus can establish a latent reservoir in this anatomic compartment [33,34], which may be distinct from the one in circulating CD4^+ T cells. The exact timing of HIV compartmentalization within the CNS is uncertain but likely occurs soon after infection in at least some individuals [2,25]. Similarly to the periphery [35–38], we hypothesized that initiation of ART during early HIV infection would reduce the size and diversity of the viral reservoir within the CNS. To test this hypothesis, we evaluated a unique cohort of 16 HIV-infected individuals with known EDI
Fig 3. Maximum likelihood phylogenetic trees of partial HIV env sequences from paired CSF cellular pellets and PBMC sampled at baseline in 8 participants. HIV DNA populations were compartmentalized in
who were sampled while receiving long-term ART and with sustained HIV RNA suppression. As previously described [39], we were able to detect HIV DNA in cells collected from the CSF, even in participants who started ART during early HIV infection (within 4 months of EDI). We did observe that early ART was associated with less molecular diversity of HIV DNA in both CSF cells and PBMC compared to late ART. Molecular diversity was not associated with age, peak viral load, CD4, CD8 and CD4/CD8 ratio.

Interestingly, although early ART initiation was associated with lower molecular diversity of provirus, most participants presented evidence of genetic compartmentalization of HIV DNA within the CSF (including 2 out of the 3 participants from the early ART group). Six participants had a clear monophyletic population of HIV DNA in the CSF. Overall, our results are consistent with previous studies reporting the presence of compartmentalized HIV RNA in CSF of HIV-infected people very early after infection [2,25]. The detection of viral compartmentalization does not necessarily imply that the populations in CSF and in blood are completely segregated, but instead, distinct subpopulations can occur in each compartment. This can occur in two different ways. First, HIV RNA populations can be sequestered from blood and populate the CNS early after infection, giving rise to a HIV RNA population within the CSF that remains genetically distinct from blood throughout the course of infection [2]. Alternatively, HIV RNA can enter the CNS early and evolve over time as a consequence of isolated replication and differential selection pressures, creating a genetically complex population within the CNS [2]. Overall, these observations suggest that the CNS compartment is permissive for HIV replication in at least a subset of persons from a very early period after infection and likely originates a distinct reservoir from that found in the blood; however, it is noted in our study that we do not know if any of these HIV DNA sequences represented replication competent proviruses.

Another open question is the cellular source of this genetically distinct HIV DNA isolated from CSF cells. In our study, we were not able to determine the exact cellular source of the HIV DNA due to technical limitations and the nature of the samples. It is possible that this genetically distinct HIV DNA population detected in CSF might be carried by macrophages or T cells into the CSF or that T-cells circulating in CSF could get infected through contact with HIV-infected macrophages residing in the brain tissue in proximity to the brain vessels [40]. Alternatively, this HIV DNA population might be originating from CD4+ T cells circulating in the CSF after crossing the blood brain barrier but this seems less likely, since HIV-infected CD4+ T cells trafficking from the periphery into the CNS should present an equilibrated viral population in comparison to blood, especially in the setting of suppressive ART. Alternatively, unrecognized isolated HIV replication within the CNS during the period before our study visit might be responsible for our observations. Unfortunately, we did not collect longitudinal CSF samples in time points previous our baseline study visits, as part of the study design.

The novelty of our study derives from the fact that we evaluated the HIV DNA populations from cells circulating in CSF and we demonstrated the presence of compartmentalized monophyletic HIV DNA populations in CSF from HIV-infected participants receiving suppressive ART, including a participant who started ART during primary infection. Both participants with longitudinal sampling showed sustained compartmentalization at all time-points, and the same monophyletic population was repeatedly sampled from CSF over a period of 5 months in one participant.
Early ART reduces CSF inflammation and HIV DNA diversity but does not prevent compartmentalization.
Despite several technical limitations (described below), our findings are important for the design of future eradication strategies and also to improve our understanding of HIV pathogenesis in the CNS. In fact, the presence of compartmentalized HIV populations has been associated with neurocognitive impairment [41]. Several studies reported associations between circulating HIV DNA levels in blood and neurocognitive impairment with and without ART [42–46]. While this observation might hold true also for HIV DNA in CNS, this has not been consistently reported especially in the setting of suppressive ART. One previous study [3], found higher levels of HIV DNA in brain tissue from people with HIV encephalitis and moderate neurocognitive impairment compared to HIV-positive controls dying without neurologic symptoms. However, this study was limited since it included autopsy material from people dying with advanced disease and variable ART exposure. Likely due to limitations in samples size and the fact that people treated early during HIV infection have overall less neurological complications, we did not find associations between HIV DNA levels and neurological impairment.
Our study also evaluated the effect of early ART initiation on selected inflammatory biomarkers in CSF and blood. Increased inflammation has been extensively reported in the CNS during HIV and was often associated with neurocognitive impairment [47–49] even during suppressive ART [49,50]. In our study, the early ART group presented significantly lower levels of IL-6 and TNF-α in CSF (but not in blood) compared to the late ART group. We also explored the possible effect of IL-6 on molecular diversity and no mediation effect was observed. These data further support the concept that early ART initiation reduces the levels of at least some inflammatory mediators in CSF.

This study has several limitations. First of all, even though we were able to collect the volumes of CSF necessary to recover sufficient cells by the lysis buffer protocol, the detection of HIV DNA from CSF has been challenging due the low number of cells typically present in CSF in the absence of neurological symptoms and when HIV is suppressed. The low number of input cells might increase the potential for error related to sampling bias, could possibly...
amplify the number of false positive events from the ddPCR assay and could affect our diversity and compartmentalization analysis. To partially evaluate its impact, we performed multiple sensitivity analysis to address a possible bias in our analysis. Although we acknowledge that the small samples size has limited our statistical evaluation, our primary predictor of interest (i.e. the time to ART initiation) appears to have a greater effect on molecular diversity than the assay-related covariates. Further, we significantly elevated the threshold of compartmentalization detection and specifically included computational tests to increase robustness against significant errors in frequency estimation. Template input was particularly low in some (but not all) CSF samples, which could negatively impact our capacity to find unique clades within the CSF: assuming we are simply resampling the most common variants, we are more likely to find that CSF sequences fall within better sampled blood clades. In contrast, despite the possible sampling bias in CSF, we were still able to observe monophyletic CSF clades at baseline in several participants. Also, the reproducibility of the phylogenetic trees with similar clades sampled across longitudinal CSF samples for one participant, suggests that our sequences are likely informative and not substantially affected by random error or sequencing bias. Despite this, and the fact that we are analyzing only a partial region of \textit{env} gene (~400 bp), we found differences in molecular diversity and in the population structure of the HIV DNA populations in CSF between the early and late ART groups.

Another limitation of the analysis is the lack of randomization for the timing of ART initiation, which might introduce some unrecognized biases in our study design. For example, people with more symptomatic infection (including the presence of neurologic symptoms, which were not tested as part of our study) will start ART earlier and might also be more likely to present compartmentalized HIV populations. The small sample size also limited our statistical power. Even though some comparisons did not reach statistical significance, effect sizes were medium to large in some cases, supporting that the study was underpowered to answer these questions.

Another limitation is inherent in all CSF studies: CSF only approximates events in the brain. Despite this, CSF has provided many important insights into brain events in HIV and other diseases [51]. A high degree of HIV DNA compartmentalization within the CSF suggests that the sampled HIV DNA is originating from brain tissue, but it could also reflect a population of cells that preferentially migrate into CSF from blood. This will need to be evaluated in future studies using larger cohorts and post-mortem brain tissues. Finally, in this study, we were also not able to determine if the HIV DNA population sampled in the CSF is replication competent.

Despite these limitations, our data provide a unique perspective by analyzing HIV DNA populations sampled using CSF prospectively collected from a unique cohort of individuals who started ART and with known EDI. Our study supports the idea that initiation of ART during early infection may limit the diversity of HIV populations and inflammation in CNS. Future studies may want to evaluate the CSF HIV DNA populations in bigger cohorts and include longitudinal assessments prior and after initiation of ART to characterize dynamics of the CNS as a HIV reservoir. Moreover, future studies need to assess the CNS replication competent HIV DNA populations. The presence of unique HIV DNA populations within the CSF during ART might be relevant for future eradication strategies.

**Material and Methods**

**Ethics statement**

The study was approved by the Institutional Review Board at the University of California. All adult participants (age ≥ 18 years) provided written informed consent. No children were included in this study.
Study cohort

Study participants were selected among HIV-infected men who enrolled in the SD PIRC between 2001 and 2012 and were still engaged in follow-up [52]. All SD PIRC participants are recruited during primary infection and followed with longitudinal blood drawn. Per protocol, visits occur at weeks 1, 2, 4, 8, 12, and 24, and then every 24 weeks thereafter. The date of infection is estimated for each participant following an established algorithm (summarized in supplementary S4 Table) [36]. Although early ART initiation is encouraged for all SD PIRC participants, implementation is based on participants’ personal decision, primary care physician input and following the current ART guidelines at the time of recruitment. Participants started ART between 2003 and 2012. Selection criteria for this study were: (1) HIV-infected males recruited during primary infection, (2) started ART during follow-up early or later during HIV-infection, (3) reached undetectable HIV RNA in blood plasma (<50 HIV RNA copies/ml) and remained undetectable during follow-up until the time of baseline CSF collection (based on our longitudinal viral loads and participant self-report) [53]. None of the participants had evidence of other inflammatory neurologic disorders or pleocytosis.

Participants were divided in early ART versus late ART groups as follow: 9 were included in the early ART group (>4 months from estimated date of infection [EDI]) and 7 in late ART group (>14 months from EDI).

Sample collection and processing

Paired blood and CSF samples were collected from each HIV-infected participant cross-sectionally. A subset of 2 participants provided a second pair of samples (3 and 5 months after their first evaluation, respectively) and one participant provided a third pair of samples (2 months thereafter).

We designed our study to maximize cellular recovery by collecting 40 ml of CSF fluid by lumbar puncture. Following standard procedures at the HIV Neurobehavioral Research Center (HNRC), the LPs were performed using atraumatic needle by an experienced physician. None of our study participants reported any complication following the CSF collection. From this larger volume, we obtained a CSF cell pellet and split it into two separate aliquots. Cell pellet lysates (containing HIV DNA) were used for ddPCR and for C2V3 env nested PCR as described below (see supplementary S1 Fig).

CSF supernatant was used to measure levels of selected markers of inflammation and neuronal damage (described below) and to measure HIV RNA by Aptima HIV RNA assay (Hologic), after concentrating 5 ml of supernatant (with single copy sensitivity). The CNS penetration effectiveness (CPE) index for the most recent ART regimen was determined as previously described [54]. For all participants, blood CD4+ T-lymphocytes were measured by flow cytometry (CLIA certified local laboratory). Levels of HIV RNA in blood plasma were quantified by the Amplicor HIV Monitor Test (Roche Molecular Systems Inc.).

Neurocognitive functioning

For each participant, neurocognitive functioning was assessed using a standardized clinical battery of seven ability areas consistent with Frascati recommendations for neuroAIDS research [55] and summarized using the validated global deficit score (GDS) [56].

Levels of inflammation and neurofilament light chain

The levels of selected markers of monocyte activation (sCD163), general inflammation (IL-6) and (TNF-α) and monocyte trafficking monocyte chemoattractive protein (MCP)-1 as well as
brain damage (NFL chains were measured in all participants. Enzyme-linked immunosorbent assay (ELISA) was used to quantify the levels of sCD163 (Trillium Diagnostics, Brewer, ME, USA) from blood plasma and CSF, and NFL in CSF (Uman Diagnostics, Sweden). Electrochemiluminescence multiplex assay (Meso Scale Diagnostics, Rockville, MD, USA) was used to quantify the levels of IL-6, TNF-α and MCP-1 in CSF and blood plasma. All assessments were performed according to the manufacturer’s procedures.

**Quantification of HIV DNA from PBMC and CSF cellular pellets**

Genomic DNA was extracted from 5 million PBMC for each participant (QIAmp DNA Mini Kit, Qiagen, CA) per manufacturer’s protocol. Genomic DNA was also extracted from 1 (out of 2) aliquot of cell pellets obtained from 20 mL of CSF (in average, there were 20,000 white blood cells/aliquot, range: 20,000–60,000) using direct lysis as previously described [22, 23]. Levels of HIV DNA (pol gene region: HXB2 coordinates 2536–2662) were measured in triplicate by (dd)PCR [57]. Briefly, 5 μL of 1:2 diluted CSF lysates or 1000 ng of DNA from PBMC per replicate was digested with BANII enzyme (New England Biolabs) prior to ddPCR. Reactions were performed with the following cycling conditions: 10 minutes at 95˚C, 40 cycles consisting of a 30 second denaturation at 94˚C followed by a 60˚C extension for 60 seconds, and a final 10 minutes at 98˚C. For DNA from CSF cell pellets, we used 5 μL (diluted 1:2) of lysate per replicate. A 1:10 dilution of the digested DNA was used for host cell RPP30 (ribonuclease P30) ddPCR and cycled with same parameters described above. Copy numbers were calculated as the mean of the three PCR replicates measurements and normalized to one million of cells (PBMC or CSF cells) as determined by RPP30 levels (total cell count). The limit of detection of the ddPCR assay for HIV DNA using the same primer-probe set was previously described as 0.7 copies per million of cells [57]. The detected number of RPP30 copies in each ddPCR reaction was used to estimate the number of cells per aliquot of CSF cellular pellet.

**Viral sequencing, sequence filtering and bioinformatics analysis**

We amplified the HIV-1 env C2-V3 (HXB2 coordinates 6928–7344) region from DNA extracted from CSF cellular pellets and PBMC by nested PCR using specific primers [58]. Sequencing was performed using NGS 454 GS FLX Titanium (454 Life Sciences, Roche, Branford, Connecticut, USA). Read (FASTA) and quality score files produced by the 454 instruments were further analyzed using a purpose-built bioinformatics pipeline [25–27]. The pipeline is available at https://github.com/veg/HIV-NGS and the key steps were summarized briefly below:

Raw data were filtered by removing sequences of low quality (q-score of less than 15) using the Datamonkey analysis tool [59] and aligned to a subtype B reference sequence [60]. High-quality reads were retained and aligned to HXB2 as a reference sequence (without generation of contigs) using an iterative codon-based alignment procedure implemented in Datamonkey. A Bayesian Dirichlet mixture of multinomials probabilistic model was used to distinguish sequencing error from true low-frequency variants (posterior probabilities of ≥99.99%). For PBMC, we obtained a median of reads of 16927.5 [13725, 23106.5] and for CSF, we obtained a median of reads of 16198 [9590, 20157.5]. All sets of representative reads were screened for evidence of recombination using GARD [29], APOBEC signatures, hypermutations and frame-shifts as part of our pipeline procedure. All sequences were screened for in-house cross-contamination using BLAST [61]. Identical sequence reads were clustered, allowing identification of non-redundant sequences. A minimum of 10 identical sequence reads were clustered into haplotypes, and the proportion of reads in each haplotype was provided. Hence, the output consists of a list of representative haplotypes and their relative frequencies. The average number of HIV DNA haplotypes recovered from the CSF is 21 (range: 11–29), while 27 (range:
9–46) haplotypes were recovered from blood. For each sample, we computed the mean of all pairwise Tamura-Nei 93 distances between reads with at least 100 overlapping base pairs to quantify nucleotide diversity [62].

Compartmentalization analysis

Viral compartmentalization was first assessed by the Fst approach defined as $F_{ST} = 1 - \frac{\pi_I}{\pi_D}$, where $\pi_I$ is the estimate of mean pairwise intra-compartment genetic distance (TN93) [28], and $\pi_D$ is its inter-compartment counterpart [63]. Both quantities were computed by comparing all reads from blood and CSF compartments, subject to the requirement that they share at least 150 aligned nucleotide positions. The large number of pairwise comparisons ($10^7$–$10^9$) was handled computationally using an efficient implementation of the TN93 distance calculator (github.com/veg/tn93), which achieves a throughput of $10^7$ distances/second on a modern multi-core desktop. Subsequently, to guard against inference of compartmentalization by skewing of allelic frequencies due to PCR amplification and other biases, we recomputed $F_{ST}$ by discarding copy number counts for read clusters (i.e. each cluster was counted as having only one sequence), i.e. all haplotypes are assigned a relative weight of 1. Statistical significance of both tests was derived via 1,000 population-structure randomization/permutation test. Finally, we performed a second tree-based Slatkin-Maddison (SM) test for compartmentalization [64]. Conservatively, we defined a CSF sample as compartmentalized only if all of the following tests were consistent and significant: (1) distance based $F_{ST}$ test, (2) sensitivity test $F_{ST}$ with collapsed haplotypes and (3) tree-based SM test.

Phylogenetic analysis of partial $env$ HIV-1 sequences

Viral haplotypes were realigned using MUSCLE [30], piped to FastTree 2 [31] for maximum likelihood trees reconstruction, and subjected to codon-based (MG94) phylogenetic analyses in HyPhy [32].

Statistical analysis

Statistical differences between groups (early versus late ART initiation) were examined for statistical significance using linear mixed-effects models with individuals included as random intercepts. The time-to ART variable was dichotomized or log transformed, and outcome variables were rank-transformed when appropriate. When residual variance differed by a specific factor in analyzing untransformed outcomes, we allowed heterogeneous variances across levels of that factor. Differences for sparse variables were detected by Fisher exact test. Whenever possible, partial $\eta^2$ ($\eta^2_p$) was provided as a measure of the strength of association. Statistical analyses were performed using the R statistical language ver 3.3 [65] and the nlme package [66].

Supporting Information

S1 Fig. CSF sample processing workflow. Forty milliliters of CSF were collected from HIV-infected people by lumbar puncture and CSF cells were pelleted down. CSF supernatant was stored at -80°C and subsequently used to measure levels of inflammation and neuronal damage as well as HIV RNA using a Single Copy Assay (Hologic). Cells were resuspended in freezing media and divided into two aliquots stored at -150°C. After thawing, cells were washed with PBS and centrifuged to obtain a dry CSF cell pellet. Cell pellets were directly lysed with lysis buffer and DNA was quantified by ddPCR and amplified by C2V3 Nested PCR. (TIFF)
S1 Table. Summary of positive samples by ddPCR and C2V3 env Nested PCR.

S2 Table. Data summary for PBMC sequences.

S3 Table. Data summary for CSF cellular pellets sequences.

S4 Table. Brief Summary of the Algorithm for Computing the Estimated Date of HIV-1 Infection.

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