Cerebrospinal fluid soluble CD30 elevation despite suppressive antiretroviral therapy in individuals living with HIV-1

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Abstract

Objectives: The aim of this study was to assess soluble CD30 (sCD30), a protein that colocalises with HIV-1 RNA and DNA in lymphoid cells and tissues, in cerebrospinal fluid (CSF) as a marker of HIV-1 infection in the central nervous system (CNS).

Methods: This was a cross-sectional study using archived samples from two clinical cohorts. Soluble CD30 concentrations were measured in paired CSF and plasma from untreated viraemic individuals (n=52), individuals on suppressive antiretroviral therapy (ART) (n=33), HIV-1 controllers (n=10), participants with CSF HIV-1 ‘escape’ (n=11) and controls without HIV-1 infection (n=16). Nonparametric tests were used to compare levels across groups and evaluate correlations with HIV-1 RNA, CSF neurofilament light chain protein (NFL) and neopterin.

Results: Compared with controls (median 30 ng/mL, interquartile range [IQR] 23–50), plasma sCD30 levels were elevated in viraemic participants (75 ng/mL, 52–116; P<0.001), but not in those on suppressive ART (38 ng/mL, 32–62). In contrast, CSF sCD30 levels were elevated in ART-suppressed individuals (34 ng/mL, 19–46; P=0.001) and in those with CSF ‘escape’ (33 ng/mL, 27–40; P=0.004) compared with controls (18 ng/mL, 11–23), but not in untreated viraemic individuals. No association was observed between CSF sCD30 and plasma HIV-1 RNA, concurrent or nadir CD4+ T cell count, duration of infection or plasma sCD30. CSF sCD30 correlated with CSF NFL (r=0.34, P=0.001).

Conclusions: In contrast to plasma, sCD30 levels are elevated in the CSF of individuals with HIV-1 infection who are on suppressive ART. Elevated levels of sCD30 in the CSF may be an indicator of persistent CNS HIV-1 infection, although the mechanism underlying this elevation warrants further investigation.

Keywords: HIV-1, reservoir, central nervous system (CNS), cerebrospinal fluid (CSF), CD30

Background

Although antiretroviral therapy (ART) has substantially reduced the morbidity and mortality rates of HIV-1 infection, the persistence of viral reservoirs continues to prevent the elimination of HIV-1 from all tissue compartments [1,2]. Consequently, attention has turned to detecting cells harbouring either latent or transcriptionally active HIV-1 in order to better characterise the HIV-1 reservoir and identify targets for viral eradication strategies.

CD30 is a member of the tumour necrosis factor (TNF) receptor superfamily [3–5] that is rarely expressed in healthy individuals [5–8]. Although CD30 is primarily expressed on lymphocytes, there is evidence that myeloid cells are also capable of expressing CD30 in vitro and in vivo [9–12]. Infection with viral pathogens including HIV-1, human T cell lymphotropic virus and Epstein–Barr virus may increase CD30 surface expression [4,13–15]. In some individuals with HIV-1, CD4+ T lymphocyte–associated HIV-1 RNA and DNA are highly enriched in cells expressing CD30 and surface expression colocalises with HIV-1 transcriptional activity in gut-associated lymphoid tissues [16]. In individuals on ART, depletion of cells expressing CD30 using a cytotoxic antibody–drug conjugate appears to reduce HIV-1 DNA and RNA in peripheral blood mononuclear cells (PBMCs) [17]. Taken together, these findings suggest that CD30 could be a marker of cells that harbour HIV-1 infection.

Surface CD30 is cleaved by metalloproteases such as TNF-α converting enzyme and released in a soluble form (sCD30) [18]. Plasma sCD30 concentrations are elevated in individuals with untreated HIV-1 infection but normal in those on suppressive ART [3,16,19–21]. This result suggests that increased sCD30 levels may be due to persistent viral replication or T cell activation within the peripheral blood compartment.

Beyond the long-lived CD4+ T cell reservoir [22], the central nervous system (CNS) may serve as an additional source of persistent infection in people living with HIV-1 [23–25]. Lymphoid cells, the primary target of HIV-1 infection, can travel into the CNS compartment [26]. HIV-1 can also infect CNS-resident myeloid cells including macrophages and microglia [27,28]. The extent to which cells of lymphoid or myeloid origin contribute to persistent HIV-1 in the setting of suppressive therapy and whether persistent CNS infection underlies continued low-level CNS inflammation and neurocognitive impairment despite ‘undetectable’ CSF HIV-1 RNA [24,27–30] remains unclear.

Direct access to CNS tissue in individuals living with HIV-1 is generally not feasible [25]. As an initial approach to evaluating sCD30 expression in CNS HIV-1 infection, we measured CSF sCD30 concentrations across a clinically relevant spectrum of
individuals and examined their correlation with plasma sCD30 levels, measures of HIV-1 replication and progression, and CSF biomarkers of inflammation and axonal injury. We hypothesised that the dynamics of sCD30 concentrations in the CSF would parallel those previously described in plasma, with viraemic individuals having elevated sCD30 levels and ART-suppressed individuals having sCD30 levels similar to uninfected controls [16,17]. We found this not to be the case.

Methods

Study design and sample

This was a cross-sectional study using archived CSF and plasma samples from a total of 134 individuals from two clinical centres, one in Gothenburg, Sweden (n = 62), and the other in San Francisco, California (n = 72). All participants had been enrolled in study protocols collecting CSF and plasma samples by lumbar puncture and phlebotomy using standard procedures. The samples were stored at −80°C [31,32]. We selected matching CSF and plasma samples from controls without HIV-1 infection and several clinically relevant groups with HIV-1 infection, including (1) untreated viraemic individuals prior to initiation of ART stratified by CD4+ T lymphocyte count; (2) ART-treated individuals who had been on suppressive therapy for at least 1 year and achieved virological suppression (plasma and CSF HIV-1 RNA below the limit of quantitation on standard clinical assays); (3) HIV-1 controllers exhibiting spontaneous viraemic (plasma viral load <500 copies/mL) or ‘elite’ control (plasma viral load below the limit of quantitation on standard clinical assays); and (4) individuals with both symptomatic and asymptomatic CSF HIV-1 ‘escape’, [33,34] defined as CSF HIV-1 RNA >50 copies/mL in the presence of plasma HIV-1 RNA <50 copies/mL (if suppressed) or CSF HIV-1 RNA greater than plasma HIV-1 RNA (if not fully suppressed) [35]. We excluded individuals with concomitant CNS infections, including neurosyphilis.

Specimen sampling, processing and analysis

Soluble CD30 levels were measured in CSF and plasma samples using the Human sCD30 Platinum ELISA Kit (Affymetrix eBioscience) according to the manufacturer’s instructions. Because this kit was designed for plasma or serum, we tested CSF specimens in duplicate, and values were averaged to obtain an adjusted CSF sCD30 value for each specimen. Quantifiable results were within the linear range of the assay. When variation between duplicates on the assay exceeded 20% (20/134 samples), the test was repeated in duplicate and the new values, if concordant, were used. If variation continued to exceed 20%, the adjusted CSF sCD30 value was calculated as the mean of the four values. Plasma sCD30 was also tested in duplicate.

Data on standard HIV-1 parameters (CD4+ T lymphocyte count, plasma HIV-1 RNA) and CSF parameters (white blood cell [WBC] count, albumin ratio, CSF HIV-1 RNA) from clinical testing were available for all individuals from paired time points. The limit of quantification of the plasma and CSF HIV-1 RNA assay was 40 copies/mL for the San Francisco site and 20 copies/mL for the Gothenburg site; we used 40 copies/mL as the limit of quantification in our analyses. Correlative data for CSF neurofilament light chain protein (NFL) were determined by immunoassay using published methods [32]. Because CSF NFL varies with age, we used a standard age correction to compare across groups and designate a consistent threshold for abnormality as previously described [36]. Neopterin concentrations were measured using a commercially available immunoassay (NEOPT-SCR.EIA 384 Det., Thermo Fisher Scientific – BRAHMS GmbH, Henningsdorf, Germany) [37].

Antibody staining and flow cytometry

Five participants were willing to provide fresh CSF and matched PBMCS for flow cytometric analyses. These samples were analysed by a multiparameter flow cytometric panel that included CD30 expression on CD4+ T cells. Full panels and antibodies used for staining are described in Supplementary Table 1. The flow cytometric panel allowed for the measurement of CD30 expression on CD4+ T lymphocytes as well as cells of myeloid origin (expressing CD14 or CD16). Samples stained with all but one fluorescent antibody were used in addition to unstained controls to adequately gate on the low-frequency cell populations while accounting for spectral overlap, even after rigorous compensation. Cells were analysed on a BD LSR II (BD Biosciences), and data were analysed in FlowJo V10 (Tree Star). Single stained beads (Life Technologies) were used for compensation. Sample gating is shown in Supplemental Figure 1. Because of the low frequency of CD30 expression and the relatively few numbers of cells that can be obtained from CSF, further characterisation of CD30+ lymphocyte or myeloid cell populations was limited.

Statistical methods

The primary outcome variable was the concentration of sCD30 in CSF specimens in the different clinical groups. Descriptive statistics (median, interquartile range [IQR]) were derived for each group and compared using Mann–Whitney or Kruskal–Wallis tests with Dunn’s pairwise comparisons where appropriate. Correlations between CSF sCD30 and other variables used nonparametric Spearman statistics. For most analyses, we evaluated correlations in the cohort as a whole and within the discrete clinical subgroups to explore relationships that might provide mechanistic clues between correlates in specific clinical scenarios. We used Stata release 15.1SE (StataCorp, LLC) to conduct all analyses and GraphPad Prism version 8.1.2 (GraphPad Software) to display results.

Study approval

The samples and related background data were all obtained between 2000 and 2018 within the context of research protocols approved by the institutional review boards of the two study sites. Informed consent was obtained from all subjects.

Results

Characteristics of participants

Table 1 describes the demographic, HIV-1 and CSF parameters of the study participants by group. There were no site-based differences in gender or race demographics, plasma or CSF HIV-1 RNA, or CD4+ T cell count; participants from San Francisco were statistically, but not substantially, older (median 48 vs 42 years, P < 0.001). The majority of participants in all groups were male, and age and race demographics were similar across groups. All participants in the ART-suppressed group had been on suppressive therapy for at least 1 year, although information regarding the composition of the ART regimen and the duration of suppression was not available. In addition to higher median plasma HIV-1 RNA, untreated viraemic individuals had higher CSF HIV-1 RNA levels, higher CSF WBC counts and lower CD4+ T lymphocyte counts at the time of the study. CSF blood albumin ratio, an index of blood–brain barrier permeability, did not differ between groups. CSF neopterin and NFL were both higher in the untreated viraemic group and in the CSF ‘escape’ group.
Table 1. Characteristics of study participants. Values reported as median (interquartile range) unless otherwise indicated

| Characteristics                  | HIV-uninfected | Untreated viraemic | ART-suppressed | Controller | CSF escape |
|----------------------------------|----------------|-------------------|---------------|------------|------------|
| Age (years)                      | 46.2 (36.8–54.8) | 43.2 (35.7–50.0)  | 47.6 (42.4–53.2) | 37.4 (30.5–44.2) | 48.2 (35.6–51.4) |
| Male sex (%)                     | 15 (94)        | 44 (85)           | 28 (85)       | 7 (70)     | 7 (64)     |
| Ethnicity (%)                    |                |                   |               |            |            |
| White                            | 7 (44)         | 30 (58)           | 22 (67)       | 3 (30)     | 5 (45)     |
| Black                            | 7 (44)         | 17 (33)           | 6 (18)        | 6 (60)     | 4 (36)     |
| Asian                            | 0 (0)          | 3 (6)             | 4 (12)        | 1 (10)     | 2 (18)     |
| Other/unknown                    | 2 (12.5)       | 2 (4)             | 1 (3)         | 0 (0)      | 0 (0)      |
| Nadir CD4+ T cell count           | n/a            | 230 (60–360)      | 207 (61–310)  | 435 (320–624) | 130 (110–170) |
| Current CD4+ T cell count         | 827 (764–940)  | 284 (97–423)      | 601 (507–702) | 568 (397–1100) | 400 (290–480) |
| Plasma HIV-1 RNA (copies/ml)      | n/a            | 41,050 (9240–186,500) | <40 (<40 to <40) | 183 (43–367) | <40 (<40 to <40) |
| CSF parameters                   |                |                   |               |            |            |
| White blood cells (cells/mm³)     | 1 (1–3)        | 5 (0–10)          | 2 (0–3)       | 1 (0–3)    | 10 (2–28)  |
| Albumin ratio                    | 6.0 (4.6–8.7)  | 5.2 (3.6–7.0)     | 4.3 (3.4–6.6) | 4.6 (2.9–5.9) | 5.2 (3.7–7.3) |
| HIV-1 RNA (copies/ml)            | n/a            | 1368 (370–7515)   | <40 (<40 to <40) | <40 (<40 to <40) | 605 (134–860) |
| Neopterin (mmol/L)               | 6.3 (4.3–8.6)  | 16.7 (9.5–24.3)   | 6.3 (4.9–9.8) | 6.3 (5.5–9.6) | 40.5 (22.5–80.2) |
| Neurofilament (age-adjusted; ng/L)| 511 (443–639) | 527 (359–834)     | 475 (357–632) | 405 (328–511) | 794 (544–1132) |

*P-values calculated using Kruskal–Wallis test evaluating whether all subgroups derive from same distribution. ART: antiretroviral therapy; CSF: cerebrospinal fluid; n/a: not applicable.

Plasma sCD30 levels are elevated in viraemic individuals and normal in participants on ART

Figure 1a shows plasma sCD30 levels in the various subgroups. Consistent with prior studies [3,16], plasma sCD30 levels were significantly elevated in viraemic participants (median 74.6 ng/mL, IQR 52.2–116.8) than in controls without HIV-1 infection (median 30.4 ng/mL, 22.5–48.5; P<0.001). Individuals on suppressive ART (median 38.1 ng/mL, iQr 32.1–62.3) had lower levels of plasma sCD30 compared with viraemic participants (P=0.001), and no significant difference was detected between this group and controls without HIV-1 infection. Individuals with CSF ‘escape’ and controllers did not have elevated plasma sCD30 levels.

Among untreated viraemic individuals, plasma sCD30 correlated with plasma HIV-1 RNA (r=0.32, P=0.019) and CD4+ T lymphocyte count (r=–0.26, P=0.05). Plasma sCD30 levels by CD4+ T cell count subgroup are summarised in Figure 2a.

CSF sCD30 levels are normal in viraemic individuals and elevated in participants on ART

Figure 1b shows CSF sCD30 levels in the various subgroups. Unexpectedly, we did not detect an elevation in CSF sCD30 in untreated viraemic participants compared with uninfected controls. Three individuals in the untreated viraemic subgroup with HIV-1-associated dementia (HAD) had relatively low CSF sCD30 levels (18.5, 28.7, 16.1 ng/mL, respectively) despite high levels of CSF HIV-1 RNA.

Unlike in plasma, and despite a lack of elevation in viraemic participants, CSF sCD30 levels were elevated in ART-suppressed individuals (median 33.8 ng/mL, IQR 19.1–46.1) compared with uninfected controls (median 19.4 ng/mL, IQR 11.4–26.9; P=0.001). A difference was also detected between ART-suppressed and untreated viraemic individuals (P=0.004).

In a sub-analysis of 10 individuals exhibiting CSF HIV-1 ‘escape’, CSF sCD30 levels were elevated (median 33.3 ng/mL, IQR 27–40). This elevation was significant when compared with controls without HIV-1 infection (P=0.004), but not with those on suppressive ART.

The controllers included in this study were a heterogeneous group that included individuals without exposure to ART who were naturally aviraemic (n=2) or exhibited low-level plasma viraemia (n=8). CSF sCD30 levels were not elevated in the combined population of controllers when compared with controls without HIV-1 infection.

Correlates of CSF sCD30

No correlation was detected between plasma and CSF sCD30 levels in the cohort or in any of the subgroups.

CSF sCD30 did not correlate with duration of infection, nadir CD4+ T lymphocyte count, plasma HIV-1 RNA, or concurrent CD4+ T lymphocyte count in the cohort or in any of the subgroups. When untreated viraemic participants were stratified by CD4+ T lymphocyte count, CSF sCD30 levels appeared to peak at moderate (200–499 cells/µL) levels of immunocompromise (Figure 2b), although the differences between groups were not statistically significant.

Within the ART-suppressed group, CSF sCD30 correlated with CSF WBCs (r=0.36, P=0.04; Figure 3a), although this association was heavily influenced by one data point and was not detected in the other cohorts. CSF sCD30 correlated with age-adjusted CSF NFL, a marker of axonal injury [36], within both the total


![Figure 1](image_url)

Figure 1. (a) Plasma sCD30 in HIV-uninfected, untreated viraemic, ART-suppressed, CSF ‘escape’ and controller subgroups. (b) CSF sCD30 in HIV-uninfected, untreated viraemic, ART-suppressed, CSF ‘escape’ and controller subgroups. I indicates statistically significant comparison between group of interest and untreated viraemic subgroup. II indicates statistically significant comparison between group of interest and ART-suppressed subgroup. ART: antiretroviral therapy; CSF: cerebrospinal fluid.

Our study suggests a source of CSF sCD30 within the CNS. The lack of association between plasma and CSF sCD30 and the CSF:sCD30:albumin index suggests scD30 production in the CNS compartment.

CD30 surface expression may be present on CSF myeloid cells

Five volunteers were willing to provide fresh PBMC and CSF specimens for analysis. Analysis of fresh PBMCs demonstrated the presence of a small population of CD30+ CD4+ T lymphocytes in the peripheral blood, consistent with previous work [16]. As seen in prior analyses of individuals with HIV-1 [16], CD30 expression on PBMCs was primarily on CD4+ T cells (0.19%, 0.13%–0.45%) rather than on myeloid cells (0.008%, 0.001%–0.13%). In contrast, although rare in the CSF, a higher proportion of CD4+ expressing cells were of myeloid origin (0.07%, 0%–0.47%) than CD4+ T lymphocytes (0.02%, 0.001%–0.03%). We were unable to perform further phenotyping of either lymphocyte or myeloid cell populations in the CSF given the rarity of CD30 expression and relatively few cells from CSF collection (8508, 7938 to 39,154 total live cells).

Discussion

This study examined soluble CD30 in the CSF across clinically relevant groups of individuals with HIV-1. In the peripheral blood compartment, sCD30 is known to be associated with persistent viral replication and immune activation [3,16,19–21]. We hypothesised that the CSF dynamics of this sCD30 marker would mirror those in the peripheral blood and that levels of CSF sCD30 would normalise in individuals on ART [16]. Instead, we found that the dynamics of sCD30 differ between the CNS and peripheral blood compartments. Despite suppressive ART, CSF sCD30 may represent ongoing HIV-1 transcriptional activity (and potentially viral replication) or bystander cell activation associated with myeloid or lymphoid cells in the CNS. These findings support the need for additional studies to delineate the location of the CNS cell type(s) responsible for producing CD30 and to determine whether these cells harbour HIV-1 genetic material.

Our study suggests a source of CSF sCD30 within the CNS. The lack of association between plasma and CSF sCD30 and the CSF:sCD30:albumin index support the likelihood of sCD30 production by a cell present within the CNS compartment and argue against diffusion of this protein across an inflamed blood–brain barrier. Metalloproteases like TNF-α converting enzyme have been identified in the CSF of individuals with cognitive impairment, suggesting that these enzymes are available in the CNS compartment to cleave surface-expressed CD30 [38].
peripheral blood compartment, suppressive ART is associated with decreased levels of plasma sCD30 [16], which was also the case in this study cohort. The presence of elevated CSF sCD30 concentrations in the ART-suppressed group suggests that certain HIV-1-specific phenomena, such as persistent activation of infected cells or potentially low-level viral replication (i.e., below the limit of quantification), may be present within the CNS compartment despite suppressive ART. Whether ART is mitigating the cytopathic
effects of HIV-1 on CD30+ cells is unknown, but such a mechanism would be compatible with the CSF sCD30 elevations observed in individuals with low-level HIV-1 RNA replication described earlier. However, ongoing CNS replication during suppressive ART is contradicted by several previous findings. There is no evidence of viral evolution or escape with resistant virus from the CNS, which would be expected after long-term treatment if low-level replication were present. Furthermore, treatment intensification studies have not shown any effect on CSF residual viral levels or inflammation [39,40].

The second unexpected finding of this study was the lack of CSF sCD30 elevations in ART-naive viremic individuals, including participants with HIV-1-associated dementia with known high viral burden in the brain. Plasma sCD30 was elevated in the untreated viremic group and correlated with plasma HIV-1 RNA, consistent with prior studies [3,19]. In contrast, CSF sCD30 was not elevated in viremic individuals, and the association between CSF sCD30 and HIV-1 RNA was strongest in those with low levels of CSF HIV-1 RNA (i.e. CSF ‘escape’). Individuals with the greatest neurological morbidity (HAD) and highest levels of CSF HIV-1 RNA had comparatively low CSF sCD30. This pattern could be seen if greater levels of HIV-1 replication resulted in higher turnover of a CD30-producing cell population owing to cytopathic effects or chemotaxis out of the CNS compartment. Alternative possibilities include decreased CD30 cleaving by metalloproteases or active trafficking of the sCD30 product out of the CNS in the setting of productive infection. Further investigation of these specific mechanisms is warranted.

Although not statistically significant, the pattern of an apparent CSF sCD30 peak at a moderate level of immune dysfunction is consistent with earlier work. A similar dynamic has been suggested in prior work looking at T cell responses across levels of plasma viraemia [41] and in unrelated studies of CSF sCD30 in multiple sclerosis, which have found that CSF sCD30 levels may be elevated with the moderate levels of immune activation characterising remission rather than the extreme levels of immune activation associated with relapse [42,43].

Within the ART-suppressed individuals, a relatively strong association was detected between CSF sCD30 and NFL, an established marker of axonal injury in HIV-1 infection [36,44,45]. This association suggests that some process, perhaps HIV-1-mediated T cell activation within the CNS compartment, could underlie neuronal damage. General immune activation may also lead to axonal injury [46].

Because the cell type responsible for the production of sCD30 in the CNS remains unclear, we conducted a proof-of-concept study to identify CD30+ cells in the CSF. In blood, the CD30+ CD4+ lymphocytes outnumbered the CD30+ myeloid cells; in the CSF, however, we detected CD30 on both lymphocytes and cells of myeloid origin. Although we observed more CD30 positivity on myeloid cells, our studies were limited by the small number of participants for which fresh CSF cells were available and the limited number of cells available within each CSF sample; further characterisation of the lymphocyte and myeloid cell populations was therefore not possible. Nonetheless, these data support the need for further in vivo and in vitro study of cellular sources of CD30 in the CNS. Of note, earlier experimental studies using cells from individuals without HIV-1 have shown the capacity for monocytoids and microglia to express CD30 [9–12]. Further work exploring CD30 in CNS tissue is needed to identify the cellular source of this soluble marker. In vivo studies are challenging given the very few numbers of cells that can be obtained from CSF and the inability to directly sample parenchymal brain tissue in living subjects.

This study has several limitations. First, although our sample size was relatively large for studies of this nature, the individual subgroups were small and the selection for individuals distributed across these groups could have introduced unknown biases. Further studies of sCD30 in more homogeneous populations will help to confirm the patterns observed here. Second, detailed HIV-1 treatment history including ART duration and regimen composition was not widely available. This limited our ability to evaluate a relationship between sCD30 and duration of ART, although no clear pattern emerged in the five individuals with two CSF samples. Further studies should evaluate how the composition or duration of different ART regimens might impact this marker. Third, it is possible that other unmeasured infectious processes within the CNS, such as Epstein–Barr virus reactivation, may have led to increased sCD30 production. However, elevations would be expected within individuals with lower CD4+ T cell counts and active viraemia, which were not observed. Fourth, because our study was meant to generate hypotheses for further investigation, we did not control for multiple statistical comparisons. Finally, the CSF flow cytometry studies were limited by the number of participants willing to provide samples and the number of cells that we could obtain. Evaluation of brain tissue was beyond the scope of this study. In the future, a longitudinal study evaluating individuals with prolonged suppression or with very low-level CSF HIV-1 RNA would further elucidate the dynamics and role of sCD30.

This study of sCD30 in the CSF of individuals with HIV-1 demonstrated that individuals on suppressive ART may have a persistent CNS source of sCD30. It also revealed important differences in the dynamics of sCD30 between the CNS and peripheral blood and suggested several potential mechanisms by which this could occur. Further work to confirm these findings and investigate the potential cellular source of sCD30 is warranted. Such studies have the potential to provide information on the presence of persistent foci of HIV-1 in the setting of suppressive ART and to help direct interventions aimed at inducing long-term suppression of HIV-1.

Acknowledgements

We are grateful to the participants in this research. This work was supported by the following grants: NIH/NIMH R21 MH116716 (PI: Henrich); NIH/NINDS R01 NS094067 (PI: Price); NIH/NAID T32 AI60530-12; the Swedish State support for Clinical Research; and ALFGBG-717531 (PI: Gisslén).

Contributions

TJH conceptualised the study. MJP, MG, RWP and TJH designed the project and obtained funding. VMA, SS, MG, SGD, and RWP managed the clinical cohorts and obtained participant samples. MJP, CT, CAP, LEH, PJD, CD, DF, HZ, MG, RWP and TJH performed or oversaw the laboratory assays. MJP, SS, PJD, CD, DF, HZ, MG, RWP and TJH analysed the data. MJP, RWP and TJH drafted the manuscript, which was reviewed and edited by all authors.

Conflicts of interest

MJP, CT, LEH, CP, VMA, SS, CD, PJD and DF have no conflicts of interest to declare. HZ has served on scientific advisory boards for Roche Diagnostics, Samumed, CogRx and Wave. HZ is also a co-founder of Brain Biomarker Solutions in Gothenburg AB, a GU Ventures-based platform company at the University of Gothenburg. MG served on the scientific advisory board of Gilead Sciences, BMS, Janssen, MSD and GSK/ViiV Healthcare; received speaker honoraria from Gilead, BMS and Janssen; and received research support from Gilead and Janssen. SGD has received grant support from Gilead and Janssen.
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Supplementary Table 1. Flow cytometric antibodies

| Antibody | Host species | Clone | Fluorochrome | Supplier |
|----------|-------------|-------|--------------|----------|
| CD30     | Mouse       | OKT4  | BV421        | BioLegend |
| CD8      | Mouse       | HIT8A | APC-Cy7     | BioLegend |
| CD3      | Mouse       | UCHT1 | FITC         | BioLegend |
| CD14*    | Mouse       | MSE2  | BV650        | BioLegend |
| CD16*    | Mouse       | 3G8   | BV650        | BioLegend |
| CD19     | Mouse       | HIB19 | BV785        | BioLegend |
| CD11B    | Mouse       | ICRF44| PE           | BD Biosciences |
| CD30     | Mouse       | BerH8 | APC          | BD Biosciences |
| LIVE/DEAD™ |            |      | Aqua         | Invitrogen |
| Brilliant Stain |        |      |              | BD Biosciences |

* The same channel was utilised for CD14 and CD16.
Supplemental Figure 1. Sample gating strategy for surface CD30 measurement.