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Cognitively unimpaired HIV-positive subjects do not have increased $^{11}$C-PiB
A case-control study

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ABSTRACT

Objectives: Diagnostic challenges exist for differentiating HIV dementia from Alzheimer disease (AD) in older HIV-infected (HIV+) individuals. Similar abnormalities in brain amyloid-$^{11}$B (A$^{42}$) metabolism may be involved in HIV-associated neuropathology and AD. We evaluated the amyloid-binding agent $^{11}$C-Pittsburgh compound B ($^{11}$C-PiB), a biomarker for A$^{42}$ deposition, in cognitively unimpaired HIV+ ($n=10$) participants and matched community controls without dementia ($n=20$).

Methods: In this case-control study, all participants had an $^{11}$C-PiB scan within 2 years of concomitant CSF studies and neuropsychometric testing. Statistical differences between HIV+ and community controls for demographic and clinical values were assessed by $\chi^2$ tests. Participants were further divided into either low (<500 pg/mL) or normal (≥500 pg/mL) CSF A$^{42}$ groups with Student t tests performed to determine if regional differences in fibrillar amyloid plaque deposition varied with CSF A$^{42}$.

Results: Regardless of CSF A$^{42}$ level, none of the HIV+ participants had fibrillar amyloid plaques as assessed by increased $^{11}$C-PiB mean cortical binding potential (MCBP) or binding potential within 4 cortical regions. In contrast, some community controls with low CSF A$^{42}$ (<500 pg/mL) had high $^{11}$C-PiB MCBP with elevated binding potentials (>0.18 arbitrary units) within cortical regions.

Conclusions: Cognitively unimpaired HIV+ participants, even with low CSF A$^{42}$ (<500 pg/mL), do not have $^{11}$C-PiB parameters suggesting brain fibrillar amyloid deposition. The dissimilarity between unimpaired HIV+ and preclinical AD may reflect differences in A$^{42}$ production and/or formation of diffuse plaques. Future longitudinal studies of HIV+ participants with low CSF A$^{42}$ and normal $^{11}$C-PiB are required. Neurology$^®$ 2010;75:111–115

GLOSSARY

A$^{42}$ = amyloid-$^{11}$B; AD = Alzheimer disease; ART = antiretroviral therapy; CDR = Clinical Dementia Rating; CHARTER = CNS Highly Activated Retroviral Therapy Effects Research; GDS = global deficit score; HAND = HIV-associated neurocognitive disorder; LP = lumbar puncture; MCBP = mean cortical binding potential; PiB = Pittsburgh compound B; ROI = region of interest; WUSTL = Washington University in St. Louis.

HIV-associated neuroinflammation can occur despite virologic control with antiretroviral therapy (ART). The prevalence of HIV-infected (HIV+) participants >50 years old has risen as life expectancy increases with ART. If current trends continue, more than 50% of all HIV+ individuals will be >50 years old by 2015. Age is a risk factor for HIV-associated neurocognitive disorder (HAND) and Alzheimer disease (AD). As HIV+ participants age, clinicians face the challenge of differentiating individuals at risk for HAND from those with AD.

Genetic, biochemical, and animal models and autopsy studies have demonstrated a critical role for brain amyloid-$^{11}$B (A$^{42}$) aggregation in AD. Similar neuropathologic abnormalities occur...
with HIV. Postmortem HIV+ subjects have increased brain Aβ42 and tau deposition compared to age-matched community controls. Decreased CSF Aβ42 is observed in subjects with AD and some unimpaired community controls with fibrillar Aβ42. Subjects with HAND have CSF Aβ42 levels similar to participants with mild AD.

Reduced CSF Aβ42 (<500 pg/mL) correlates with increased fibrillar amyloid deposition using the PET amyloid binding agent N-methyl-[11C]2-(4-methylaminophenyl)-6-hydroxybenzothiazole (11C-PiB) in subjects with AD and unimpaired community controls with preclinical AD. It remains unknown if a similar relationship exists for HIV. We investigated if low CSF Aβ42 levels were predictive of increased 11C-PiB binding potentials in cognitively unimpaired HIV+ participants.

METHODS Participants. HIV+ participants (n = 10) (39–59 years of age) with confirmed serologic status were selected from the CNS Highly Activated Retroviral Therapy Effects Research (CHARTER) cohort at Washington University in St. Louis (WUSTL). Four participants with low CSF Aβ42 levels (<500 pg/mL) and 6 with normal CSF Aβ42 levels (≥500 pg/mL) were contacted. We selected community controls (n = 20) (44–63 years of age) of similar sex and education from memory and aging studies at the WUSTL Alzheimer’s Disease Research Center (2 controls for every HIV+ subject). We received approval from the WUSTL ethical standards committee on human experimentation for experiments using human subjects. In this case-control study, written informed consent was obtained from all subjects participating in this study. The recommendations of the Strengthening the Reporting of Observational Studies in Epidemiology criteria were followed whenever applicable.

All participants had an 11C-PiB scan within 2 years of concomitant lumbar puncture (LP) and neuropsychometric testing. For HIV+ participants, cognition was assessed at the time of scan and approximately 2 years prior to LP. Cognition was evaluated in HIV+ subjects using the previously validated global deficit score (GDS) with impairment deemed significant if GDS ≥0.5. For community controls, impairment was assessed by the Clinical Dementia Rating (CDR) scale with impairment noted if CDR >0.

CSF evaluation. CSF collection used previously described methods. CSF Aβ42 was analyzed using a commercial enzyme-linked immunosorbent assay (Innogenetics, Ghent, Belgium). Samples were kept on ice with assays performed on aliquots after a single thaw.

Imaging. Participants underwent 11C-PiB as previously described. Tracer was injected into the antecubital vein with a 60-minute 3-dimensional dynamic PET scan performed. Each subject had a T1-weighted anatomic scan with 11C-PiB images corrected for head motion and registered to this scan. The cerebellum was used as a reference as amyloid deposition has not been observed within this area in community controls. Logan graphical analyses were performed and 11C-PiB distribution volume calculated for the prefrontal, lateral temporal, precuneus, and gyrus rectus. 11C-PiB binding potentials for each region of interest (ROI) and the mean cortical binding potential (MCBP) were calculated.

Statistical analysis. Statistical differences between HIV+ and community controls for demographic and clinical values were assessed by χ² tests. Participants were divided into either low (<500 pg/mL) or normal (≥500 pg/mL) CSF Aβ42 groups using previously defined criterion with excellent sensitivity (100%) and good specificity (84%) for predicting subjects at risk for dementia. An analysis of variance with Bonferroni correction for multiple comparisons assessed if regional differences in fibrillar amyloid plaque deposition varied with CSF Aβ42.

RESULTS Demographic and clinical variables were similar (table). Neither group had significant cognitive impairment. HIV+ participants, even those with low CSF Aβ42 (<500 pg/mL), did not have increased fibrillar amyloid plaques using 11C-PiB (figure 1A). In contrast, community controls with low CSF Aβ42 had more fibrillar amyloid plaques (figure 1B). Several community controls had 11C-PiB measures indistinguishable from a typical AD pattern. These unimpaired community controls may have preclinical AD.

We assessed the relationship between fibrillar amyloid deposition using 11C-PiB and CSF Aβ42 for HIV+ participants and community controls. A 2 × 2 matrix was created using CSF (Aβ42 <500 pg/mL) and 11C-PiB MCBP (<0.18 arbitrary units) (figure 2A). All HIV+ participants were located in the left upper and lower quadrants. Community controls fell within 3 boxes: left upper and lower quadrants and right lower quadrant. Half of the community controls had low CSF Aβ42 and high

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Table: Clinical and laboratory values for HIV-infected (HIV+) and community control participants

<table>
<thead>
<tr>
<th></th>
<th>HIV+ participants (n = 10)</th>
<th>Community control participants (n = 20)</th>
<th>p Value</th>
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<tbody>
<tr>
<td><strong>Demographics</strong></td>
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<tr>
<td>Age, y, mean ± SE</td>
<td>52 ± 6</td>
<td>47 ± 6</td>
<td>0.06</td>
</tr>
<tr>
<td>% Men</td>
<td>45</td>
<td>80</td>
<td>0.07</td>
</tr>
<tr>
<td>Education, y, mean ± SE</td>
<td>16 ± 3</td>
<td>15 ± 2</td>
<td>0.25</td>
</tr>
<tr>
<td>% Taking combination antiretroviral therapy</td>
<td>90</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td><strong>Laboratory studies</strong></td>
<td></td>
<td></td>
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<tr>
<td>CD4, cells/mm³, mean (quartiles)</td>
<td>456 (308, 540)</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Nadir CD4, cells/mm³, mean (quartiles)</td>
<td>152 (20, 200)</td>
<td>NA</td>
<td></td>
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<tr>
<td>Log viral load, copies/mm³, mean (quartiles)</td>
<td>2.47 (1.69, 2.64)</td>
<td>NA</td>
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</tr>
<tr>
<td>CSF Aβ42, mean ± SE</td>
<td>696 ± 333</td>
<td>595 ± 226</td>
<td>0.40</td>
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</table>
A mismatch existed between CSF Aβ42 and 11C-PiB MCPB for HIV+ participants and community controls (left lower quadrant). Binding potentials were assessed within ROIs to determine degree of variation in fibrillar amyloid deposition. Binding potentials were elevated for community controls with low CSF Aβ42 compared to other groups within all areas. HIV+ participants including those with low CSF Aβ42 (n = 4) had binding potentials similar to community controls with normal CSF Aβ42 (figure 2B).

DISCUSSION We observed that cognitively unimpaired HIV+ participants, even with low CSF Aβ42 (<500 pg/mL), did not have increased 11C-PiB that might indicate fibrillar brain amyloid deposition. However, community controls with a low CSF Aβ42 were more likely to have elevated 11C-PiB MCPB (>0.18 arbitrary units). Unimpaired community controls with increased 11C-PiB MCPB may have preclinical AD. Within a 2-year retrospective interval during which we followed the HIV+ participants, even those with low CSF Aβ42 had no significant changes in cognition (GDS = 0.18 at LP and GDS = 0.31 at subsequent 11C-PiB). Our findings suggest that 11C-PiB MCPB differs in cognitively unimpaired HIV+ individuals compared to community controls with low CSF Aβ42. In the setting of HIV, low CSF Aβ42 may not reliably predict fibrillar Aβ brain deposits as it does in preclinical AD. As the HIV+ population ages, this distinction could be...
diagnostically important. It remains necessary to understand whether fibrillar Aβ seen with increased $^{11}$C-PiB is present in patients with HAND. This would assist in differentiating HAND from AD. While APOE status was not determined for participants, future studies investigating the impact of genetic risk factors on $^{11}$C-PiB MCBP and CSF $\alpha$42 in HIV+ participants are required.2

Both $^{11}$C-PiB and CSF $\alpha$42 levels are biomarkers of brain amyloid deposition in patients with AD and antecedent measures of impairment in community controls with preclinical AD.8 A strong inverse correlation exists between these biomarkers. The lack of correlation between CSF $\alpha$42 and $^{11}$C-PiB MCBP in unimpaired HIV+ participants could result from decreased Aβ42 production, increased intraneuronal Aβ42 deposition leading to reduced extracellular concentrations, or more extracellular Aβ42 amyloid but in a diffuse, non-fibrillar Aβ form.4,9,10 In each instance, relatively normal $^{11}$C-PiB would occur. Future longitudinal examination, especially a larger sample of HIV+ participants with low CSF Aβ42 and normal $^{11}$C-PiB, are required to understand whether observed low CSF Aβ42 represents an aggregation of diffuse oligomeric forms ($^{11}$C-PiB–negative) that eventually become substantial fibrillar ($^{11}$C-PiB–positive) deposits,1,5 or simply the low normal end of CSF Aβ42 in HIV+ participants.9 Our findings reinforce the importance of understanding amyloid metabolism in HIV-associated neuropathology, while confirming that low CSF Aβ42 is not simply a manifestation of early fibrillar Aβ deposition in the brain.

AUTHOR CONTRIBUTIONS
Statistical analysis was conducted by Dr. Beau Ances and Dr. Chengjie Xiong.
ACKNOWLEDGMENT

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DISCLOSURE

Dr. Ances receives research support from the NIH (NIMH K23MH801176 [PI]), the Foundation for AIDS Research, and from the Dana Foundation. Dr. Teshome reports no disclosures. Mr. Christensen serves as Staff Scientist from the Washington University School of Medicine; and receives research support from the NIH (1R01DC009950 [Staff Scientist], 5P50NS0683341 [Staff Scientist], 5P50NS0683340 [Staff Scientist], 5P01AG02627604 [Staff Scientist], 5P00AG00568152 [Staff Scientist]) and SU1AG0324802 (Staff Scientist). Ms. Taylor reports no disclosures. Dr. Xiong serves as an Associate Editor of Biostatistics; and receives research support from the NIH (NIA K25 AG025189 [PI], NIA P01 AG26276-01 [Biostatistics Component Leader], NIA 5 P01 AG03991 [Biostatistics Core Director], NIA 5P50 AG05681 [Biostatistics Core Director], NIA U01 AG1024344 [Biostatistics Core Director], and R01 AG029672 [Subcontract PI]), and from the Alzheimer Association. Ms. Aldea reports no disclosures. Dr. Fagan serves on a speakers’ bureau for the Alzheimer’s Association. Dr. Holtzman serves on scientific advisory boards for Sanofi Pharmaceuticals and EnVivo Pharmaceuticals; serves as an Associate Editor of Annals of Neurology, the Journal of Neuroscience, Neurobiology of Disease, and Experimental Neurology; may accrue revenue on pending US Patent 20080145941 (filed 6/18/08); Methods for Measuring the Metabolism of Neurotrophin Derived Biomolecules in Vivo, pending US Patent 20090074775 (filed 3/19/09): Use of Anti-AB Antibody to Treat Traumatic Brain Injury, pending US Patent 20090032598 (filed 2/5/09); Methods to Treat Alzheimer’s Disease or Other Amyloid Beta Accumulation Associated disorders; US Patent 7,159,761 (issued 3/27/07); Humanized antibodies that sequester abeta peptide, US Patent 7,015,044 (issued 3/21/06); Diagnostic for early stage Alzheimer’s disease, US Patent 6,465,195 (issued 10/15/02); Predictive diagnostic for Alzheimer’s disease; serves as a consultant to Merck Serono, Eli Lilly and Company, Takeda Pharmaceutical Company Limited, Abbott, Comerit, Inc., Eisai Inc., and AstraZeneca; is cofounder of and receives board of directors compensation from C2N Diagnostics LLC; receives research support from AstraZeneca, Pfizer Inc., Eli Lilly and Company, Elan Corporation, Forest Laboratories, Inc., the NIH (NIA R53 AG19596 [PI], NINDS 1R01NS057195 [PI], NINDS P01-NS535902 [PI of project 8], NINDS P01-NS532636 [PI of project 3], NIA P01-AG026276 [Co-I], NIA R01-AG025824 [J], NINDS R01-NS034467 [J], NIA U01-AG1024348 [Co-I], NIA P01-AG03991 [PI of project 2]), Care Alzheimer’s Fund, and Fidely Foundation; has received compensation from Washington University from license revenue received for licensing of patent application entitled “Methods for Measuring the Metabolism of Neurotrophin Derived Biomolecules in Vivo” to C2N Diagnostics LLC; and may receive future royalty payments for Washington University licensing patent entitled “Methods for Measuring the Metabolism of Neurotrophin Derived Biomolecules in Vivo” to C2N Diagnostics LLC, and could receive future royalty payments from Washington University for licensing patent entitled “Humanized antibodies that sequester abeta peptide” US Patent 7,159,761 to Eli Lilly and Company. Dr. Morris serves on scientific advisory boards for AstraZeneca, Bristol-Myers Squibb, Genentech, Inc., Merck Serono, Novartis, Pfizer Inc., Schering-Plough Corp., Eli Lilly and Company, Wyeth, and Elan Corporation; serves on the editorial advisory board of Alzheimer's Disease and Associated Disorders; receives royalties from publishing Mild Cognitive Impairment and Early Alzheimer's Disease (John Wiley and Sons, 2008), Demen-

tia (Clinical Publishing, 2007), Handbook of Dementing Illnesses, 2nd edition (Taylor & Francis, 2006), and for an editorial in Lancet Neurology (Elsevier, 2008); and receives research support from Elan Corporation, Wyeth, Eli Lilly and Company, Novartis, Pfizer Inc, Avid Radiopharmaceuticals, the NIH (NIA P50AG05681 [PI], P01AG03991 [PI], P01AG026276 [PI], U01AG032438 [PI], U10AAG024904 [Neuropathology Core Leader], R01AG16335 [Consultant], and P50NS068333 [Investigator]), and from the Dana Foundation. Dr. Mintun serves as a consultant for Avid Radiopharmaceuticals, Inc. and receives research support from the NIH (R01AG103664-05 [PI], R30 AG019153 [Co-PI], 1R01AG03991-26 [Director of Imaging Core], P01 AG026276 [Co-I], P50 AG056801-22 [Project of Project 3], U50 AG032438-02 [Director, Imaging Core], P01 NS078683 [Co-PI], R01 DC009095-03 [Co-I], P01 CA091842 [Co-I], U1L RR024992 [Director, Imaging Unit], R01NS059566-03 [Co-I], and U54CA136598-02 [Director of the Human Imaging Core]). Dr. Clifford serves has served on scientific advisory boards for Biogen Idec, Elan Corporation, Roche, Forest Laboratories, Inc., Genentech, Inc., GlaxoSmithKline, Millennium Pharmaceuticals, Inc., Schering-Plough Corp., Bristol-Meyers Squibb, and Genzyme Corporation; received speaker honorarium and fund-
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REFERENCES


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