Functional Compartamentalization of Antibodies in the Central Nervous System During Chronic HIV Infection

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The central nervous system (CNS) has emerged as a critical HIV reservoir. Thus, interventions aimed at controlling and eliminating HIV must include CNS-targeted strategies. Given the inaccessibility of the brain, efforts have focused on cerebrospinal fluid (CSF), aimed at defining biomarkers of HIV-disease in the CNS, including HIV-specific antibodies. However, how antibodies traffic between the blood and CNS, and whether specific antibody profiles track with HIV-associated neurocognitive disorders (HAND) remains unclear. Here, we comprehensively profiled HIV-specific antibodies across plasma and CSF from 20 antiretroviral therapy (ART) naive or treated persons with HIV. CSF was populated by IgG1 and IgG3 antibodies, with reduced Fc-effector profiles. While ART improved plasma antibody functional coordination, CSF profiles were unaffected by ART and were unrelated to HAND severity. These data point to a functional sieving of antibodies across the blood-brain barrier, providing previously unappreciated insights for the development of next-generation therapeutics targeting the CNS reservoir.

Keywords. cerebrospinal fluid; plasma; antibody functions; complement deposition; phagocytosis; NK activation; Fc receptors; ART; HIV-associated neurocognitive disorders.

Acute human immunodeficiency virus (HIV) infection is marked by a rapid burst of systemic viral replication, seeding diverse tissues, including the central nervous system (CNS) [1]. Detection of HIV-RNA within the cerebrospinal fluid (CSF) has been used as evidence of viral infection in the brain [2, 3]. However, whether this represents active infection of CNS resident cells or passage of infected cells remains unclear. Progression of HIV-associated neurocognitive disorders (HAND), despite systemic viral suppression by antiretroviral therapy (ART) [4, 5], has been proposed to be caused by productive infection of brain-resident cells. Moreover, although increased levels of some inflammatory chemokines decline in blood following ART, they remain elevated in CSF [6–9], suggesting the persistence of a neuroinflammatory environment.

Beyond elevated inflammatory chemokines in the CSF, emerging data suggest the importance of virus-specific antibodies as critical biomarkers of disease. Specifically, persistence of high CSF HIV antibodies has been proposed as an indicator of ongoing viral replication in the CNS [10]. However, whether CSF antibodies are transferred from blood and whether they are functionally compartmentalized remains unclear. Here, we applied systems serology to comprehensively dissect humoral immune profiles in the plasma and CSF of chronically infected persons with HIV. Striking brain-specific antibody signatures were identified, marking unique compartmentalization of the humoral response within the CNS.

METHODS

Participants, and Clinical and Biological Markers

Paired plasma and CSF were collected between 2004 and 2012 from 20 persons with HIV infected for less than 5 years: 9 were ART naive, and 11 were virally suppressed (HIV-RNA <200 copies/mL) on ART. Participants were identified from the CNS HIV Antiretroviral Therapy Effects Research (CHARTER) study, which aimed to determine the frequency of HAND in a prospective cohort of over 1000 persons with HIV [11]. All participants provided written informed consent and underwent clinical evaluation, routine laboratory tests (no information about CSF oligoclonal bands was available), and neuropsychological testing. HAND severity was classified based on the Frascati criteria [11, 12] (asymptomatic neurocognitive impairment [ANI], mild neurocognitive disorder [MND], and HIV-associated dementia [HAD]), and assessed using Global Clinical Rating and demographically corrected cognitive test...
scores (T-scores) [12, 13]. A conversion table from T-scores to clinical rating has been previously defined [13].

**Antibody Assays**

Paired plasma and CSF samples were used for antibody testing, in a blinded fashion. Results were obtained from duplicates and by averaging 2 independent experiments. Antibody assays included measurements of pathogen-specific antibody isotype/subclass levels, binding capacity to Fc-receptors (FcRs) and complement, and ability to drive antibody-effector functions, as described below. For all assays, events were detected using a Bio-Plex System (IntelliCyt, iQue Screener Plus).

**Selection of HIV Antigens and Controls**

Antibody responses to 3 different HIV antigens (gp120, IT-001-0024p; gp140, IT-001-0024TMp; and p24, IT-001-020p; ImmuneTech) and non-HIV antigens (herpes simplex virus 1 [HSV1], IT-005-055p; HSV2, IT-005-011p; Epstein-Bar virus [EBV], IT-005-035p, ImmuneTech; cytomegalovirus [CMV], CMV-PENT-100; Cederlane) were assessed. Influenza virus A (IT-003-SW12p and IT-003-001p; ImmuneTech) and Ebola (0501-015; IBT) were used as positive and negative controls, respectively.

**Isotype, Subclass, FcR/Complement Binding Analysis**

HIV-specific and other pathogen-specific antibody isotype, subclasses, and FcR-binding and complement-binding capacity were measured using a customized multiplexed Luminex assay, as reported [14]. Briefly, HIV or control antigens were biotinylated, coupled to red fluorescent neutravidin microspheres, then incubated with plasma or CSF samples (dilution 1:30 or 1:3, respectively). Immune complexes were washed, incubated with guinea pig complement (CL4051; Cedarlane), then washed with 15 mM EDTA. Complement deposition was detected by fluorescein-conjugated goat IgG to guinea pig complement C3b (855385; MP Biomed) and analyzed using a phosphatase-buffered saline level.

**Determination of Antibody-Mediated Functions**

1. Antibody-dependent cellular phagocytosis (ADCP) and neutrophil phagocytosis (ADNP) were assessed using a flow cytometry, microsphere-based phagocytic assay, as described [15, 16]. Briefly, HIV or control antigens were biotinylated and coupled to yellow-green fluorescent neutravidin microspheres (F8776; Thermo Fisher), then incubated with plasma or CSF (dilution 1:100 and 1:10, respectively). Microspheres-bound immunocomplexes were then washed and incubated with a human monocyte cell line (THP-1) to assess ADCP or with human neutrophils obtained from healthy donors’ blood to assess ADNP. Cells in ADNP assays were then stained with anti-CD66b Pac blue antibody (305112; BioLegend) to identify CD66b+ neutrophils. In both ADCP and ADNP assays cells were fixed with 4% paraformaldehyde (PFA) and identified by gating on single cells and microsphere-positive cells. Microsphere uptake was quantified as a phagocytosis score, calculated as the (percentage of microsphere-positive cells) × (MFI of microsphere-positive cells) divided by 100,000. 

2. Antibody-dependent complement deposition (ADCD) assessed the ability of antigen-specific antibodies to bind complement component C3b [17], as described [18]. Briefly, HIV or control antigens were biotinylated, coupled to red fluorescent neutravidin microspheres (F8775; Thermo Fisher), and then incubated with plasma or CSF samples (dilution 1:30 or 1:3, respectively). Immune complexes were washed, incubated with guinea pig complement (CL4051; Cedarlane), then washed with 15 mM EDTA. Complement deposition was detected by fluorescein-conjugated goat IgG to guinea pig complement C3b (855385; MP Biomed) and analyzed gating on single microspheres and C3b+ events.

3. Antibody-dependent natural killer cell activation (ADNKA) quantified the ability of antigen-specific antibodies to activate human NK cells to produce CD107α, IFN-γ, and CCL4 (MIP-1β), as described [19]. Briefly, enzyme-linked immunosorbent assay (ELISA) plates were coated with HIV or control antigens. NK cells were isolated from buffy coats, obtained from healthy blood donors, using RosetteSep NK enrichment kit (15065; Stem Cell Technologies), and rested overnight with interleukin 15 (IL-15), as reported [20]. Antigen-coated ELISA plates were then incubated with plasma or CSF (dilution 1:100 or 1:10, respectively). NK cells were stained with anti-CD107a (555802; BD) and treated with a protein-transport inhibitor (554724; BD Bioscience), and with brefeldin A (B7651; Sigma) to block degranulation. NK cells were then added to the immune-complexes, labeled with surface staining including anti-CD3 (558082; BD) and treated with guinea pig complement (CL4051; Cedarlane), then stained with anti-CD107a (555802; BD) and analyzed gating on single microspheres and CD3+ events.

**Statistical Analyses**

Univariate comparative analyses of antibody features between groups or compartments were performed using nonparametric tests (Mann-Whitney, Kruskal-Wallis, and Friedman test) or
Spearman coefficients in GraphPad Prism with corrections for multiple comparisons.

Multivariate analyses were performed with R software, version 4.0.0. Luminescent data were log_{10}^{+}-transformed. All data were z-scored. Missing values were imputed using R package “DMwR.” A multilevel partial least square discriminant analysis (M-PLSDA) with least absolute shrinkage selection operator (LASSO) was used to select antibody features that contributed most to discriminate plasma versus CSF antibodies [21]. Antibody features were ranked according to their variable importance in projection, for which higher values indicate higher contribution to the model. Five-fold cross validation was performed. The model was validated by comparing it to a null model, for which the modeling approach was repeated 100× with random shuffling of labels (for a total of 10 replicates of cross validation).

RESULTS

Clinical Characteristics
We profiled plasma- and CSF-derived antibodies from a cohort of 20 persons with HIV. Demographics, laboratory values, and clinical and neurocognitive assessments are shown in Table 1. Demographics were similar between the 9 ART-naive and 11 ART-treated individuals, with expected differences in plasma and CSF HIV-RNA levels, CD4⁺ counts, and CSF white blood cell counts (Table 1).

Distinct HIV-Specific Antibody Profiles Across Plasma and CSF
HIV-specific IgG targeting gp120, gp140, and p24 were detected in both plasma and CSF (Figure 1A). However, although all Ig isotypes (IgG, IgM, and IgA) and subclasses (IgG1, IgG2, IgG3, and IgG4, and IgA1 and IgA2), were detected in plasma (Figure 1B–1D, data not shown for IgA), CSF was populated mainly by IgG1 and IgG3 (Figure 1B and 1D), with little IgM (Figure 1C). These data show compartmentalization of antibodies with specific IgG subclasses, and absence of IgM, within the CSF. Generally, antibodies in the CSF are thought to be primarly translocated from (and recycled to) the peripheral circulation via neonatal Fc receptor (FcRn)-mediated transport across the blood-brain barrier (BBB) [22–24]. Alternatively, in disease settings, CSF-antibodies may derive from circulating B cells that gain access to the brain and mature to antibody-
producing plasma cells [25–27]. Thus, our data suggest that HIV-specific antibodies are unlikely to be produced within the CNS, as more variation in isotype and classes would be expected. It is possible that the restricted Ig isotypes and classes observed in the CSF may result from IgG subclass selectivity by FcRn (IgG1 > IgG4 > IgG3 > IgG2) [28, 29], and might explain why IgG1 and IgG3 predominated in CSF, whereas IgG2 and IgM were not detected.

**HIV-Specific Antibodies Are Enriched in CSF Compared to Non-HIV-Specific Antibodies**

To determine whether the observed Ig isotype/subclass exclusion from the CSF was unique to HIV-specific antibodies, or reflected overall differences in humoral compartmentalization across plasma and CSF, we analyzed antibody responses to several additional viruses (influenza virus, HSV1, HSV2, CMV, and EBV) in the same cohort. While diverse isotype/subclass antibody profiles were observed across other virus-specific responses in plasma, CSF antibodies were dominated again by a focused IgG1 response (Figure 2A). However, CSF to plasma ratios for individual IgG isotype/subclasses or IgM highlighted the presence of higher ratios of HIV-specific (all antigens combined) antibodies in CSF, especially for IgG1 (Figure 2B), pointing to an enrichment of HIV-specific compared to other virus-specific antibodies in the CSF. These data are in line with previous observations of persistently high anti-HIV responses in the CSF of persons with HIV [10].
Weak CSF HIV-Specific Antibody Binding to FcRn Suggests Reduced Brain Clearance and Antibody Retention

We next aimed to assess whether the observed CSF-specific antibody-subclass profiles were related to differences in FcRn binding capacity by HIV or other pathogen-specific antibodies. FcRn binding was corrected for total pathogen-specific antibody titers to capture qualitative as well as quantitative differences. We observed strong FcRn binding by HIV-specific plasma antibodies, in contrast to weaker binding by CSF-antibodies (Figure 2C). Interestingly, HIV-specific CSF to plasma FcRn binding ratios were lower than those for influenza virus, HSV1, HSV2, CMV, and EBV (Figure 2D), and these differences were mainly driven by herpes viruses, and in particular CMV and EBV (Supplementary Figure 1).

Overall, these data suggest selective retention within the CNS of HIV-specific antibodies (but not antibodies to other viruses) through reduced FcRn-mediated clearance in the context of active HIV disease.

CSF HIV-Specific Antibodies Are Less Functional Than Those in Plasma

To further understand the level of sieving across the compartments, we next explored the functional characteristics of HIV-specific antibodies across the plasma and CSF. Specifically, we explored differences in the capacity of HIV-specific antibodies to interact with diverse FcRs and to drive antibody-mediated functions (ADCD, ADCP, ADNP, and ADNKA). Given that IgG1 and IgG3 are the most...
functional antibody subclasses in humans [29], we hypothesized that the CSF, enriched in our study of these 2 IgG subclasses, would harbor robust antibody effector functions. However, HIV-specific antibodies in CSF exhibited little to no function compared to plasma antibodies (Figure 3A and 3B), with notably lower levels of ADCD and ADNP in CSF (Figure 3B). This reduced functionality was linked to lower binding to both activating (FcγR2A, 3A, 3B) and inhibitory (FcγR2B) FcRs, at similar levels across all HIV antigens (gp120, gp140, p24) (Figure 3C and 3D). These data point to the selective transfer of IgG1 and IgG3 antibodies with attenuated Fc-effector functions into the CNS, possibly reflecting exclusion of functional antibodies aimed at protecting the brain from potentially pathological antibodies. Along these lines, complement activation has been implicated in brain tissue damage in some acute infectious encephalitis [30–32], but is not observed in other chronic conditions [33], suggesting that ADCD-inducing antibodies may penetrate the brain only under particular conditions, and are otherwise excluded from the brain antibody arsenal during infections marked by persistent low-level viral replication, such as HIV infection of the CNS.

Figure 3. Antibody signatures in plasma and CSF. A, Violin plots of gp120-specific antibody functions in plasma (P) and CSF (C), including ADCD, ACP, ADNP, and antibody-dependent NK cell activation with expression or production of CD107 (NK CD107), IFN-γ (NK IFN-γ), and CCL4 (NK CCL4). B, Bar plots of mean differences in antibody functions between plasma and CSF, across all 3 HIV antigens (gp120, gp140, p24). Bars on the right (or left) show features higher in plasma (or CSF). Only difference in mean between groups that are statistically significant (P < .05) are plotted. C, Violin plots of Fc receptor binding for gp120-specific antibodies in plasma (P) and CSF (C), including Fcγ receptors (FcγR2A, FcγR2B, FcγR3A, FcγR3B), Fcα receptor (FcαR), and binding to complement component C1q. D, Bar plots of mean differences in antibody binding to Fc receptors between plasma and CSF, across all 3 HIV antigens (gp120, gp140, p24). Bars on the right (or left) show features higher in plasma (or CSF). Only difference in mean between groups that are statistically significant (P < .05) are plotted. A–D, normalized values, Z-scores, Friedman test with Dunn correction, **P < .01, ***P < .001, ****P < .0001. Abbreviations: ADCD, antibody-dependent complement deposition; ACP, antibody-dependent cellular phagocytosis; ADNP, antibody-dependent neutrophil phagocytosis; CSF, cerebrospinal fluid; HIV, human immunodeficiency virus; IFN-γ, interferon-γ; NK, natural killer; ns, not statistically significant.

Defining Multivariate Compartment-Specific Antibody Profiles and Functional Coordination

To define the features that distinguished plasma- and CSF-antibodies, we next applied multivariate models (LASSO/PLSDA) to all combined HIV-specific antibody data [21]. Clear separation was observed between plasma and CSF antibody profiles (Figure 4A; model validation P < .01, accuracy 100%). In agreement with the univariate analysis, gp120-specific ACP, ADCD, and FcRs binding (particularly, FcγR2B and FcRn) were enriched in plasma (Figure 4B). These data clearly demonstrate the exclusion of functional antibodies...
Figure 4. Multivariate antibody signatures and functional correlations across plasma and CSF. A and B, Multivariate analysis of antibody signatures in plasma and CSF. M-PLSDA on LASSO-selected features was used to resolve antibody profiles in plasma:CSF pairs. Dots represent individual samples (plasma [P] CSF [C]) across all 3 HIV antigens (gp120, gp140, p24). Five-fold cross validation was performed, resulting in 100% cross-validation accuracy. Bar graph shows LASSO-selected features across all 3 HIV antigens (gp120, gp140, p24) ranked by their variable importance in projection. All selected features are enriched in plasma. C and D, Correlation matrix across gp120-specific antibody functions (ADCD, ADCP, ADNP, NK CD107, NK IFN-γ, NK CCL4) Ig classes and subclasses (total IgG, IgG1, IgG2, IgG3, IgG4, IgM, IgA1, IgA2, and Fc receptor binding (FcγR2A, FcγR2B, FcγR3A, FcγR3B, FcRn, FcγR, C1q) in plasma and CSF where correlation strength is proportional to color intensity ($r$ from $-1$ = negative correlation, blue to $1$ = positive correlation, purple). Z-scores, Spearman $r$ test, Benjamini-Hochberg correction for multiple comparisons; *$P<.05$, **$P<.01$, ***$P<.001$. Abbreviations: ADCD, antibody-dependent complement deposition; ADCP, antibody-dependent cellular phagocytosis; ADNP, antibody-dependent neutrophil phagocytosis; CSF, cerebrospinal fluid; HIV, human immunodeficiency virus; IFN-γ, interferon-γ; Ig, immunoglobulin; LASSO, least absolute shrinkage selection operator; M-PLSDA, multilevel partial least square discriminant analysis; NK, natural killer.
Figure 5. Effect of ART treatment on antibody profiles in plasma and CSF. A, Violin plots of IgG subclasses and IgM in plasma and CSF, comparing ART-treated and untreated individuals. B, Violin plots of antibody functions in plasma and CSF, comparing ART-treated and untreated individuals. A and B, Normalized values, Z-scores, Friedman test with Dunn correction, \( P > .05 \). C and D, Correlation matrix between clinical variables (age, years of education, duration of HIV infection, plasma HIV load, CSF HIV load, count of CD4+ T cells/mL in blood at the time of sample, nadir of CD4+ T-cell count, total protein mg/dL in CSF, white blood cells/mL in CSF, Global Clinical Rating, Global Cognitive T-scores) and gp120-specific antibody features (Ig classes [IgG, IgM, IgA], Ig subclasses [IgG1, IgG2, IgG3, IgG4, IgA1, IgA2], functions [ADCD, ADCP, ADNP, NK CD107, NK IFN-\( \gamma \), NK CCL4], and Fc receptor binding [Fc\( \gamma \)2A, Fc\( \gamma \)2B, Fc\( \gamma \)3A, Fc\( \gamma \)3B, FcRn, Fc\( \alpha \)R, C1q]) in plasma and CSF of ART-treated versus untreated individuals, where correlation strength is proportional to color intensity: \( r \) from \(-1\) = negative correlation, blue to 1 = positive correlation, purple. Z-scores, Spearman \( r \) test, Benjamini-Hochberg correction for multiple comparisons, \( ^* P < .05, ^{**} P < .01, ^{***} P < .001 \). Abbreviations: ADCD, antibody-dependent complement deposition; ADCP, antibody-dependent cellular phagocytosis; ADNP, antibody-dependent neutrophil phagocytosis; ART, antiretroviral therapy; CD4Absol, count of CD4+ T cells/mL in blood at the time of sample; CD4Nadir, nadir of CD4+ T cells count; CSF, cerebrospinal fluid; CSFprot, total protein mg/dL in CSF; CSFVL, CSF HIV load; CSFWBC, white blood cells/mL in CSF; Educat, years of education; GClinR, Global Clinical Rating; GcognTS, Global Cognitive T-scores; HIV, human immunodeficiency virus; HVIdurat, duration of HIV infection; IFN-\( \gamma \), interferon-\( \gamma \); Ig, immunoglobulin; NK, natural killer; PlasmaVL, plasma HIV load; T, ART-treated; U, untreated.
Figure 6. CSF-antibody signatures in HAND. A, Dot-bar plot showing CSF-HIV load (expressed as number of copies/mL), CSF leukocytes count, blood CD4+ T-cell count (expressed as number of CD4+ cells/mL), and CSF-HIV-specific IgG titers in HAND (green) versus cognitively unimpaired (salmon) individuals. Mann-Whitney test. B and C, Violin plots of FcR binding and functions in CSF, comparing HAND (green) and cognitively unimpaired (salmon) individuals. Z-scores, Friedman test with Dunn correction, $P < .05$. D, Correlation matrix of gp120-specific functions (ADCD, ADCP, ADNP, NK CD107, NK IFN-$\gamma$, NK CCL4 Ig classes and subclasses (total IgG, IgG1, IgG2, IgG3, IgG4, IgM, IgA1, IgA2), Fc receptor binding (Fcγ2AR, Fcγ2BR, Fcγ3AR, Fcγ3BR, FcγR, FcRn, FcR, C1q), and clinical variables (duration of ART treatment, age, years of education, estimated duration of HIV, plasma HIV load, CSF HIV load, CD4+ T-cell count at time of sample, CD4+ T-cell count at nadir, CSF total protein level, CSF white blood cell count/mL, Global Cognitive Rating [higher levels indicate higher cognitive impairment], Global Cognitive T-Scores [higher levels indicate lower cognitive impairment]) in CSF of HAND and CN individuals. Correlation strength is proportional to color intensity: $r$ from $-1$ = negative correlation, blue to 1 = positive correlation, purple). Z-scores, Spearman $r$ test, Benjamini-Hochberg correction for multiple comparisons, $^* P < .05$, $^{**} P < .01$, $^{***} P < .001$. Abbreviations: ADCD, antibody-dependent complement deposition; ADCP, antibody-dependent cellular phagocytosis; ADNP, antibody-dependent neutrophil phagocytosis; ARTdurat, duration of ART treatment; CD4Nadir, CD4+ T-cell count at nadir; CSF, cerebrospinal fluid; CSFprot, CSF total protein level; CSFVL, CSF HIV load; CSFWBC, CSF white blood cell count/mL; CU, cognitively unimpaired; educat, years of education; GCRn, Global Cognitive Rating; GCS, Global Cognitive T-Scores; HAND, HIV-associated neurocognitive disorder; HIV, human immunodeficiency virus; HIVdurat, estimated duration of HIV; IFN-$\gamma$, interferon-$\gamma$; Ig, immunoglobulin; MFI, median fluorescence intensity; NK, natural killer; ns, not statistically significant; PlasmaVL, plasma HIV load, WBC, white blood cell.
from the CSF, far beyond our traditional understanding of isotype/subclass sieving into this unique compartment.

Also, we explored the coordination between HIV-specific antibody functions, isotypes, subclasses, and FcR binding. While antibody responses were highly correlated in plasma, CSF antibodies exhibited lower coordination in all but FcR binding profiles (Figure 4C and 4D). These data further highlight the presence of different qualities of antibodies across compartments, and the sieving of antibodies from the periphery, rather than local production of antibodies in the CNS that would be anticipated to exhibit a broader and more coordinated functional profile.

**ART Treatment Results in a More Strongly Coordinated HIV-Specific Plasma Antibody Response**

Previous studies have shown that ART induces a strongly coordinated humoral immune response, potentially linked to reduced immune activation in the setting of lower viremia [34]. Whether ART also affects humoral responses in the CSF remains unclear. Here, we compared antibody profiles, viral markers, and clinical features in plasma and CSF across the 11 virally suppressed ART-treated and 9 ART-naive individuals (Table 1). As expected, untreated individuals exhibited higher viral loads in both plasma and CSF and lower CD4+ counts in blood. Additionally, untreated individuals had higher leukocyte counts in the CSF, suggesting persistent neuroinflammation. Otherwise, no differences in neurocognitive scores (Table 1), Ig isotype/subclass distribution (Figure 5A), function (Figure 5B), and FcR binding (not shown) were observed between treated and untreated individuals. As previously reported [34, 35], enhanced HIV-specific antibody functional coordination in the plasma was observed in the ART-treated compared to untreated individuals (Figure 5C). In contrast, in the CSF, antibody functional coordination remained low and more variable, regardless of ART treatment (Figure 5D), suggesting functional segregation of antibody effector profiles across the compartments even in the setting of successful ART. Thus, these data suggest that ART-induced reduction in peripheral immune activation may enhance systemic humoral coordination, but may have a more limited effect on CSF antibody profiles.

**HAND Is Associated With Higher Viral Burden, Neuroinflammation, and Antibody Functional Coordination**

We next aimed to define the relationship between viral burden or antibody features and HAND (Supplementary Table). Individuals with HAND exhibited higher CSF HIV loads and leukocyte counts (Figure 6A), suggesting a role for CNS viral burden and neuroinflammation in the development of neurocognitive dysfunction, as supported by previous studies [11, 36, 37], even in some persons with milder forms (ANI and MND). As expected, we observed a relationship between low blood CD4+ counts and presence of HAND (Figure 6A) [38, 39], likely reflecting advanced HIV disease. Additionally, a logistic regression model with HAND as the outcome and ART along with antibody features as predictors suggested ART was not a significant predictor of HAND status ($P = .99$), in accordance with univariate analysis (Supplemental Table). Interestingly, while there were no significant differences in antibody titers, FcR binding, and function between cognitively unimpaired and impaired individuals in plasma and CSF (Figure 6B and 6C, and Supplementary Figure 2), coordination of FcR binding, antibody titers, and functions were higher in the CSF of individuals with HAND compared to cognitively unimpaired individuals (Figure 6D), pointing to an inflammatory shift in antibody profiles within the brain of cognitively impaired individuals. Whether this coordination results in distinct inflammatory cascades within the CNS compartment that contributes to progressive neuroimpairment remains unclear but could provide insights into possible mechanisms to prevent or delay HAND.

**DISCUSSION**

There is increasing evidence that humoral responses, in particular highly functional antibodies, are important correlates of HIV viral control [16, 40]. These responses have been well-established in the peripheral blood. However, whether these antibodies can transit to the CNS or other privileged sites where the virus can escape traditional immune control remains unclear. Thus, to begin to define the nature of antibody transfer to the CNS, we deeply profiled the humoral immune response across plasma and CSF in a cohort of ART-treated and untreated persons with HIV. Striking compartmentalization was observed in antibody profiles, marked by a unique and highly selective enrichment of poorly functional IgG1 and IgG3 antibodies in the CSF. Moreover, HIV-specific antibodies were transferred at higher levels than antibodies to other pathogens. Given that FcRn is unlikely to bind and selectively transport antibodies, these data suggest that an alternative transfer mechanism may contribute to antibody crossing through the BBB that is augmented in the setting of HIV infection. Thus, this study provides previously unappreciated insights into antibody transfer to the brain that may support the design of future therapeutic interventions to cure HIV and other infections in the CNS.

If B cells were producing HIV-specific antibodies locally in the CNS, they would switch and produce multiple antibody isotypes or subclasses. Instead, in our study we observed a striking sieving across the BBB, marked by selection of IgG1 and IgG3 in the CSF for both HIV and non–HIV–specific antibodies. Given enhanced affinity of FcRn for IgGs, with marked transfer efficiency differences across subclasses (IgG1 > IgG3 > IgG2) [28, 29], these data are consistent with an FcRn-mediated
transfer mechanism across the BBB. However, the highly selective exclusion of FcγR-binding and functional antibodies suggests sieving mechanisms beyond FcRn. For instance, FcRn may act with other Fc-receptors at the BBB or other barriers. Along these lines, recent data showed collaboration between FcγR3a and FcRn in the placenta [19, 41], to ensure selection of optimal IgG, or to restrict passage of inflammatory antibodies that could otherwise cause localized pathology. Alternatively, selective exclusion of inflammatory IgGs from the brain may also occur via enzymatic modification of Fc-glycosylation of CSF antibodies, as documented in the liver [42]. However, whether similar mechanisms are at play in the CNS or whether a noncanonical Fc-receptor may work in concert with FcRn to selectively enrich less inflammatory antibodies in the brain is unclear.

During early infection, anti-HIV humoral responses evolve within 2–3 weeks after initial virus entry [43], with antibody levels increasing earlier in the blood than in CSF [10]. During chronic infection, ART treatment results in a significant decrease in HIV loads [44], systemic immune activation [45], and systemic antibody levels [46]. However, whether ART has similar effects in the CSF compartment remains a matter of debate. In our study, we did not observe differences in antibody titers, IgG subclasses, functions, or FcγR binding of CSF-antibodies between treated and untreated individuals. However, while coordination of antibody effector functions improved in the peripheral circulation with ART, the same improvement in antibody function was not observed in the CSF. These data highlight the persistent dysfunction in the CSF, which may relate to reduced ART activity in the CNS. These results are in line with recent observations that in chronically infected patients, ART does not significantly affect CSF antibody levels [10], and highlights that HIV-specific antibody quality may in fact reflect persistent viral load in the CNS.

HAND remains one of the main drivers of poor quality of life in people living with HIV, and its pathogenesis is multifactorial [36]. Our study supports a role of HIV infection of the CNS and neuroinflammation, even in some individuals with only mildly impaired cognition (ANI and MND), by showing a direct correlation between severity of cognitive impairment, level of HIV replication in CSF, and neuroinflammation, indicated by an increased number of infiltrating leukocytes in CSF. This is in line with other studies reporting elevated inflammatory brain metabolites [47, 48] and white matter changes in HIV-infected individuals with more severe forms of HAND [49]. However, our observation that antibody titers and Fc-effector functions did not correlate with severity of HAND suggests that antibody-mediated innate immune functions are not likely to be major contributors to HAND pathogenesis.

The current study has several limitations, including the small cross-sectional sample size, which limited statistical power to perform subgroup analyses, such as the comparison of HAND severity by ART treatment. Additionally, the study lacked information on oligoclonal bands in the CSF to support either the selective antibody-transfer across the BBB or the potential for intrathecal antibody synthesis. Also, we did not perform genetic analyses of HIV in the blood and CSF to support the role of compartmentalized immune responses in the emergence/selection of unique CNS HIV variants that might contribute to HAND pathogenesis [50]. Nevertheless, our study provides the first functional characterization of brain-specific HIV antibody signatures in the CNS, suggesting a unique compartmentalization of the anti-HIV humoral response during chronic infection. The selective sieving of antibodies with reduced functionality might help to prevent pathology within the brain, but may also allow the virus to persist unrestricted by traditional antiviral immune mechanisms. These novel observations highlight the importance of deeper longitudinal analyses of antibody trafficking to the brain in HIV and across neuroinfections and could provide critical insights for the design of next-generation therapeutics that can gain access to the brain through mechanisms that go beyond FcRn-mediated transfer to selectively target and eliminate viruses from this immune-privileged site.

### Supplementary Data

**Supplementary materials** are available at *The Journal of Infectious Diseases* online (http://jid.oxfordjournals.org/). Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyedited. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

### Notes

**Author contributions.** M. S., D. G., and G. A. conceived and designed the experiments. M. S., S. S., and Y. D. performed the experiments. M. S., D. C., C. L., and D. A. L. analyzed the data. M. S., D. C., C. L., M. J. G., E. R., N. W., S. S. M., D. G., and G. A. contributed to interpretation of data. M. S. and G. A. wrote the paper. All authors revised the paper for intellectual content.

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