Dietary patterns and β-amyloid deposition in aging Australian women

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Abstract

Introduction: Evidence indicates that associations between diet and Alzheimer’s disease may occur through biomarker pathways such as amyloid-β (Aβ); however, few studies have investigated dietary/Aβ relationships, and no study has investigated this relationship in women.

Methods: Dietary patterns were extrapolated for 115 participants from the Women’s Health Aging Project. Aβ deposition was measured via in vivo F-18 florbetaben positron emission tomography scanning.

Results: Participants were, on average, aged 70 years (±2.63 SD), had 13 years of education (±3.57 SD), a BMI of 28 kg/m² (±5.46 SD), and a daily energy intake of 5161 kJ (±1679.03 SD). Four dietary patterns were identified: high fat, Mediterranean, junk food, and low fat. Adherence to the junk food diet was a significant predictor of Aβ deposition (β = .10, P = .03).

Discussion: This study highlights the potential of diet to influence neurodegenerative disease and as a potential modifiable lifestyle risk factor for Alzheimer’s disease.

Keywords: Biomarkers; Alzheimer’s disease; Neuropathology; β-amyloid protein; Diet; Nutrition; Dietary pattern; Factor analysis; Women

1. Introduction

Diet may play a substantial role in the Alzheimer’s disease (AD) symptomatology and offer great potential for non-pharmacological prevention. Epidemiological evidence has suggested increased adherence to a Mediterranean diet [1], low glycemic index [2,3], and higher consumption of omega-3 polyunsaturated fatty acids [4] were associated with a decrease in AD biomarker burden. Systematic review found 50 out of 64 studies revealed an association between diet and AD incidence [5]; however, only one study has used a priori analysis to analyze dietary associations with the hallmark cerebral protein implicated in AD, β-amyloid (Aβ). In this study, dietary pattern analysis identified a pattern characterized by a higher intake of fresh fruit, vegetables, whole grains, fish, and low-fat dairy, and a lower intake of sweets, fried potatoes, processed meat, and butter was negatively associated with in vivo cerebral Aβ [6].

Furthermore, male and mixed cohort studies predominate the research, and to date, no study has investigated this relationship specifically in women. Women are more likely than men to develop AD [7], have a higher penetrance for the apolipoprotein ε-4 (APOE-ε4) allele [8], and are more likely to progress from mild cognitive impairment to AD [8]. Impacts of higher male mortality, vascular risk factors, and the postmenopausal loss of estrogenic neuroprotection suggest females are 1.5 times more likely to develop AD than men [9]. Given sex differences in AD risk, research is needed for those at greater risk of disease.

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Studies investigating in vivo AD biomarkers are needed to clarify how nutrition promotes healthy brain aging and to identify neuroprotective patterns for those at the greatest risk of AD. The objectives of this study were to identify dietary patterns using an a priori approach and investigate their associations with Aβ deposition in healthy aging Australian women. We previously reported on a lack of a relationship between a healthy Mediterranean diet and Aβ deposition [10] and hypothesized this was due to limitations of the self-reported food frequency questionnaire in measuring the potentially beneficial phytochemicals in olive oil. Given high-fat, high-glycemic diets have been associated with increased AD biomarker burden, we hypothesized that a dietary pattern characterized by high-fat, high-sugar content would be associated with an increase in cerebral Aβ pathology.

2. Methods

2.1. Study population

Participants were sought from the 2012 follow-up of the Women’s Health Ageing Project (WHAP), an epidemiologically sourced prospective study of healthy aging Australian women. WHAP is an extension of the Melbourne Women’s Midlife Health Project. Briefly, 438 women within the Melbourne metropolitan area were identified by random digit dialing in 1991. Women were eligible for the cohort if they were Australian-born, aged 45–55 years, had menstruated in the three months before recruitment, and were not taking estrogen-containing hormone replacement therapy. In 2012, participants were re-contacted and invited to participate in a late-life health study. Clinical assessments were conducted on 252 participants by trained field researchers. The clinical assessments included a battery of validated measures of physical health, sociodemographics, lifestyle, cognitive function, psychological health, and biomarkers. A complete methodology has been published elsewhere [11].

2.2. Diet

Participants completed a validated food frequency questionnaire entitled the Dietary Questionnaire for Epidemiological Studies Version 2 (DQES v2) [12]. The DQES v2 incorporates 80 food items with frequency response options on 74 of these items. The DQES v2 covers five types of dietary intake: cereals/sweets/snacks, dairy/meat/fish, fruits, vegetables, and alcoholic beverages. Data collected by the DQES v2 was used to calculate daily energy and nutrient intakes by the Cancer Council of Victoria based on Australian nutrient composition data from NUTTAB95, collated via the Composition of Foods, Australia [13]. Individuals were removed if their energy intake was reported as below 3000 kJ/day or above 20,000 kJ/day. All food items were reported in grams per day and placed into 33 food groups defined a priori (Table 1) that was similar to those used by others [14,15]. Dietary patterns were extrapolated from food groupings using iterated principal factor analysis with oblique varimax rotation due to the presumed intercollinearity and nonindependence of dietary patterns.

2.3. Imaging

In the 2012 follow-up, all WHAP participants were offered the opportunity to have cerebral imaging. Aβ deposition was measured via in vivo F-18 Florbetaben positron emission tomography.
emission tomography (PET) at the Austin Health Centre for PET in Victoria, Australia. Participants received 250 MBq of 18F-florbetaben intravenously, with a 20-minute acquisition commencing 90 minutes after injection. Standardized uptake values (SUVs) were calculated for all brain regions examined, and standard uptake value ratios (SUVRs) were generated by normalizing regional SUVs by the cerebellar cortex with atrophy correction from structural magnetic resonance imaging. Neocortical SUVR, a global index of Aβ burden, is expressed as the average SUVR of the area-weighted mean. Area-weighted means were calculated for each participant by averaging the frontal, superior parietal, lateral temporal, lateral occipital, and anterior and posterior cingulate regions. This protocol has been described elsewhere [11].

2.4. Covariates

Age (in years), education (in years), and body mass index (BMI) were collected as part of the clinical assessments in 2012. Total energy intake in kilojoules was calculated by the Cancer Council of Victoria from the DQES v2. The Consortium to Establish a Registry for Alzheimer’s Disease (CERAD) Savings Score was used as a valid indicator of cognitive ability [16]—it has been suggested as the most reliable index in differentiating cognitively normal individuals from AD [17]. Participants’ APOE genotype was determined by direct sequencing and were dichotomized as an ε4/ε4 carrier (APOE ε4/ε4), APOE ε3/ε4, and APOE ε4/ε4 or a noncarrier. Adherence to identified dietary patterns was converted from weighted factor loadings to binary presence of the APOE ε4 allele.

2.5. Statistical analysis

All analyses were conducted in STATA software on Windows operating system. Complete data were available for 115 WHAP participants, and there were no significant differences between the included (n = 115) and excluded (n = 137) cohorts. PET SUVRs displayed a positive skew that was rectified using 1/square transformation; therefore, results should be interpreted as inverse coefficient derivatives. Generalized linear models were used to assess associations between Aβ deposition and dietary patterns scores. Generalized linear models were adjusted for age in years, education in years, cognition (CERAD Savings), and binary presence of the APOE ε4 allele.

3. Results

Four dietary patterns were identified: high fat, Mediterranean, junk food, and low fat. Factor loadings (Table 2 and Fig. 1) indicated that the high-fat diet loaded heavily on food groups such as processed meats, fried fish, red meats, fried potatoes, and poultry. The Mediterranean style diet loaded chiefly on whole grains, vegetables, nuts, fish, and wine as the main source of alcohol. The unhealthy junk food pattern was characterized by high consumption of takeaway foods, added sugar, confectionary and cakes, biscuits, and sweet pastries, whereas the low-fat diet loaded heavily on low-fat dairy products, vegetables, and unsaturated spreads.

Participants’ characteristics are found in Table 3. Participants in the Mediterranean diet group (n = 31) displayed the highest level of education (14.10 ± 3.87 years), highest CERAD Savings score (72.97 ± 31.07), and lowest level of Aβ deposition (PET SUVR 1.0834 ± 0.14). Daily energy intake was highest in the high-fat group (5443.46 ± 2116.50 kJ/day) and lowest in the Mediterranean group (4677.26 ± 1242.79 kJ/day). Significant group differences were observed in education, energy intake, and CERAD

### Table 2

<table>
<thead>
<tr>
<th>Dietary Pattern</th>
<th>Food Group</th>
<th>Loadings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High Fat</strong></td>
<td>Saturated dairy products</td>
<td>0.2053</td>
</tr>
<tr>
<td></td>
<td>Full-fat dairy products</td>
<td>0.1773</td>
</tr>
<tr>
<td></td>
<td>Low-fat dairy products</td>
<td>0.1212</td>
</tr>
<tr>
<td></td>
<td>Refined grains</td>
<td>0.2868</td>
</tr>
<tr>
<td></td>
<td>Red meats</td>
<td>0.1956</td>
</tr>
<tr>
<td></td>
<td>Processed meats</td>
<td>0.2089</td>
</tr>
<tr>
<td></td>
<td>Alcohol–wine</td>
<td>0.1769</td>
</tr>
<tr>
<td><strong>Mediterranean</strong></td>
<td>Nuts</td>
<td>0.4485</td>
</tr>
<tr>
<td></td>
<td>Legumes</td>
<td>0.2923</td>
</tr>
<tr>
<td></td>
<td>Another vegetables</td>
<td>0.1550</td>
</tr>
<tr>
<td></td>
<td>Confectionery</td>
<td>0.2117</td>
</tr>
<tr>
<td><strong>Junk Food</strong></td>
<td>Added sugar</td>
<td>0.1263</td>
</tr>
<tr>
<td></td>
<td>Takeaway foods</td>
<td>0.2021</td>
</tr>
<tr>
<td></td>
<td>Fried fish</td>
<td>0.4662</td>
</tr>
<tr>
<td></td>
<td>Other fish</td>
<td>0.4029</td>
</tr>
<tr>
<td></td>
<td>Fried potatoes</td>
<td>0.4195</td>
</tr>
<tr>
<td><strong>Low Fat</strong></td>
<td>Other potato</td>
<td>0.1803</td>
</tr>
</tbody>
</table>

Rotated factor loadings for iterated principal factor analysis with oblique promax rotation. Blanks represent absent loadings (<0.1). Alcohol–spirits not shown due to not loading (>0.1) on any factor.
Savings and were therefore adjusted for in all generalized linear models.

Adherence to the junk food diet (Table 4) was a significant predictor of Aβ deposition ($\beta = .10, P = .036$) as was binary presence of the APOE ε4 allele ($\beta = .11, P = .004$). No significant interaction effects were observed in the combined effect of diet and APOE ε4 on Aβ deposition ($P = .59$). All other dietary patterns were not associated with Aβ deposition. Age, education, and cognition were also not significantly associated with Aβ deposition.

4. Discussion

In this cross-sectional study in Australian women, adherence to the junk food was a significant predictor of cerebral Aβ deposition. These results suggest that higher adherence to a high-fat, high-sugar style diet may be associated with an increased deposition of AD biomarkers and a higher risk for disease.

We observed similar cognitive status between dietary groups. However, women adhering to the Mediterranean dietary pattern displayed significantly higher cognitive scores than the other dietary groups. In the longitudinal Nurse’s Health Study, women with higher Mediterranean diet adherence had significantly higher overall cognitive status [18].

Given evidence for the cardiovascular determinants of cognitive decline [19,20], there is clear evidence for an inverse relationship between Mediterranean diet adherence and cognition; however, the cross-sectional nature of this study limits our ability to address this relationship.

Our results contribute to the growing body of evidence linking diet with AD. A high-glycemic diet has been associated with greater amyloid burden in the brain [2] and cerebrospinal fluid measures [3,21,22]. A principal component analysis on nutrient intake patterns showed consumption of omega-3 fatty acids, zinc, vitamin B-12, and vitamin D was associated with decreased amyloid deposition [6,23]. Consumption of omega-3 fatty acid supplementation has been shown to be related to tau (phosphorylated and total) and amyloid biomarkers of AD in cerebrospinal fluid [24]. Serum docosahexaenoic acid has also been inversely associated with cerebral amyloid burden [25].

Research has established that diets with higher consumption of sugar, carbohydrates, and high-glycemic foods are
associated with impaired glucose metabolism [26]. Disrupted glucose metabolism affects the production and clearance of Aβ and tau phosphorylation [27], and both insulin resistance [28] and type-2 diabetes [29] are risk factors for AD. Several animal studies have illustrated that a high-fat diet causes brain Aβ accumulation in wild-type rabbits [30] and transgenic mice [31,32]. Furthermore, human APOE isoforms have been shown to modulate glucose and metabolic pathways, with the APOE ε3/ε4 variants showing markedly reduced glucose uptake and metabolism in mouse models [33]. APOE ε2 brains demonstrated a more robust metabolic profile than APOE ε3/ε4, suggesting a physiological mechanism for its protective role against AD [33].

We speculate that the relationship observed between a high-fat, high-sugar diet and increased cerebral Aβ deposition may be modulated by impaired glucose metabolism in this female-only cohort. We believe our results suggest an impaired glucose metabolic pathway interacting with an APOE-Ab physiological mechanism. Research has been shown that APOE ε4 confers a greater risk in women than men [8]. Women with a single APOE ε4 allele have up to a four-fold increase in risk when compared with women homozygous for APOE ε3; however, men with a single APOE ε4 allele have little to no increase in risk [34]. Given animal model evidence for an APOE-mediated glucose metabolism [33], females may experience greater AD risk due to a mechanistic action in their glucose metabolism. Further research is required to elucidate the physiological mechanisms that underpin this relationship, for example, to replicate animal evidence of glucose metabolism in human models of APOE ε4 isoforms.

Our findings strengthen the hypothesis of diet being a modifiable risk factor for AD by linking amyloid deposition with an unhealthy-type diet in a female-only cohort. These findings suggest a metabolic pathway linking diet with cerebral Aβ deposition and should motivate investigations into dietary impacts on glucose metabolism by variations in presence of the APOE ε2/ε3/ε4 alleles.

Table 3
Descriptive statistics for the included participants grouped by adherence to dietary patterns identified using IPFA

<table>
<thead>
<tr>
<th>Variable</th>
<th>High fat (n = 24)</th>
<th>Mediterranean (n = 31)</th>
<th>Junk food (n = 24)</th>
<th>Low fat (n = 35)</th>
<th>Total (n = 115)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (in years)</td>
<td>69.79 ± 2.42</td>
<td>69.45 ± 2.23</td>
<td>70.41 ± 3.19</td>
<td>69.57 ± 2.70</td>
<td>69.76 ± 2.63</td>
</tr>
<tr>
<td>Education (in years)</td>
<td>12.88 ± 3.67</td>
<td>14.10 ± 3.87</td>
<td>11.50 ± 2.96</td>
<td>12.63 ± 3.36</td>
<td>12.84 ± 3.57</td>
</tr>
<tr>
<td>BMI</td>
<td>28.58 ± 6.75</td>
<td>27.43 ± 5.48</td>
<td>27.14 ± 5.58</td>
<td>29.29 ± 4.26</td>
<td>28.18 ± 5.46</td>
</tr>
<tr>
<td>Energy (kJ/day)</td>
<td>5443.46 ± 2116.50</td>
<td>4809.79 ± 1145.51</td>
<td>6035.40 ± 1993.21</td>
<td>4677.26 ± 1242.79</td>
<td>5160.53 ± 1679.03</td>
</tr>
</tbody>
</table>

Table 4
Generalized linear model for independent variables (PET SUVR) and four dietary patterns identified using iterative principal factor analysis. (95% CIs shown; n = 114)

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient</th>
<th>Std. Err.</th>
<th>CI lower</th>
<th>CI higher</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET SUVR</td>
<td>-0.00705</td>
<td>0.04372</td>
<td>-0.09273</td>
<td>0.07864</td>
</tr>
<tr>
<td>Mediterranean</td>
<td>0.06390</td>
<td>0.04349</td>
<td>0.142</td>
<td>-0.02135</td>
</tr>
<tr>
<td>Junk food</td>
<td>-0.09740</td>
<td>0.04511</td>
<td>-0.18582</td>
<td>-0.00898</td>
</tr>
<tr>
<td>Low fat</td>
<td>-0.02338</td>
<td>0.04092</td>
<td>-0.085428</td>
<td>0.10103</td>
</tr>
<tr>
<td>Age (in years)</td>
<td>-0.00120</td>
<td>0.00702</td>
<td>-0.01495</td>
<td>0.01256</td>
</tr>
<tr>
<td>Education (in years)</td>
<td>0.00139</td>
<td>0.00502</td>
<td>0.00845</td>
<td>0.01125</td>
</tr>
<tr>
<td>BMI</td>
<td>-0.00076</td>
<td>0.00326</td>
<td>-0.00714</td>
<td>0.00563</td>
</tr>
<tr>
<td>Energy (kJ/day)</td>
<td>-0.000001</td>
<td>0.00001</td>
<td>-0.00003</td>
<td>0.000001</td>
</tr>
<tr>
<td>APOE Presence</td>
<td>-0.10916</td>
<td>0.03919</td>
<td>-0.18598</td>
<td>-0.03233</td>
</tr>
<tr>
<td>CERAD Savings Score</td>
<td>0.000125</td>
<td>0.00065</td>
<td>0.005</td>
<td>0.00002</td>
</tr>
</tbody>
</table>

Abbreviations: APOE, apolipoprotein E; BMI, body mass index; CERAD, Consortium to Establish a Registry for Alzheimer’s Disease; CI, confidence interval; PET, positron emission tomography; SUVR, standard uptake value ratio.

NOTE. Bold indicates statistical significance (P < 0.05). CIs are for coefficient. Analysis adjusted for age in years, education in years, cognition (CERAD Savings score), and binary presence of the APOE ε4 allele.

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C.S. has provided clinical consultancy and been on scientific advisory committees for the Australian Commonwealth Scientific and Industrial Research Organization, Alzheimer’s Australia, University of Melbourne, and other relationships that are subject to confidentiality clauses. She has been a named chief investigator on investigator-driven collaborative research projects in partnership with Pfizer, Merck, Bayer, and GE. She may accrue revenues from patent in pharmacogenomics prediction of seizure recurrence. The other authors have no conflict of interest to report.

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