

Structural and Functional Brain Abnormalities in Human Immunodeficiency Virus Disease Revealed by Multimodal Magnetic Resonance Imaging Fusion: Association With Cognitive Function

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Background. Human immunodeficiency virus (HIV)-associated neurocognitive impairment remains a prevalent comorbidity that impacts daily functioning and increases morbidity. While HIV infection is known to cause widespread disruptions in the brain, different magnetic resonance imaging (MRI) modalities have not been effectively integrated. In this study, we applied 3-way supervised fusion to investigate how structural and functional coalterations affect cognitive function.

Methods. Participants (59 people living with HIV and 58 without HIV) completed comprehensive neuropsychological testing and multimodal MRI scanning to acquire high-resolution anatomical, diffusion-weighted, and resting-state functional images. Preprocessed data were reduced using voxel-based morphometry, probabilistic tractography, and regional homogeneity, respectively. We applied multimodal canonical correlation analysis with reference plus joint independent component analysis using global cognitive functioning as the reference.

Results. Compared with controls, participants living with HIV had lower global cognitive functioning. One joint component was both group discriminating and correlated with cognitive function. This component included the following covarying regions: fractional anisotropy in the corpus callosum, short and long association fiber tracts, and corticopontine fibers; gray matter volume in the thalamus, prefrontal cortex, precuneus, posterior parietal regions, and occipital lobe; and functional connectivity in frontoparietal and visual processing regions. Component loadings for fractional anisotropy also correlated with immunosuppression.

Conclusions. These results suggest that coalterations in brain structure and function can distinguish people with and without HIV and may drive cognitive impairment. As MRI becomes more commonplace in HIV care, multimodal fusion may provide neural biomarkers to support diagnosis and treatment of cognitive impairment.

Keywords. neuroHIV; magnetic resonance imaging; multimodal fusion.

With the advent of combination antiretroviral therapies, people living with human immunodeficiency virus (PLWH) have nearly average life expectancies [1]. Yet, HIV-associated neurocognitive disorder (HAND) remains a prevalent comorbidity, with rates ranging from 15% to 55% across populations [2]. While milder forms of HAND now predominate, these impairments have real-world impacts on daily functioning (eg, employment, medication adherence) and are predictive

of increased morbidity and mortality [3]. Even with sustained viral suppression, HIV reservoirs in the central nervous system may lead to brain injury via chronic inflammation [4].

Noninvasive magnetic resonance imaging (MRI) has yielded important insights into the neuropathology of HIV. Structural MRI studies that quantify gray matter morphology have demonstrated HIV-related atrophy predominantly in the frontal lobes and striatum [5]. Diffusion-weighted imaging (DWI), which maps the diffusion of water molecules across white matter, has revealed decreased integrity of multiple projection, association, and callosal fibers in PLWH compared with controls not living with HIV [6]. Studies of resting-state functional MRI (rs-fMRI), which measures the temporal correlation of spontaneous changes in blood flow across spatially distributed regions, have reported diminished connectivity within and between major neural networks in PLWH [7]. While there is no specific profile of neuroHIV, likely due to multiple mechanisms,

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the growing literature supports the role of brain abnormalities in the development of HAND.

While each MRI modality offers unique and complementary information related to neurologic disease, when evaluated independently, unimodal studies may provide an incomplete characterization of brain structure and function. By contrast, innovative fusion approaches jointly analyze multiple neuroimaging datasets to reveal interrelated patterns across modalities. Specifically, multimodal canonical correlation analysis with reference plus joint independent component analysis (MCCAR + jICA) is a fusion method that flexibly captures multimodal interactions and generates independent spatial components [8]. This analytic framework has been successfully applied to examine cognitive dysfunction related to multiple neuropsychiatric disorders, exhibiting superior estimation accuracy and reliability relative to its alternatives [9, 10].

To date, MRI studies in neuroHIV have not been adequately integrated to yield a comprehensive view of the structural and functional coalterations associated with neurocognitive impairment. Our aim in this study was to identify neural biomarkers acquired using multimodal MRI that are predictive of cognitive function in PLWH. Given the complexity of high-dimensional datasets, our supervised model used cognitive function as a reference to identify linked multimodal components.

METHODS

Sampling

Adults aged 18–55 years were recruited from infectious diseases clinics and the community via advertisements in local newspapers, websites, and nonprofit organizations. For individuals with known HIV diagnosis, HIV status was verified by medical record review. PLWH had to have been diagnosed for >3 months and prescribed antiretroviral medications. For others, an oral rapid antibody test (OraSure ADVANCE HIV-1/2) was conducted; all participants not living with HIV had a nonreactive result. Exclusion criteria were English nonfluency or illiteracy, less than an eighth grade education, severe learning disability, unresolved neurological disorders or neuroinfections, severe head trauma with loss of consciousness for >30 minutes and persistent functional decline, bipolar I or psychotic disorder, acute psychiatric symptoms that interfere with functioning, MRI contraindications, and/or impaired mental status. Current nicotine, alcohol, and marijuana use was permissible, but participants could not meet criteria for alcohol or marijuana dependence. For other drugs, individuals were excluded for a history of dependence, lifetime regular use for >2 years, any use in the past 30 days, and/or a positive urine drug screen. These exclusions are consistent with current guidelines for classifying contributing and confounding conditions to HAND [11].

Procedures

Participants were enrolled in 1 of 3 protocols with shared procedures. The proportion of PLWH was equivalent across protocols

(55%, 47%, and 50%; $\chi(2)^2 = 0.42$, $P = .81$). Participants provided written informed consent, and procedures were approved by the Duke University Health System Institutional Review Board. An in-person screening assessed medical, psychiatric, and substance abuse histories. Eligible participants returned for neuropsychological testing, MRI scanning, and additional assessments. Participants abstained from alcohol, marijuana, and illicit drugs for ≥ 4 hours prior to the visit. To minimize nicotine withdrawal, participants were allowed to smoke up to 30 minutes prior to the MRI.

Assessments

Screening. The Addiction Severity Index-Lite, a structured clinical interview, was used to assess current and lifetime functioning across multiple domains, including substance use, psychiatric symptoms, and medical conditions [12]. The Mini International Neuropsychiatric Interview was used to identify Diagnostic and Statistical Manual of Mental Disorders, fourth edition (DSM-IV) mood and psychotic disorders [13]. Module E of the Structured Clinical Interview for DSM-IV was used to identify substance use disorders [14]. Recent drug use was assessed using a urine toxicology screen for amphetamine, barbiturates, benzodiazepines, cannabis, cocaine, methadone, methamphetamine, opioids, and oxycodone. Medical records were reviewed to ensure no exclusionary conditions.

Neuropsychological Testing. The battery was used to assess 7 domains relevant to HAND, including executive functioning, processing speed, working memory, verbal fluency, learning, motor, and memory (see [Supplementary Table 1](#) for description of tests). Raw scores were converted to standardized T scores (mean = 50, standard deviation = 10) that correct for age and, when available, other demographic factors. T scores for each test within a domain were averaged to create a domain T score, and domain T scores were averaged to create a global T score.

HIV Clinical Data. The following data were abstracted from the medical record: date of HIV diagnosis, nadir and current CD4⁺ T-cell count, current antiretroviral regimen, and most recent HIV viral load. All participants had laboratory testing within 8 months of their MRI (M = 2.03 months, SD = 1.71). Viral suppression was defined as <50 copies/mL. To normalize the data for analysis, CD4⁺ T-cell counts were square-root transformed.

MRI Data Acquisition

Data were acquired with a single 3.0T GE Discovery MR750 whole-body scanner using an 8-channel head coil. High-resolution T1-weighted (T1w) structural images were acquired with the following parameters: repetition time (TR) = 8.10 ms, TE = 3.18 ms, FOV = 25.6 cm, 256*256 matrix, 12° flip, 166 interleaved slices of 1 mm thickness). DWI was acquired in the axial plane using single-shot spin-echo

echo-planar imaging (FOV = 25.6 cm, 128*128 matrix, 90° flip, 2-mm interleaved slices). Additional parameters differed slightly for protocol 1 (b-factor = 900 s/mm², TR/TE = 10 000/83.2 ms, 73 slices), protocol 2 (b-factor = 800 s/mm², TR/TE = 8000/77.9 ms, 67 slices), and protocol 3 (b-factor = 800 s/mm², TR/TE = 8000/78.2 ms, 67 slices). Data were acquired in 30 directions for the first 2 protocols and 64 directions for the third protocol. The 64-direction protocol was downsampled using a MATLAB dot() function (inner product) that identified the diffusion-encoding direction most similar to those in the 30-direction protocol. For the rs-fMRI data, whole-brain BOLD images were collected using T2*-weighted echo-planar imaging (TR = 2000 ms, FOV = 24 cm, 64*64 matrix, 3.8 mm interleaved slices, voxel size 3.75*3.75 mm*3.8 mm). Additional parameters differed slightly between the first protocol (TE = 27 ms, 77° flip, 39 slices) and the other 2 protocols (TE = 25 ms, 90° flip, 35 slices). Data harmonization consisted of ensuring that all scans had 148 volumes.

Image Preprocessing

The DWI data was denoised [15], motion- and eddy-corrected using DTIPrep 3.1 [16], and preprocessed using tools from FSL 5.0.9 (B1 bias-correction, global DWI intensity normalization, rigid-body registration between participants' mean B0 images and T1w data, and nonlinear registration between T1w data and the standard Montreal Neurologic Institute [MNI brain]) [17]. The probabilistic tractography was conducted in MRtrix 3.0 using an anatomically constrained procedure and a diffusion tensor. One million tracks were generated by seeds from the mask image in 0.2 mm steps [18, 19]. Fractional anisotropy (FA) was then calculated within all white matter voxels.

The T1w images were skull-stripped using custom thresholds and analyzed with FSL-VBM [20]. Participant-level maps of gray matter volume (GMV) were created.

For rs-fMRI, the first 6 volumes were excluded to ensure steady-state sampling. Data processing was conducted using FSL, including motion correction using rigid-body transformation, slice-timing correction, high-pass temporal filtering using a Gaussian filter at 0.01 Hz, and signal intensity normalization. White matter and cerebrospinal fluid means were regressed out. Motion and physiological-related components were removed using independent component analysis-automatic removal of motion artifacts (ICA-AROMA) [21]. Images were registered to the 2-mm MNI template using nonlinear registration [22]. Registration of functional data to the T1w acquisition was done using a 12-parameter affine transformation. After preprocessing and warping to MNI space, regional homogeneity (ReHo) was calculated using the AFNI v20.2.16 tool 3dReHo using a 27-voxel neighborhood [23]. ReHo evaluates the similarity between the time series of a voxel to a predefined cluster of its nearest neighbors [24].

Statistical Analyses

Feature Normalization. Images were upsampled to 3 mm³ and spatially smoothed using a Gaussian kernel with a full width at half maximum of 6 mm³. The 3-dimensional images were reshaped into a 1-dimensional vector and stacked, forming a matrix ($N_{\text{participant}} \times N_{\text{voxel}}$) for each modality. The matrices were normalized to have the same average sum of squares. A single normalization factor was used for each data type; thus, the relative scaling within a given data type was preserved, but the units between data types were the same (in a least-squares sense). Age, intracranial volume, nicotine use, and protocol were regressed out from the ReHo, GMV, and FA features.

Multimodal Fusion. The preprocessed MRI features were fed into the MCCAR + jICA pipeline with Global T score as the reference (Figure 1). After MCCAR optimization, we obtained the canonical variants that were most correlated with the reference in each modality and across participants between modalities. Joint ICA was applied to the concatenated maps to keep the modality linkage while maximizing the spatial independence of components. Based on the minimum description length criterion [25], 23 independent components (ICs) were estimated with corresponding participant-wise loadings derived for each modality. ICs from the same index across all 3 modalities are considered joint ICs. Our analysis selected the joint IC, denoted IC_{ref}, that correlated maximally with the reference across all modalities [8]. Multivariate analysis of variance was used to compare the groups on the IC_{ref} loadings. The identified brain regions are described using the Harvard-Oxford Atlas for gray matter regions [26] and the IIT Human Brain Atlas for white matter tracts [27]. Pearson correlation was used to examine the relationship between IC_{ref} loadings and CD4⁺ T-cell counts, and independent sample *t* tests were used to compare loadings by HIV suppression.

RESULTS

Participant Characteristics

Table 1 includes the demographic characteristics of the 59 PLWH and 58 controls not living with HIV. The majority of PLWH were on a regimen of dual nucleoside reverse transcriptase inhibitor plus a nonnucleoside reverse transcriptase inhibitor (42.4%), an integrase strand transfer inhibitor (33.9%), or a protease inhibitor (15.3%); 8.5% were on a different combination. Duration since HIV diagnosis ranged from 4 months to 29 years (Median = 7.17, interquartile range [IQR] = 11.70). The median nadir CD4⁺ T-cell count was 250 (IQR = 297), and 41% had a nadir <200. Currently, the majority (80%) had a suppressed plasma HIV RNA, and the median CD4⁺ T-cell count was 598 (IQR = 424). Participants with unsuppressed viral load were significantly more likely to have a current CD4⁺ T-cell count <200 (67% vs 0%; $\chi^2(1) = 36.25$, $P < .001$). PLWH were

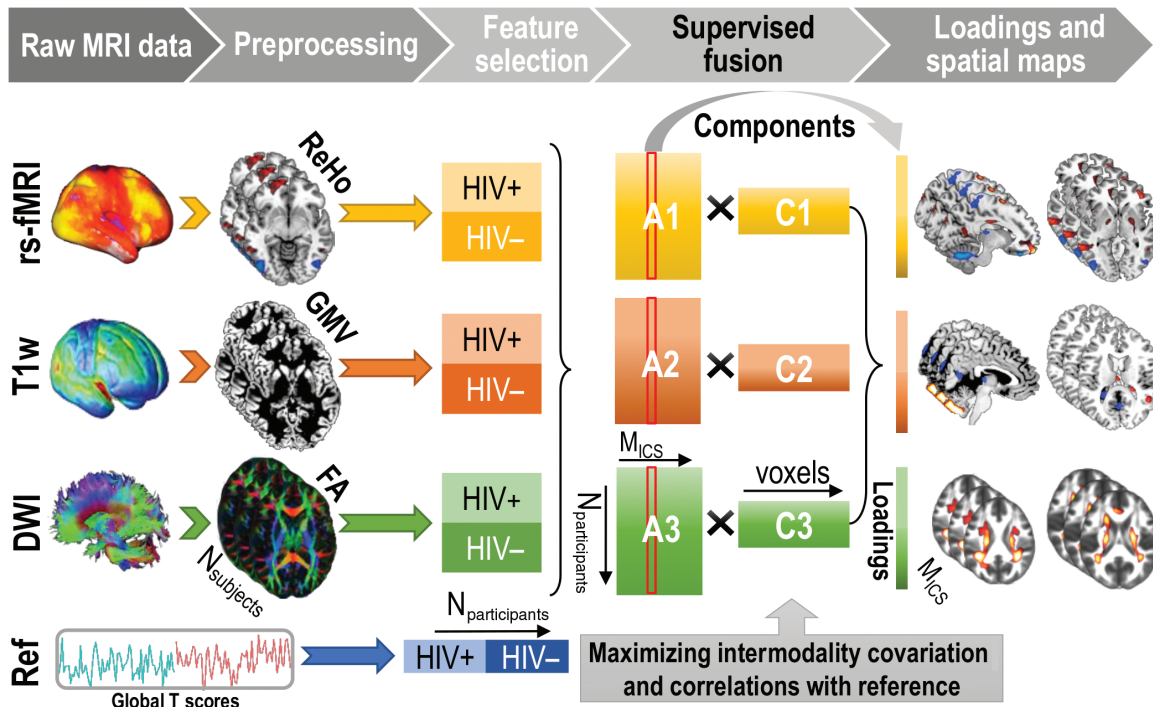


Figure 1. Flow chart of the multimodal canonical correlation analysis with reference plus joint independent component analysis fusion pipeline. The preprocessed MRI features (ie, ReHo from resting-state fMRI, GMV from T1-weighted, and FA from DWI) are entered with global cognitive scores as the reference. After completing the cognition-guided fusion, component loadings and spatial maps for each modality are generated. Abbreviations: DWI, diffusion-weighted imaging; FA, fractional anisotropy; GMV, gray matter volume; HIV, human immunodeficiency virus; M_{ics} , M independent components; MRI, magnetic resonance imaging; Ref, reference; rs-fMRI, resting-state functional magnetic resonance imaging; ReHo, regional homogeneity.

older and more likely to smoke cigarettes daily but they were otherwise comparable to the group not living with HIV. As expected, cognitive functioning was significantly lower in PLWH compared with those without HIV.

Joint Independent Component

Figure 2 displays representative spatial maps for IC_{ref} , the joint component that maximally correlated with global T score across all 3 modalities. This component was characterized by the following linked regions: FA in bilateral corpus callosum (splenium, body, and genu), short and long association fiber tracts,

and corticopontine fibers; GMV in bilateral visual cortex, medial orbitofrontal cortex, posterior parietal cortex, and thalamus; and ReHo in bilateral posterior parietal cortex; dorsomedial and ventromedial prefrontal cortex; precuneus, precentral, and postcentral gyri; and visual cortex. Supplementary Table 2 details the regions comprising the component. As shown in Figure 2, IC_{ref} correlated significantly with global T score for FA, GMV, and ReHo.

There was a significant group difference for IC_{ref} ($F[3, 113] = 2.985, P = .034$; Wilk's $\Lambda = .927$). PLWH had lower loadings compared with controls not living with HIV for FA

Table 1. Sample Characteristics by Human Immunodeficiency Virus Status

	Living With HIV	Not Living With HIV	Statistic	PValue
Characteristic	N = 59	N = 58		
Age, mean (SD), y	41.24 (8.54)	37.84 (9.47)	$t(115) = 2.04$.044
Female gender, n (%)	14 (23.73)	20 (34.48)	$\chi^2(1) = 1.64$.200
African-American race, n (%)	45 (76.27)	38 (65.52)	$\chi^2(1) = 1.64$.200
Education, mean (SD), y	14.08 (2.08)	14.72 (2.14)	$t(115) = 1.64$.104
Nicotine use in past 30 days, n (%)	22 (37.29)	10 (17.24)	$\chi^2(1) = 5.92$.015
Alcohol use in past 30 days, n (%)	34 (57.63)	34 (58.62)	$\chi^2(1) = 0.01$.913
Marijuana use in past 30 days, n (%)	12 (20.34)	11 (18.97)	$\chi^2(1) = 0.04$.852
Global cognitive function, mean (SD)	45.31 (6.47)	49.15 (6.52)	$t(115) = 3.20$.002

Abbreviations: HIV, human immunodeficiency virus; SD, standard deviation.

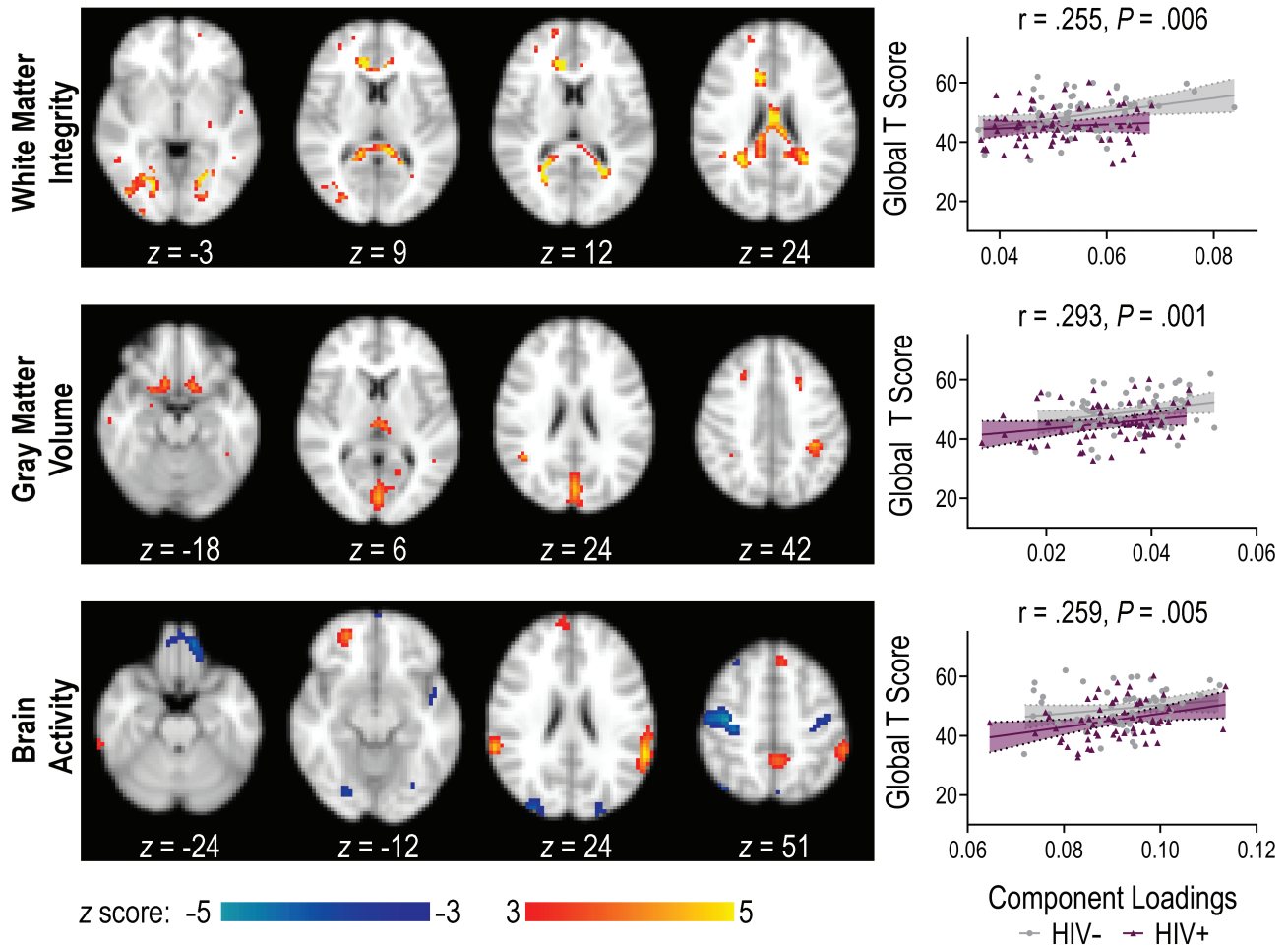


Figure 2. Illustration of the joint independent component for each magnetic resonance imaging modality. The brain maps show the regions that comprise the multimodal component with brain maps visualized at $|z| > 3.0$. The regional homogeneity map includes regions with positive activation (lighter shading) and negative activation, or de-activation (darker shading). The scatter plots to the right illustrate the correlation of the component loadings with cognitive performance (global T score). Abbreviation: HIV, human immunodeficiency virus.

and GMV, meaning the component was expressed less strongly (Table 2).

Association With HIV Clinical Measures

For IC_{ref} markers of immunosuppression correlated with the component loading for FA but not GMV or ReHo. Specifically, lower FA loadings correlated with lower current ($r = .272, P = .037$) and nadir ($r = .338, P = .009$) $CD4^+$ T-cell

counts. In contrast, HIV viral suppression was unrelated to IC_{ref} loadings.

In a sensitivity analysis that excluded participants with unsuppressed HIV, there were similar group differences in IC_{ref} loadings for FA ($F[1103] = 4.922, P = .029$) and GMV ($F[1103] = 5.606, P = .020$). However, the correlation between FA loadings and $CD4^+$ T-cell counts were no longer significant for current ($r = .124, P = .406$) or nadir ($r = .268, P = .068$) with the restricted range.

Table 2. Group Comparison on Component Loading for IC_{23}

Modality	Living With HIV	Not Living With HIV	Statistic	P Value
	N = 59	N = 58		
Gray matter volume, mean (SD)	0.032 (0.0094)	0.036 (0.0083)	$F(1115) = 6.357$.013
White matter integrity (fractional anisotropy), mean (SD)	0.051 (0.0090)	0.056 (0.0087)	$F(1115) = 7.359$.008
Brain activity (regional homogeneity), mean (SD)	0.091 (0.0088)	0.092 (0.0094)	$F(1115) = .572$.451

Abbreviations: HIV, human immunodeficiency virus; IC, independent component; SD, standard deviation.

DISCUSSION

By applying an innovative analytic approach that “fuses” multimodal MRI data, we identified coalterations in brain structure and function that correlated with cognitive function. Specifically, global T score was linked to reduced volume in the thalamus and visual, posterior parietal, and orbitofrontal cortices; reduced white matter integrity throughout the corpus callosum and association fibers; and altered activity in frontal-parietal and occipital networks.

PLWH had lower component scores for both FA and GMV, suggesting that HIV-associated alterations in brain structure drive neurocognitive impairment. A unique feature of multimodal fusion is its ability to discover linked alterations in spatially distinct brain regions, even when there is no direct morphologic connection. Building on a literature dominated by unimodal analyses, our results support the role of diffuse coalterations in brain structure and function in neuroHIV [5–7].

Supporting a linkage between white and gray matter structure, we found HIV-associated reductions in thalamic volume and integrity of thalamic projection fibers correlated with worse cognitive function. The thalamus is a major “hub” region that acts as a relay station between subcortical areas and the cerebral cortex [28]. Prior DWI studies in PLWH have documented microstructural abnormalities in the white matter fibers of the thalamus [29–31], while structural MRI studies have identified HIV-related atrophy in adjacent gray matter [5].

Effective neural communication may be constrained by the structural composition of the brain. The ReHo component linked to cognitive function displayed activations in posterior parietal and prefrontal cortices, regions typically associated with executive function [32], with corresponding deactivations in the precuneus and medial prefrontal cortex. Such divergent activity in neural function is characteristic of healthy cognition [33]. This pattern of local functional connectivity was linked to reduced integrity in the corpus callosum, a white matter fiber that forms interhemispheric connections, and reduced volume in major hubs, including the thalamus and precuneus. In addition, we found both functional and structural abnormalities in visual processing regions that were negatively correlated with global cognitive function. While there were no group differences for ReHo, the overall component loading may have masked abnormalities in the neural activation patterns linked to cognitive function, as HIV has been associated with both hypoactivation in hippocampal and visual processing regions and compensatory hyperactivation in frontostriatal regions to preserve cognitive function [7]. Future studies might consider the potentially divergent implications of positive versus negative functional homogeneity in relation to HIV-associated neurocognitive impairments.

Clinically, our results suggest that HIV-related immunosuppression may increase vulnerability to white matter degradation, consistent with unimodal studies [5, 34]. The FA component

was correlated with both nadir and current CD4⁺ T-cell counts, such that participants with greater immunosuppression had reduced white matter integrity. While the so-called legacy effect, which links historical severe immunosuppression with brain structure and cognitive function, has been widely reported [2, 3, 7], our results underscore the neurologic benefit of restoring and maintaining strong immune function. Unsuppressed viral load was unrelated to the joint component, and differences in brain structure linked to cognitive function were observed even when people with unsuppressed HIV were excluded. Prior studies have consistently found that viral load is a poor predictor of brain integrity, which may result from cumulative inflammatory processes [35]. However, we were unable to tease out the extent to which immunosuppression during periods of untreated HIV may be driving our findings. In contrast, GMV and ReHo metrics in the identified multimodal component were unrelated to HIV disease characteristics, suggesting that degradations in white matter may precede alterations in gray matter regions, possibly serving as an early marker of HAND [36].

Despite the innovative analytic approach and robust findings, the following limitations should be mentioned. First, the reference-guided fusion strategy uses selected MRI features rather than original data (eg, using ReHo instead of 4D fMRI data). Since the time series of the fMRI data were not used, there may have been some loss of the temporal information. However, compared with the high-dimensional raw data, features tend to be more tractable and provide a more concise signature to link [10, 37]. Future investigations may incorporate alternative features such as mean diffusivity, cortical thickness, and task-evoked neural activation. Another potential limitation is the cross-sectional design. Longitudinal analyses are needed to verify the temporal relationship between structural and functional changes in the brain and their causal relationship to cognitive impairment in larger samples of PLWH with more generalizable eligibility criteria. Finally, the MCCAR + jICA framework was designed to identify brain regions that maximally covary across modalities and that strongly correlate with the reference. There may be additional HIV-related alterations in 1 or more MRI modalities that were not revealed by this approach because they are unrelated to cognitive function or are not linked across modalities.

In summary, our innovative supervised data fusion approach revealed multimodal neural signatures of HIV that correlated with cognitive function. To the best of our knowledge, this is the first attempt to holistically investigate the impact of brain function and structure on HIV-associated neurocognitive impairment. Given that cognitive decline in PLWH is associated with increased morbidity and mortality, there is a critical need for accurate biomarkers of HAND to aid diagnosis, treatment planning, and long-term monitoring. The current standard for HAND diagnosis involves lengthy neuropsychological testing

that can lack precision and sensitivity. Multimodal fusion, which reveals covariation across imaging and clinical modalities, holds great promise as a more reliable and cost-effective biomarker of HAND.

Supplementary Data

Supplementary materials are available at *Clinical Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes

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