Vascular Health is Associated with Amyloid-β in Cognitively Normal Older Adults

Yumei Liua, Sophy J. Perdomoa, Jaimie Warda, Eric D. Vidonib, Jason F. Sisantea,∗, Kiersten Kirkendolla, Jeffrey M. Burnsb and Sandra A. Billingera

aDepartment of Physical Therapy and Rehabilitation Science, University of Kansas Medical Center, Kansas City, KS, USA
bUniversity of Kansas Alzheimer’s Disease Center, Fairway, KS, USA
cDepartment of Physical Medicine and Rehabilitation, University of Kansas Medical Center, Kansas City, KS, USA

Handling Associate Editor: Jack de la Torre

Accepted 7 May 2019

Abstract

Background: Vascular health is closely related to Alzheimer’s disease (AD). Vascular function measured by flow mediated dilation (FMD) or pulsatility index (PI) can be used as marker of peripheral and central vascular health but is poorly characterized in those at risk for AD.

Objective: To assess the relationship of peripheral and central vascular function with amyloid-β (Aβ) and white matter lesion burden among cognitively normal older adults.

Methods: We enrolled participants 65 years of age and older. Using Doppler ultrasound, we assessed brachial artery FMD, and middle cerebral artery (PI). Global Aβ burden, quantified using [18F] Florbetapir PET imaging, and white matter lesion volume (WML) were used as measures of AD pathology and vascular brain injury.

Results: After adjusting for age and cardiovascular risk factors, the data (n = 83) showed a negative association between FMD and Aβ burden (β = −0.03, p < 0.001). FMD at a cut-off of 4.45% had 88% specificity and 75% sensitivity to elevated Aβ (AUC = 0.86, 95%CI: 0.77–0.95). FMD was not related to WML volume (p = 0.8), and PI was unrelated to Aβ burden or WML volume (0 > 0.4).

Conclusions: Among cognitively normal older adults, blunted peripheral vascular function, as measured by brachial artery FMD, is associated with Aβ burden. These findings provide support for further exploration into the pathophysiological relationship of vascular health and AD risk as measured by Aβ.

Keywords: Amyloid-β protein, pulsatile flow, transcranial doppler sonography, vascular endothelium

INTRODUCTION

Although amyloid-β (Aβ) accumulation is a hallmark of Alzheimer’s disease (AD) pathology, increasing evidence indicates vascular health plays an important role in the development of AD [1–5]. People with AD have increased cerebral pulsatile flow, which can be measured as pulsatility index (PI) using transcranial Doppler ultrasound [6]. Higher PI measured in the middle cerebral artery (MCA) suggests increased resistance distal to the vessel being measured and can result in reduced cerebral blood flow and tissue damage [7]. Recently, individuals

ISSN 1387-2877/16/$35.00 © 2019 – IOS Press and the authors. All rights reserved
This article is published online with Open Access and distributed under the terms of the Creative Commons Attribution License (CC-BY 4.0).
with a diagnosis of hypertension who demonstrated impaired cognition also demonstrated a significantly higher PI (above 1.1), while the group with hypertension and normal cognition had a lower PI (below 1.0). Higher PI is closely related to higher white matter lesion (WML) burden [8], a marker of small vessel ischemic disease associated with aging and cognitive decline [9–12]. Despite the current evidence, the relationship between PI with Aβ burden in cognitively normal older adults is largely unknown.

Cardiovascular risk factors (e.g., hypertension, diabetes, hyperlipidemia) damage the endothelium leading to arterial stiffness and over time may reduce cerebral blood flow [7, 13]. This vascular pathway may be synergistically contributing to the development of AD [13]. Peripheral vascular dysfunction, resulting from damage to the endothelium, could be the link between cardiovascular disease risk factors and AD [1, 14, 15] and may be the earliest event that triggers AD pathology [14, 16]. For example, recent published work reported aortic stiffness (using cardiac magnetic resonance imaging technique) was related to lower whole brain cerebral blood flow in individuals (n = 155) without cognitive impairment [17] and highlights the importance maintaining vascular health in older adults. Therefore, identifying peripheral vascular systemic indicators would be advantageous.

Peripheral vascular dysfunction is considered to be an important contributor or event in the early stages of atherosclerosis and can be assessed using a technique called flow mediated dilation (FMD) typically in the brachial artery [18]. FMD is used as a biomarker of vascular disease [19] and has predictive value for future cardiovascular events [20]. Our prior work has shown that individuals with high cardiovascular risk (stroke, diabetes) and associated cardiovascular risk factors (hypertension, dyslipidemia) have lower brachial artery FMD values [21–23] than healthy individuals (low cardiovascular risk) due to deleterious effects on vascular function. Older adults with elevated Aβ (a risk factor for AD) often have higher cardiovascular risk than those non-elevated [21, 24, 25]. Therefore, linking peripheral vascular health via FMD to brain health could provide valuable information. Currently, there is a paucity of evidence as to whether older adults with elevated Aβ have impaired peripheral vascular function. To bridge this gap in knowledge, we sought to evaluate the relationship of peripheral (FMD) and central (PI) vascular function with Aβ and white matter lesion burden among cognitively normal older adults. Lower brachial artery FMD is considered an indicator of peripheral vascular dysfunction [18, 20], while higher MCA PI is considered an indicator of vascular resistance in the smaller distal arteries [7]. Therefore, we expected that brachial artery FMD would be negatively associated with Aβ burden and WML burden. Conversely, we hypothesized that MCA PI would be positively associated with Aβ and WML burden.

METHODS

Participants

We recruited individuals as part of an ongoing Alzheimer’s and brain aging research program at the University of Kansas Alzheimer’s Disease Center (KU ADC) to characterize cerebral Aβ burden in cognitively normal older adults. For this study, the KU ADC recruited a convenience sample of individuals from the community who were interested in participating in brain aging research. Recruited individuals had no concomitant diagnosis of other neurologic conditions and no evidence of stroke or major head trauma on MRI. Inclusion criteria for the program were: 1) 65–90 years of age; 2) classified as cognitively normal/non-demented with a Clinical Dementia Rating = 0 and no clinically significant evidence of stroke or deficits on neuropsychological testing (defined as no scores greater than 1.5 standard deviations below the mean on two or more tests in the National Alzheimer’s Coordinating Center Uniform Data Set); 3) sedentary or underactive lifestyle defined as less than the recommended 150 min of moderate intensity exercise per week [26]; and 4) completion of [18F] Florbetapir positron emission tomography (PET) scan within 6 months of our experimental procedures. Exclusion criteria were: 1) Diagnostic and Statistical Manual of Mental Disorders-IV defined drug or alcohol abuse within the prior 2 years; 2) clinically significant depression or anxiety; 3) insulin-dependent diabetes; 4) myocardial infarction or symptoms of coronary artery disease within the prior 2 years; 5) acute decompensated congestive heart failure or class IV heart failure; 6) major orthopedic disability; and 7) inability to exercise due to pain or restrictions from physician.

The University of Kansas Medical Center Human Subjects Committee reviewed and approved the study. All participants provided institutionally approved written consent prior to study procedures.

Participants...
Flow mediated dilation

Brachial artery FMD is an assessment of endothelial-dependent function in response to reactive hyperemia [18]. FMD is calculated as the peak increase in arterial diameter from the baseline value in response to an acute increase in blood flow (reactive hyperemia) and expressed as percent change [27, 28]. We followed our previous published methods [21, 27, 29] and adhered to published recommendations for FMD [18]. Participants refrained from caffeinated beverages for 12 h, physical activity for 24 h, and fasted overnight. All morning medications for blood pressure were withheld [21]. All study procedures began between 7:30–9:00 am with FMD being conducted first. Participants rested in the supine position for 20 min in a quiet and dimly lit room. Room temperature was kept between 22–24°C. HR was monitored continuously using v5 on the electrocardiogram.

Images of the brachial artery were obtained with a 7.5 MHz linear array transducer (Siemens Medical Solutions, Malvern, PA). Once a satisfactory image of near and far arterial walls was obtained, the transducer was stabilized. Doppler flow angle was corrected at 60°. Baseline values for vessel diameter and blood velocity were recorded continuously for 1 min. The automated cuff (SC5D 6 × 83 cm, D.E. Hokanson, Bellevue, Washington) with a rapid inflation system (D.E. Hokanson, Bellevue, Washington) was placed distal to the olecranon process and then inflated to 200 mmHg and maintained for 5 min. Diameter and blood velocity recordings resumed 20 s prior to cuff deflation and continued for 3 min post deflation. All images were stored on a computer and analyzed (Brachial Analyzer, Medical Imaging Applications, Coralville, Iowa).

Transcranial doppler ultrasound

After the FMD procedure, participants sat upright and were fitted with a head frame and TCD ultrasound 2 MHz probe (RobotoC2MD, Multigon). If left MCA signal was not obtainable, the right side was used. Once the optimal signal was identified, recording commenced for 8 min. Data were sampled at 500 Hz. To analyze, the data were divided by R-to-R cardiac interval. For each cardiac cycle, systolic and diastolic velocities were identified, and the MCAv mean was calculated as area under the curve (cm/s). The MCAv data were then interpolated to 0.5 Hz using shape-preserving, piecewise cubic interpolation. The PI was calculated as the difference between MCA systolic and diastolic velocity divided by mean velocity for each cardiac cycle [30] during the 8 min data acquisition. Study personnel performing the TCD and FMD data collection were blinded to the Aβ status of participants.

[18F] Florbetapir PET and Aβ burden analysis

PET images were obtained on a GE Discovery ST-16 PET/CT scanner. Two 5 min duration PET brain frames were acquired continuously, approximately 50 min after [18F] Florbetapir (370 MBq) administration. Head movement was minimized by use of self-adherent wrap across the forehead. PET frames were summed and attenuation corrected. To increase tissue uptake specificity, we calculated standardized uptake value ratios (SUVR) in several individualized regions using a custom processing pipeline in SPM12 (http://www.fil.ion.ucl.ac.uk/spm). First, the PET image was co-registered to the native T1-weighted MRI (see next section). Next, we performed VBM8 unified segmentation and normalization of the anatomical T1-weighted MRI. Images were then smoothed using a 5 × 5 × 5 mm FWHM kernel. General regions of interest (ROI) from the Wake Forest Pick Atlas (anterior cingulate, posterior cingulate, precuneus, inferior medial frontal cortex, lateral temporal cortex, superior parietal cortex) were chosen for their relationship to Aβ accumulation [31]. The general ROI were transformed to native anatomical space using the inverse parameters from the unified segmentation process, then crossed with each individual’s gray matter probability map (threshold >0.5). The co-registered PET image was divided by the mean SUV of the whole cerebellum, creating an SUVR image. Mean SUVR in each individualized gray matter ROI was extracted. A global Aβ burden index was calculated as the mean of the 6 ROI. Individuals were classified as elevated if they had global Aβ burden >1.1 as this threshold has been associated with moderate-to-frequent neuritic plaques [32, 33].

MRI and WML

On a separate day, high-resolution MRI were acquired using a Siemens Skyra 3.0 Tesla scanner, including T1 (MP-RAGE; 1.0*1.0*1.2 mm voxels; TR = 2300 ms, TE = 2.98 ms, TI = 900 ms, FOV 256 mm, 9° flip angle) and T2-weighted images (FLAIR 0.9*0.9*5.0 mm voxels; TR: 9000 ms, TE = 91 ms, TI = 2500 ms, FOV 240 mm, 150° flip angle). We then used the Lesion Segment Tool,
Version 2.0.13 [34], to generate total lesion volume maps. Briefly, the LST algorithm co-registered FLAIR images to the native space anatomical image. FLAIR intensity distribution was calculated for each tissue class, and these distributions were summed to create belief maps. Based on pilot work and visual inspection, as suggested by the developer, we selected an appropriate threshold, $k = 0.125$, as the cutoff point on the gray matter belief map. This threshold was independently corroborated in a separate cohort on the same scanner (data not shown). The LST algorithm generates the white matter lesion segments and the primary WML burden measures used for analysis.

**Statistical analysis**

The data were analyzed using SPSS version 24.0 (SPSS Inc., Chicago, IL, USA). All data were tested for normal distribution using Shapiro-Wilk. Chi-square and Welch tests were used to analyze between group differences (Aβ elevated, non-elevated) for categorical and continuous variables. If the data had skewed distribution, we used Mann-Whitney U tests.

Multiple linear regression analyses were performed by making adjustment for participant age, sex, and a summed score of cardiovascular risk factors as defined by the American College of Sports Medicine (hypertension, family history of heart disease, obesity, sedentary lifestyle, age referenced to sex, diabetes, hyperlipidemia, and current smoking) [36] to determine the independent effect of FMD and PI on Aβ burden. Continuous variables with normal distribution were expressed as the mean ± standard deviation. $p$ values <0.05 were considered statistically significant. Previous work suggested that a FMD value below 5% is impaired and a marker of cardiovascular disease [37]. Therefore, we used receiver operating characteristic curves (ROC) to determine if a specific threshold of FMD would prove sensitive and specific for elevated Aβ status.

**RESULTS**

**Participant characteristics**

Participants included 83 adults who had complete datasets for all outcome measures (MRI, PET, FMD, and TCD). The time between the PET scan and the subsequent experimental protocol for TCD and FMD was 58.3 ± 43.1 days. Those with elevated Aβ were older ($p = 0.03$) with no difference between cardiovascular risk factors (Table 1).

**Flow mediated dilation**

FMD was negatively associated with Aβ burden ($\beta = -0.03, p < 0.001$) after adjusting for cardiovascular risk factors.

| Participant demographics |
|-----------------|-----------------|-----------------|---|
| Age, y          | 71.1 ± 5.2      | 73.7 ± 6.1      | 70.3 ± 4.6 | 0.03 |
| Female, n (%)   | 56 (67.5)       | 14 (70.0)       | 42 (66.7) | 0.78 |
| Education, y    | 16.7 ± 2.7      | 16.5 ± 3.3      | 16.8 ± 2.5 | 0.65 |
| BMI, kg/m²      | 26.7 ± 4.0      | 26.1 ± 4.1      | 26.8 ± 4.0 | 0.53 |
| Systolic BP (mmHg) | 132.5 ± 15.3    | 135.0 ± 16.5    | 131.7 ± 14.9 | 0.41 |
| Diastolic BP (mmHg) | 76.0 ± 8.0      | 75.5 ± 6.8      | 76.2 ± 8.4 | 0.73 |
| Mean arterial BP (mmHg) | 73.8 ± 11.6    | 73.1 ± 10.3     | 74.0 ± 12.0 | 0.69 |
| Total cholesterol (mmol/L) | 183.8 ± 33.5   | 172.6 ± 27.9   | 186.8 ± 34.4 | 0.13 |
| Hypertension, n (%) | 31 (37.3)       | 10 (50.0)       | 21 (33.3) | 0.18 |
| Diabetes, n (%)  | 3 (3.6)         | 1 (5.0)         | 2 (3.2)  | 0.70 |
| CVD, n (%)       | 5 (6.0)         | 3 (15.0)        | 2 (3.2)  | 0.05 |
| Dyslipidemia, n (%) | 46 (55.4)      | 10 (50.0)       | 36 (57.1) | 0.58 |
| Smoking, n (%)   | 2 (2.4)         | 0 (0)           | 2 (3.2)  | 0.42 |
| ACE inhibitors, n (%) | 6 (7.2)        | 2 (10.0)        | 4 (6.3)  | 0.58 |
| Calcium channel blocker, n (%) | 10 (12.0)   | 3 (15.0)        | 7 (11.1) | 0.64 |
| Beta blockers, n (%) | 12 (14.5)      | 5 (25.0)        | 7 (11.1) | 0.12 |
| Angiotensin II antagonist, n (%) | 10 (12.0) | 3 (15.0)    | 7 (11.1) | 0.64 |
| Alpha adrenergic blocking, n (%) | 2 (2.4)      | 0 (0)           | 2 (3.2)  | 0.42 |
| Antihyperlipidemic, n (%) | 39 (47.0) | 12 (60.0)    | 27 (42.9) | 0.18 |
| Sex hormones, n (%) | 14 (16.9)      | 1 (5.0)         | 13 (20.6) | 0.10 |
| Thyroid, n (%)   | 21 (25.3)       | 8 (40.0)        | 13 (20.6) | 0.08 |
| Bronchodilator, n (%) | 5 (6.0)       | 1 (5.0)         | 4 (6.3)  | 0.83 |

BMI, body mass index; BP, blood pressure; CVD, cardiovascular disease; ACE, angiotensin converting enzyme.
Fig. 1. Multiple linear regression of flow mediated dilation (FMD) and pulsatility index (PI) against Aβ burden. A) There was a negative correlation between FMD and Aβ burden across groups (p < 0.001, β = –0.03) after adjusting age, sex, and cardiovascular risk. B) There was no significant correlation between PI and Aβ burden across groups (p = 0.415) after adjusting age, sex, and cardiovascular risk.

Table 2
Flow-mediated dilation between two groups

<table>
<thead>
<tr>
<th></th>
<th>All cohort n = 83</th>
<th>Aβ elevated n = 20</th>
<th>Aβ non-elevated n = 63</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline diameter (mm)</td>
<td>3.7 ± 0.6</td>
<td>3.7 ± 0.6</td>
<td>3.7 ± 0.6</td>
<td>0.79</td>
</tr>
<tr>
<td>Peak FMD (%)</td>
<td>6.1 ± 2.8</td>
<td>3.8 ± 1.8</td>
<td>6.8 ± 2.7</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Time to peak FMD (s)</td>
<td>58.5 ± 35.2</td>
<td>72.1 ± 36.4</td>
<td>54.1 ± 33.9</td>
<td>0.05</td>
</tr>
<tr>
<td>Peak Velocity (m/s)</td>
<td>1.4 ± 0.3</td>
<td>1.4 ± 0.2</td>
<td>1.4 ± 0.3</td>
<td>0.21</td>
</tr>
</tbody>
</table>

FMD, flow mediated dilation.

Pulsatility index

PI was not associated with Aβ burden (p = 0.67, Fig. 1B) or WML volume (p = 0.42), PI was not significantly different between Aβ groups, 1.18 versus 1.06, p = 0.14 (Table 3).

DISCUSSION

The main finding of this study revealed impaired brachial artery FMD was significantly associated with increased Aβ burden among cognitively normal older adults. For every 1% FMD increase, the Aβ SUVR decreased by 0.3 independent of age and vascular disease risks. A 1% decrease in FMD is meaningful given that after the 4th and 5th decade...
of life, FMD has been reported to decrease by 0.21–0.49% with each additional year of normal aging [38]. Additionally, we found that an FMD of 4.45% was sensitive and specific to Aβ elevation. This value is close to the 5% FMD threshold, which is used as the biomarker for cardiovascular disease. Our data suggests that peripheral vascular function is concomitantly impaired with higher brain Aβ accumulation.

There is some existing evidence that brachial artery FMD is impaired in those with AD. One study compared 25 patients with AD and 24 older adult control participants [39]. Individuals enrolled were free of cardiometabolic disease, hypertension, and stroke, and were not taking medications that affected arterial vasodilation. Their results showed significantly impaired FMD in AD patients compared to controls (3.4% versus 8.4%, p < 0.01). The authors concluded that peripheral vascular dysfunction exists in those with AD without existing cardiovascular disease and associated risk factors further supporting that AD may be a “consequence of poor vascular pathology” [39] such as endothelial dysfunction. Further, these findings were independent of age and an inverse relationship between FMD and stage of AD using the Clinical Dementia Rating scale.

In the present study, our findings support and extend previous work to include peripheral vascular dysfunction in cognitively normal individuals with and without elevated Aβ. We found those with the elevated Aβ (highest risk for AD) had significantly attenuated FMD when compared to the non-elevated group (3.8% versus 6.8%, p < 0.01). Although our FMD results are similar to prior work [39] our participants presented with existing cardiovascular disease and associated risk factors, which integrate the role of lifestyle and vascular health.

Globally, there is increasing interest in the role of lifestyle factors (physical activity) and vascular health (blood pressure) to promote healthy brain aging [40]. This topic is of interest since injury to cerebral blood vessels including cerebral vessel pathology over the lifespan negatively affects the blood brain barrier and can increase inflammation [41]. Over time, blood brain barrier dysfunction and the alteration of neurovascular function such as cerebral endothelial dysfunction may accelerate Aβ accumulation by reducing perivascular clearance from the brain [41, 42]. The role of a healthy cerebrovascular system is essential to Aβ clearance. Reports suggest that the cerebrovascular system clears ~80–85% of Aβ peptides and that damage to brain endothelial cells particularly at the capillary level results in a loss of key clearance Aβ receptors such as p-glycoprotein and lipoprotein receptor-related protein 1, which leads to Aβ accumulation in the brain and further cerebrovascular impairment [41, 43–45].

Our previous work suggests in older cognitively intact adults with varying Aβ levels, a negative inverse relationship exists between cerebrovascular response during exercise and global Aβ burden, which may suggest cerebrovascular impairment is present before changes in cognition [46]. However, we only studied middle cerebral artery velocity and not other measures of cerebrovascular impairment such as PI. A higher PI has been reported in most of intracranial arteries in those with cognitive impairment [7] and AD [47, 48] when compared to nondemented individuals. In our study, PI was not associated with global Aβ burden and PI was not significantly different in those with elevated Aβ. A potential explanation may be that PI was not sensitive to detect differences in those in the earliest stages of pathology or the number of individuals with highest global Aβ burden was too small. We further acknowledge that our overall sample size was small and more individuals were characterized as non-elevated (n = 63) compared to those with elevated Aβ (n = 20).

Aging and arterial stiffness of the cerebral vessels can lead to increased PI and result in WML [8, 49, 50] especially with increased duration (in years) of hypertension [7]. The possibility exists that the individuals in this present study had experienced hypertension for a shorter duration of time or were well-controlled by medication resulting in no differences for PI and WML burden, which could be influenced by the presence of higher education and lack of racial diversity in our sample of participants. This topic warrants further exploration.
Our study has several strengths. First, our participants were well characterized as cognitively normal older adults, and we used PET imaging for to quantify global Aβ burden. Second, we used a gold standard measurement (FMD) for peripheral vascular function and used a rigorous, well-controlled FMD protocol that followed the published recommendations [18]. FMD has the advantage of being noninvasive and feasible in clinical ultrasound settings. Additionally, study personnel who conducted FMD and TCD measurements were blinded to Aβ status. In our analyses, we controlled for potential confounders including age, sex, and cardiovascular disease. As reported previously, advancing age may influence peripheral vascular function and is an important factor to control in data analysis [39].

Despite these strengths, our study has limitations that should be considered. Although these data are cross sectional, the possibility exists that higher Aβ burden may be a downstream consequence of poor vascular health. Future research should target middle age cohorts free of cognitive impairment and assess vascular health with longitudinal follow up. This would allow for a more robust assessment of vascular health and whether the result in Aβ accumulation over time exists. Second, our conclusions are limited by the homogeneity of our sample which was predominantly white, educated, cognitively typical and free of severe large vascular disease, stroke, transient ischemic attack, or traumatic brain injury. Third, we were unable to evaluate the importance of genetic risk factors such as the APOE4 allele, the strongest genetic risk factor for late-onset, sporadic AD [51]. APOE4 carriers demonstrate more evidence of neurovascular insult such as white matter hyperintensities and microbleeds than non-carriers [52]. APOE appears key to maintaining cerebrovascular integrity independent of Aβ deposition [53]; and the APOE4 allele may accelerate blood-brain barrier degradation [54]. Future work in this area should look at the role of APOE4 and other genetic factors on both central and peripheral vascular function.

Conclusions

In conclusion, the present study provides support that impaired peripheral vascular function was significantly associated with increased Aβ burden. Those with lower FMD are more likely to have elevated Aβ than those considered non-elevated. These findings provide further support for the important role of vascular health in older adults, and the need for additional exploration of the pathophysiological relationship of vascular health and Aβ accumulation.

ACKNOWLEDGMENTS

We would like to thank Torrie Cross (University of Kansas), Kelsy Schoen (University of Kansas Medical Center), Andrea Sander (University of Kansas Medical Center), and Dana Stumpff (University of Kansas Medical Center) for their help in data collection. We would also like to thank Allegra Morton (University of Kansas Medical Center) and Carolyn Kaufman (University of Kansas Medical Center) for their assistance with organizing data and tables. This study was funded in part by K01HD073718 from the Eunice Kennedy Shriver National Institute of Child Health and Human Development (Dr. Billinger) and by the American Heart Association Grant 16GRNT30450008 (Dr. Billinger). Dr. Sisante was supported in part by T32HD057850 from the Eunice Kennedy Shriver National Institute of Child Health and Human Development. This project was supported by the University of Kansas Alzheimer’s Disease Center (P30 AG035982) and by an Institutional Clinical and Translational Science Award, NIH/NCATS Grant Number UL1TR000001. Avid Radiopharmaceutical, Eli Lilly and Co., and the NIA (R01 AG043962) provided funds that supported the florbetapir imaging procedure. Its contents are solely the responsibility of the authors and do not necessarily represent the official views of the NIH or other funding agencies. Dr. Billinger receives support from the Wohlgemuth Faculty Scholar Award. The Georgia Holland Research in Exercise and Cardiovascular Health (REACH) laboratory space was supported by the Georgia Holland Endowment Fund.

Authors’ disclosures available online (https://www.j-alz.com/manuscript-disclosures/18-1268r2).

REFERENCES


