No Influence of Age-Related Hearing Loss on Brain Amyloid-β

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Abstract.

Background: Hearing loss is independently associated with a faster rate of cognitive decline in older adults and has been identified as a modifiable risk factor for dementia. The mechanism for this association is unknown, and there has been limited exploration of potential casual pathology.

Objective: Our objective was to investigate whether there was an association between degree of audiometrically measured hearing loss (HL) and brain amyloid-β (Aβ) in a pre-clinical sample.

Methods: Participants of the Australian Imaging and Biomarker Longitudinal Study (AIBL; \( n = 143 \)) underwent positron emission tomography (PET) imaging and objective measurement of hearing thresholds within 5 years of imaging, as well as cognitive assessment within 2 years of imaging in this observational cohort study.

Results: With one exception, study participants who had cognitive assessments within 2 years of their PET imaging (\( n = 113 \)) were classified as having normal cognition. There was no association between cognitive scores and degree of hearing loss, or between cognitive scores and Aβ load. No association between HL and Aβ load was found once age was controlled for. As previously reported, positive Apolipoprotein E4 (\textit{APOE4}) carrier status increased the risk of being Aβ positive (\( p = 0.002 \)).

Conclusion: Degree of HL was not associated with positive Aβ status.

Keywords: Amyloid, cognition, dementia, hearing loss, neuroimaging

INTRODUCTION

Hearing loss (HL) occurs in up to 80% of adults aged over 65 years [1, 2]. A pure tone average (PTA) of greater than 20 dB hearing level is defined as hearing loss in accordance with World Health Organization criteria [3]. Epidemiological data shows HL to be associated independently with cognitive decline and dementia in older adults [4, 5], with the risk of cognitive decline increasing with increased HL such that compared to an individual with no HL, older adults with mild HL have twice the risk of incident dementia, and those with severe HL have a five-fold risk [6]. Recent population-based cohort studies also report that individuals with onset of HL in middle age (45–64 years) have the highest dementia risk compared with older adults with normal hearing (adjusted hazard ratios of 1.90 and 1.70, respectively [7, 8]). In many older adults, HL can be ameliorated with hearing aid or cochlear implant use, showing this risk factor for dementia to be potentially modifiable.

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[6]. Despite the strong evidence for their association, the processes that link HL to faster cognitive decline in older adults remain unknown. One primary hypothesis is that HL may arise from the same early neurodegenerative processes that give rise to dementia. However, there is limited research examining relationships between HL and biomarkers associated with dementia in preclinical populations.

Alzheimer’s disease (AD) is the most common form of dementia, with estimates suggesting it accounts for 60–80% of cases [9]. Advances in understanding AD biology have allowed the application of amyloid biomarkers to identify the disease in the preclinical stages and confirm its presence in symptomatic individuals. While adults with preclinical AD do not have cognitive impairment, they do show subtle but relentless cognitive decline, faster atrophy of cortical brain areas, and an increased rate of progression to clinically recognizable mild cognitive impairment or dementia [10, 11]. Therefore, studying HL in preclinical AD provides an opportunity to understand the nature and magnitude of relationships between AD-specific neurodegeneration and HL in the absence of cognitive impairment. If AD-related neurodegeneration is related to HL, then indices of HL will be greater in preclinical AD than in age-matched controls. The current study aimed to examine the association between degree of HL and amyloid-β (Aβ) using objective audiological measurement of HL.

**MATERIALS AND METHODS**

**Participants**

Participants in this study were older adults (aged 50 years and over) enrolled in the Australian Imaging Biomarkers and Lifestyle (AIBL) Study, a prospective longitudinal cohort study of aging and AD [12]. Individuals from this cohort who did not have a mild cognitive impairment or dementia diagnosis underwent positron emission tomography (PET) imaging, had an objective audiological assessment of their hearing within 5 years of their scans, had cognitive assessment within 2 years of their PET scan, and had no diagnosed mild cognitive impairment or dementia were selected (n = 143).

**Objective assessment of hearing**

Pure tone audiometry was used to measure air and bone conduction thresholds at 500, 1000, 2000, and 4000 Hz bilaterally. Bone conduction testing was conducted when an interaural difference of > 10 dBHL in air conduction thresholds was identified. Testing was conducted by an audiologist using portable equipment in participants’ homes. Background noise levels, measured using a sound level meter, were no greater than 40 dBSPL, as per recommended practice. Participants’ ears were also examined for wax occlusion and collapsed ear canals. PTAs (average of hearing thresholds at 500, 1000, 2000, and 4000 Hz) were calculated. A PTA of greater than or equal to 20 dBHL was defined as HL, according to the World Health Organization criteria [3]. Participant self-reported hearing aid use was also noted (yes/no) and APOE genotyping was undertaken.

**PET imaging methods and analysis**

Aβ PET imaging was conducted using the Aβ tracers: 11C-PiB, 18F-florbetapir, or 18F-flutemetamol. PET acquisitions were performed 40–70 min post-tracer injection (PI) for 11C-PiB, 50–70 min PI for 18F-florbetapir, and 90–110 min PI for 18F-flutemetamol. PET images were not corrected for partial volume. All Aβ PET scans were quantified using CapAIBL [13] and the Aβ level was expressed in Centiloids (CL) as described previously [14, 15]. Aβ level was classified according to five categories: <15 Negative, 15–25 CL uncertain, 26–50 CL moderate, 51–100 CL high, 100 CL very high. 3T MRI three-dimensional magnetization-prepared rapid gradient-echo (3DMPRAGE) was used to measure hippocampal volume (HV) corrected for intracranial brain volume. A sagittal 3D FLAIR image was used to quantify white matter lesion size based on spatial location (total, lobar, and deep) [16].

**Cognitive assessment**

Cognition was assessed by psychologists using the Cogstate Cognitive Battery [17–20] which comprises assessments of psychomotor function, attention, working memory, and visual learning. The Cogstate Battery is a computerized test battery developed for repeated assessment of cognitive performance. The battery is highly reliable (test-retest reliability for each measure ranges between 0.84–0.94), facilitates minimal practice effects [17], and is relatively quick to administer (approximately 30 min, depending on ability). The Cogstate Battery can detect decline in cognitive function that does not affect function in
everyday life over even a 6-month period [21]. The battery is visually presented and is therefore suitable for people with HL. Speed and accuracy of responses are recorded and transformed on a centralized platform to give normalized data distributions [17, 18]. A composite working memory/visual learning score was derived to classify participants as cognitively normal or abnormal. In older adults, Cogstate measures of information processing speed, attention, and memory are highly sensitive to cognitive dysfunction and longitudinal cognitive decline [22, 23].

Statistical analysis

Participant characteristics were described using means, SDs, and minima and maxima for continuous variables, and counts, percentages, and cross tabulations for categorical variables. Pearson’s correlations were used to examine the associations between HL and cognitive function and between Aβ load and cognitive function. Linear regression analysis was used to investigate the relationship of Aβ load with HL, age, and APOE e4 status. Two sensitivity analyses of the regression results for the continuous measurement of Aβ load were carried out—one categorizing Aβ load as normal (less than 15) or abnormal (above 15) and fitting a Probit model with HL, age, and APOE e4 status, and a second with Aβ load in ordered categories negative/uncertain/moderate/high/very high as above and fitting an Ordered Probit model with HL, age, and APOE e4 status. Further regression models for Aβ using hearing loss by individual hearing frequencies separately (500, 1000, 2000, and 4000 Hz) were also conducted as a sensitivity analysis. A reported sex effect in the relationship between hearing loss and cognition [24] motivated an additional sensitivity analysis. Controlling for sex in each of the statistical models (Table 2) did not alter the results.

Data availability

All data has been published within this article. Data from AIBL are available to the public on request. Requests for audiometric data will be considered by the authors.

Standard protocol approvals, registrations, and patient consents

This study was carried out in accordance with the recommendations of the Australian National Health and Medical Research Council guidelines for ethical research conduct. The study protocol was approved by the University of Melbourne Behavioural and Social Sciences Human Ethics Sub-Committee (Ethics ID: 1646925). All participants gave written informed consent in accordance with the Declaration of Helsinki.

RESULTS

Clinical and pathologic characteristics of the cohort

Cohort characteristics are detailed in Table 1 according to hearing status. Participant ages ranged from 65.5 to 96.6 years. Approximately two thirds of the sample had HL (67.1%), which was on average mild in magnitude (calculated based on average PTAs re ANSI-1969 for 500, 1000, 2000, and 4000 Hz) [25], consistent with prevalence statistics for older adults in this age range. Only 27% used hearing aids. A significantly greater number of males had HL (75.9%; \( p = 0.037 \)), with 59.3% of females with HL. There was no difference in prevalence of APOE4 or cardiovascular conditions between males and females. Forty-six percent of participants had more than 15 years of education. Only one of the 113 participants with cognitive assessment data within 2 years of their PET scan had cognitive impairment. Given this, there was no difference in composite working memory/visual learning score between the groups with HL and normal hearing. There was also no association between cognitive scores and degree of hearing loss, or between cognitive scores and Aβ load. Both HL and Aβ pathology increased with age (\( r = 0.50, p = 0.000 \); \( r = 0.31, p = 0.000 \) respectively), as shown in Figs. 1 and 2. There was no significant difference in Aβ pathology between participants with and without HL (26% with HL were Aβ positive, versus 19.1% without HL, \( p > 0.05 \)). This result indicates that HL could be due to age in participants who
Relationship between HL and Aβ levels

Amyloid levels were not different between samples classified according to the presence or absence of HL (Table 1). Similarly, indices of HL were not different between samples classified according to whether amyloid levels were normal or abnormal (Table 2). However, with amyloid level and HL defined as continuous measures a significant positive relationship between the two was observed ($r = 0.23$, $p = 0.005$; Fig. 3). However, this association was not evident when the common effect of age was entered into the model. Sensitivity analyses of hearing loss by hearing frequency yielded the same result (Table 3). Analyses examining the association between amyloid level and HL with age controlled are summarized in Table 2. Model A, a linear regression for the continuous measure of Aβ load in CL, found HL was not a significant predictor for Aβ load after controlling for age and APOE4 status. This same outcome was observed in sensitivity analyses that classified amyloid level in terms of a binary classification (i.e., normal versus abnormal; Table 3, Model B), or according to the five ordinal categories (14, 15; Table 3, Model C).

### Table 2
Statistical models for Aβ using pure tone average

|                | Model A: Regression | Model B: Probit | Model C: Ordered |
|----------------|---------------------|-----------------|-----------------|
| Intercept      | −115.28             | −3.04           | 0.01            |
|                | (0.005)             | (0.103)         |                 |
| PTA (dB)       | 0.24                | 0.01            | 0.01            |
|                | (0.376)             | (0.553)         | (0.519)         |
| Age (y)        | 1.52                | 0.02            | 0.04            |
|                | (0.008)             | (0.383)         | (0.141)         |
| APOE4          | 25.23               | 1.02            | 1.01            |
|                | (0.000)             | (0.000)         | (0.000)         |
| HA use         | 8.45                | 0.47            | 0.41            |
|                | (0.255)             | (0.169)         | (0.208)         |
| n              | 137                 | 137             |                 |
| R-squared      | 0.218               |                 |                 |

For all models, statistical model coefficients are reported, with $p$ values in brackets. In model A, Aβ is treated as a continuous measure. For model B, Aβ is classified according to positive or negative status. In model C, Aβ status is classified according to the 5 categories of Centiloid status.

### Table 3
Statistical models for Aβ using hearing loss by hearing frequency

|                | Model A: Regression | Model B: Probit | Model C: Ordered |
|----------------|---------------------|-----------------|-----------------|
| Intercept      | −82.50              | −2.21           | NA              |
|                | (0.055)             | (0.270)         | (NA)            |
| 500 Hz         | 0.04                | 0.02            | 0.02            |
|                | (0.920)             | (0.381)         | (0.391)         |
| 1000 Hz        | −0.11               | −0.03           | −0.02           |
|                | (0.813)             | (0.418)         | (0.275)         |
| 2000 Hz        | 0.02                | 0.01            | 0.01            |
|                | (0.951)             | (0.546)         | (0.692)         |
| 4000 Hz        | 0.22                | 0.01            | 0.01            |
|                | (0.283)             | (0.360)         | (0.355)         |
| Age (y)        | 1.06                | 0.01            | 0.02            |
|                | (0.080)             | (0.725)         | (0.451)         |
| APOE4          | 26.40               | 1.03            | 1.03            |
|                | (0.000)             | (0.000)         | (0.000)         |
| HA use         | 9.47                | 0.56            | 0.47            |
|                | (0.212)             | (0.114)         | (0.163)         |
| R-squared      | 0.207               |                 |                 |

Analyses examining the association between amyloid level and HL with age controlled are summarized in Table 2. Model A, a linear regression for the continuous measure of Aβ load in CL, found HL was not a significant predictor for Aβ load after controlling for age and APOE4 status. This same outcome was observed in sensitivity analyses that classified amyloid level in terms of a binary classification (i.e., normal versus abnormal; Table 3, Model B), or according to the five ordinal categories (14, 15; Table 3, Model C).
DISCUSSION

In this cross-sectional cohort of participants with and without HL who underwent PET examination, HL was not associated with brain Aβ once the age of individuals was controlled, irrespective of whether amyloid level was considered as a continuous, binary, or five-level ordinal measure. Without age controlled in the analyses, a weak association between amyloid level and HL was observed. While neither amyloid level nor HL were used as inclusion criteria for this study, the number of participants with abnormally high amyloid did not differ significantly between those with and without HL. We observed in these cognitively normal older adults that both HL and amyloid level increased with increasing age. These relationships have been described previously in many studies (e.g., [2, 26–28]), therefore should be controlled in investigations of the relationship between brain Aβ and HL. The fact that both Aβ deposition and HL are correlated with age confounded the current analysis, as there was likely collinearity between them. Despite the distribution of severity of HL in this sample being representative of the general population of older adults, as the average severity of HL in this sample was only mild, statistical power to detect any effect of HL on brain Aβ may have been limited. Only 27% of participants with HL used HAs, which may have limited power to detect any effect of HA use in this sample. Strengths of this study are that it is the largest sample to date of preclinical older adults with HL (n = 142; [29–32]), objective audiometric assessment of HL, objective non-screening cognitive assessment data obtained through visual, not auditory, presentation, and the use of the Centiloid scale to allow comparison across different tracers and sites.

The data from this study and three of four others examining the relation between HL and Aβ suggest that indices of Aβ are not increased in older adults with HL. Three initial studies (two imaging, one via autopsy) investigated whether there was an association between HL and Aβ, with all reporting no association [30–32]. However, all three studies were subject to significant methodological limitations. Primary limitations include small numbers of preclinical participants with HL (e.g., n = 27, 28; number unknown for the third study, but 55 hearing aid users of unknown cognitive status), unreliable HL data for two of the studies (HL was defined according to the opinions of others who had interacted with participants in one study, and the method of HL diagnosis was unknown in another, even to the investigators), unrepresentative HL prevalence in samples, and the use of auditorily presented screening cognitive assessments to participants with HL, which may have confounded cognitive results. Given these limitations, the results of these studies do not provide a firm basis for the development of brain behavior models of HL in dementia [33].

Only one further study has examined the association between HL and Aβ, and in contrast reported a weak baseline association between HL and brain Aβ (equivalent to an increase of approx. 3 CLs for participants with HL) [29], in a younger sample (64 ± 3.5 years). The reported relationship between global amyloid SUVR (using a different radiisotope) and suprathreshold speech perception results appears likely to be driven by two poorly performing outliers, with only 15/98 participants not showing a ceiling effect on the speech perception task. However, the PTA data also demonstrated this relationship, despite the small number of participants with HL and average mild HL (n = 23; PTA = 33 dBHL). The cohort had a narrow age range (SD ± 3.5 years), therefore the usual association between HL and age observed in the current study was not seen. While this study used objective measurement of HL, the generalizability of the results is again limited by small sample size (only 23 participants had HL, and only 2 used hearing aids) and a lack of cognitive data to determine whether participants were cognitively healthy.

Models of pre-clinical disease show that the accumulation of the amyloid protein in the brain is an early initiating event in the AD cascade [27, 34]. Although predicted by positive Aβ pathology, the
onset of dementia can take decades to develop [27, 35], giving a window of time for potential modification of the cognitive outcome. Mixed [36], small [37], and sometimes no association [38] between Aβ pathology and cognitive impairment have been reported, with Aβ plaques commonly observed in the brains of clinically normal individuals, indicating that Aβ pathology is not the only driver of dementia. The sequence of events in the AD cascade, as well as the mechanisms by which Aβ pathology impacts on neuronal integrity and subsequent clinical dementia symptoms, are still unknown despite decades of research.

Previous volumetric MRI studies of brain structure as a measure of neurodegeneration have reported changes in brain structure associated with HL. Human and animal neuroimaging studies of individuals with hearing loss have shown reduced auditory cortex volume [39], white matter tract dysfunction and decline [40, 41], and accelerated whole brain atrophy [42] in degenerative patterns similar to those of AD patients. However, despite many reports of the independent association between HL and dementia, the mechanisms for this association also remain unknown. These are likely to be multiple, and there are several hypotheses. The common cause hypothesis postulates degeneration of both cognitive and non-cognitive processes due to a common neuropathic origin, while the cascade hypothesis proposes that sensory degradation causes a reduction in auditory signaling, leading to the atrophy of brain structures. The cognitive load theory hypothesizes that increased cognitive load due to perceptual difficulties that occur with HL causes diversion of cognitive resources towards speech processing rather than working memory tasks. A fourth theory proposes that dementia is caused by psychological sequelae of HL such as social isolation, depression, and loneliness, which are known risk factors for dementia [43]. A recent study has postulated a genetic relationship between HL and AD, supporting mounting evidence for the importance of inflammatory pathways in AD and suggesting that the relationship between hearing loss and AD may be due, in part, to shared genes and immune response pathways between both conditions [33]. A final recent theory postulates that changes in brain activity related to listening difficulties with HL cause irreversible molecular degenerative damage, increasing the protein pathologies associated with AD [44]. It is unlikely that dementia could cause cochlear dysfunction, as the number of adults with HL who do not have clinical pathology far exceeds the number with both pathologies, it is widely reported that peripheral HL generally precedes dementia [4–6, 45], and neuropathologic evidence of AD has not been found in the peripheral auditory pathway to date [46].

The current lack of evidence for an association between peripheral HL and AD pathology, however, does not exclude other potential explanations for the widely observed association between HL and dementia. Firstly, it is important to note that dementia is not equivalent to AD, and that as most previous studies of HL and cognitive decline have not utilized biomarkers of AD, it is possible that the observed association between HL and dementia is being driven by vascular and/or other comorbid pathologies. Further, AD pathology is not simply equivalent to brain amyloid, as is evident in the null association between Aβ load and cognitive function in the current study. There is mounting recent evidence that tau pathology may play a synergistic role in driving the clinical expression of AD [47, 48]. The timeframe for the development of clinical AD in participants of this study who are Aβ positive is currently unknown, which also leaves open the possibility that HL may influence the spread of tau pathology (age-related HL is reported to be associated with elevated cerebrospinal fluid tau levels and atrophy of the entorhinal cortex [32]), and therefore the subsequent development of AD. This theory may be proven with longitudinal follow-up of the participants in this study and in others including measurement of brain tau deposition.

Finally, it is important to note that pure tone audiometry is only one form of measurement of hearing impairment and may not necessarily give the best estimate of real-life hearing function, given the use of pure tones, not speech, in a quiet listening environment in this assessment method. All major neurodegenerative pathologies are associated with auditory brain dysfunction and have clinically relevant phenotypes of central hearing impairment [51]. In particular, auditory scene processing deficits, apparent in poor performance on speech-in-noise tests, are a common prodromal marker of more generalized cognitive decline and increased risk of AD [49–50]. Auditory dysfunction is present in AD in terms of auditory scene analysis, in Lewy body disease in the form of auditory hallucinations, impaired auditory scene analysis, tone or rhythm processing, and in the frontotemporal dementias as deficits in rhythm, pitch and timbre perception [51]. As argued previously, it is therefore plausible that some of the
The observed association between HL and dementia may be reflective of central auditory processing dysfunction/decline [51]. Also as noted previously, a focus on audiometric measures of HL may underestimate the association between auditory dysfunction and cognitive decline, particularly since impaired auditory scene analysis may not be evident in assessments conducted in quiet listening environments and can be present in the absence of any elevation of pure-tone audiometric thresholds [52]. Given the lack of central auditory processing data in the current study (as with all cross-sectional biomarker studies conducted to date), we were unable to detect whether HL in participants was peripheral only, involved central auditory processing deficits in addition to elevated audiometric thresholds, or involved only the former in the absence of audiometric HL.

The question of how HL may be related to dementia currently remains unanswered. Given the limited state of knowledge of dementia causes and the relationship between HL and neuropathologic changes, including Aβ deposition, further investigations are needed. Future studies of the association between HL and neurodegenerative biomarkers of AD (potentially including plasma neurofilament light) as a prognostic marker of both cognitive decline and changes measured via neuroimaging [53]) should include larger sample sizes (including greater numbers of participants with HL, a wider spread of HL, and hearing aid users), and neuroimaging follow-up coincident with longitudinal objective assessments of HL and cognition. Such studies would ideally follow individuals through to clinical dementia and should include the assessment of central auditory processing abilities.

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