Mediterranean Diet and Risk for Alzheimer’s Disease

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Abstract

Objective—Previous research in Alzheimer’s disease (AD) has focused on individual dietary components. There is converging evidence that composite dietary patterns such as the Mediterranean diet (MeDi) is related to lower risk for cardiovascular disease, several forms of cancer, and overall mortality. We sought to investigate the association between MeDi and risk for AD.

Methods—A total of 2,258 community-based nondemented individuals in New York were prospectively evaluated every 1.5 years. Adherence to the MeDi (zero- to nine-point scale with higher scores indicating higher adherence) was the main predictor in models that were adjusted for cohort, age, sex, ethnicity, education, apolipoprotein E genotype, caloric intake, smoking, medical comorbidity index, and body mass index.

Results—There were 262 incident AD cases during the course of 4 (±3.0; range, 0.2–13.9) years of follow-up. Higher adherence to the MeDi was associated with lower risk for AD (hazard ratio, 0.91; 95% confidence interval, 0.83–0.98; p = 0.015). Compared with subjects in the lowest MeDi tertile, subjects in the middle MeDi tertile had a hazard ratio of 0.85 (95% confidence interval, 0.63–1.16) and those at the highest tertile had a hazard ratio of 0.60 (95% confidence interval, 0.42–0.87) for AD (p for trend = 0.007).

Interpretation—We conclude that higher adherence to the MeDi is associated with a reduction in risk for AD.

Diet may play an important role in the causation and prevention of AD.¹,² Dietary restriction in animals extends their life span³ and increases the resistance of neurons to degeneration.³ However, epidemiological data on diet and AD have been conflicting. Higher intake of vitamin C,⁴–⁶ vitamin E,⁴–⁸ flavonoids,⁹ unsaturated fatty acids,¹⁰–¹² fish¹²,¹³; higher levels of vitamin B12¹⁴–¹⁶ and folate¹⁵,¹⁶; modest to moderate ethanol¹⁷–²¹; and lower total fats²²,²³ have been related to a lower risk for AD or slower cognitive decline. At the same time, other studies have found that the risk for AD or cognitive decline is not associated to intake of antioxidants, such as vitamin C,²⁴ vitamin E,²⁴ and carotenes,²⁴ fats,²⁵ or levels of vitamin B12.²⁶

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There is paucity of data regarding the effect of composite dietary patterns (rather than individual foods or nutrients) on the risk for AD. Dietary pattern analysis in relation to many other diseases (ie, cirrhosis or various cancers) has recently received growing attention because individuals do not consume foods or nutrients in isolation, but rather as components of their daily diet. Defining diet by dietary patterns has the ability to capture its multidimensionality whereas reducing its apparent complexity because patterns can integrate complex or subtle interactive effects of many dietary constituents and bypass problems generated by multiple testing and the high correlations that may exist among these constituents.27 Dietary patterns can be developed a priori from previous knowledge concerning a favorable or adverse health effect of various dietary constituents.

One such dietary pattern is the Mediterranean diet (MeDi), which has received increased attention in recent years because of converging ecological,28 analytical-observational,29–33 and interventional34,35 evidence relating it to lower risk for cardiovascular disease, several forms of cancer, and overall mortality. The MeDi is characterized by high intake of vegetables, legumes, fruits, and cereals; high intake of unsaturated fatty acids (mostly in the form of olive oil), but low intake of saturated fatty acids; a moderately high intake of fish; a low-to-moderate intake of dairy products (mostly cheese or yogurt); a low intake of meat and poultry; and a regular but moderate amount of ethanol, primarily in the form of wine and generally during meals.31 Therefore, the MeDi appears to include many of the components reported as potentially beneficial for AD and cognitive performance.

We examined the association between MeDi and AD using data from the Washington Heights-Inwood Columbia Aging Project (WHICAP). We hypothesized that higher adherence to the MeDi would be associated with lower risk for AD.

**Subjects and Methods**

**Sample and Procedures**

This study included participants of two related cohorts recruited in 1992 (WHICAP 1992) and 1999 (WHICAP 1999), which were identified (via ethnicity and age stratification processes) from a probability sample of Medicare beneficiaries residing in an area of three contiguous census tracts within a geographically defined area of northern Manhattan.23,24,36,37 The same assessments and study procedures were used in both cohorts. At entry, a physician elicited each subject’s medical and neurological history and conducted a standardized physical and neurological examination. All available ancillary information (medical charts, computed tomography scans, or magnetic resonance images) was considered in the evaluation. A global summary score on the Clinical Dementia Rating (CDR)38 scale also was assigned. Each subject also underwent a structured in-person interview including an assessment of health and function and a neuropsychological battery.39 The neuropsychological battery contained tests of memory (short- and long-term verbal and nonverbal), orientation, abstract reasoning (verbal and non-verbal); language (naming, verbal fluency, comprehension, and repetition), and construction (copying and matching).

A consensus diagnosis for the presence or absence of dementia was made at a diagnostic conference of neurologists and neuropsychologists where information of all the above evaluations was presented. Evidence of cognitive deficit (based on the neuropsychological scores as described above), evidence of impairment in social or occupational function (as assessed by the Blessed Dementia Rating Scale, the Schwab and England Activities of Daily Living Scale, and the physician’s assessment), and evidence of cognitive and social-occupational function decline compared with the past were the criteria used for the diagnosis of dementia as required by the Diagnostic and Statistical Manual of Mental Disorders, Revised Third Edition (DSM-III-R). The type of dementia was subsequently determined. For
the diagnosis of probable or possible AD, the criteria of the National Institute of Neurological and Communicative Disorders and Stroke-Alzheimer’s Disease and Related Disorders Association were used. Because in these criteria stroke does not preclude the diagnosis of AD (unless cerebrovascular disease was considered the primary cause of the dementia), the diagnosis of AD with concomitant stroke also was assigned. Dietary data were not available to the consensus panel and were not considered in the diagnostic process.

Subjects were followed up at intervals of approximately 1.5 years, repeating the baseline examination and consensus diagnosis at each follow-up. The initial sample for this study was 4,166 individuals, of whom 3,436 were without dementia at initial evaluation (Fig 1). Because the dietary assessment component was added after initiation of the study, dietary information was missing for some subjects (n = 527). An additional 24 subjects were excluded due to incomplete dietary information. From the remaining 2,885 subjects, follow-up was not available for 627 (including 101 because they died relatively soon after the baseline assessment). Thus, the initial analytic sample for the survival analyses comprised 2,258 subjects (1,037 [46%] from the WHICAP 1992 and 1,221 [54%] from the WHICAP 1999 cohort).

Evaluation

**PREDICTORS**

**Diet:** Dietary data regarding average food consumption over the past year were obtained using a 61-item version of Willett’s semiquantitative food frequency questionnaire (SFFQ; Channing Laboratory, Cambridge, MA). Trained interviewers administered the SFFQ in English or Spanish. We have previously reported validity (using two 7-day food records) and reliability (using two 3-month frequency assessments) of various components of the SFFQ in WHICAP. We followed the method that Trichopoulou and colleagues described previously for the construction of the MeDi score. More specifically, we first regressed caloric intake (measured in kilocalories) and calculated the derived residuals of daily gram intake (as Willet and Stampfer recommended) for each of the following seven categories (which define the components of the MeDi that Trichopoulou and colleagues defined previously): dairy, meat, fruits, vegetables, legumes, cereals, and fish. A value of 0 or 1 was assigned to each of these seven groups, using sex-specific medians as cutoffs. For beneficial components (fruits, vegetables, legumes, cereals, and fish), individuals whose consumption was below the median were assigned a value of 0, and individuals whose consumption was at or above the median were assigned a value of 1. For components presumed to be detrimental (meat and dairy products), individuals whose consumption was below the median were assigned a value of 1, and individuals whose consumption was at or above the median were assigned a value of 0. For fat intake (the eighth food category), we used the ratio of daily consumption (in grams) of monounsaturated lipids to saturated lipids (again using sex-specific median cutoffs for assignment values of 0 for low and 1 for high). For alcohol intake (the ninth food category), subjects were assigned a score of 0 for either no (0gm/day) or more than moderate (≥30gm/day) consumption, and a value of 1 for mild-moderate alcohol consumption (>0 to <30gm/day). This agrees with Trichopoulou’s previous reports that consider moderate amount of alcohol consumption as another characteristic component of the MeDi. We classified alcohol consumption dichotomously, because of the skewed distribution of alcohol in our population (68% reporting no alcohol intake, 31% reporting less than 30gm/day [mild-to-moderate intake], and 1% reporting ≥30gm/day [heavy intake]). The MeDi score was generated for each participant by adding the scores in the food categories (theoretically ranging from 0–9) with the higher score indicating stricter adherence to the MeDi. Median intake of consumption for MeDi food...
categories is presented in Table 1. We used MeDi score from the baseline visit as the main predictor in the survival analyses.

**Covariates:** Age (years), education (years), caloric intake (kcal), and body mass index (weight in kilograms divided by height in square meters \([\text{kg/m}^2]\)) were used as continuous variables. We also considered cohort (1992 cohort as reference), sex (men as reference), and smoking status at baseline evaluation (no smoking as reference). Ethnic group was based on self-report using the format of the 1990 census. Participants then were assigned to one of four groups: Black (non-Hispanic), Hispanic, White (non-Hispanic), or Other. Ethnicity was used as a dummy variable with White (non-Hispanic) as the reference. Apolipoprotein E (APOE) genotype was used dichotomously: absence of \(\varepsilon_4\) allele versus presence of either 1 or 2 \(\varepsilon_4\) alleles.

A modified version of the Charlson Index of Comorbidity (referred to as “comorbidity index”) was included as a continuous variable. It included items for myocardial infarct, congestive heart failure, peripheral vascular disease, hypertension, chronic obstructive pulmonary disease, arthritis, gastrointestinal disease, mild liver disease, diabetes, chronic renal disease, and systemic malignancy from the baseline visit. All items received weights of one, with the exception of chronic renal disease and systemic malignancy, which were weighted two.

**STATISTICAL ANALYSES**—Baseline characteristics of patients by availability of follow-up, outcome of interest, and MeDi tertiles were compared using \(t\) test or analysis of variance for continuous variables and \(\chi^2\) test for categorical variables.

**Mediterranean Diet Score Stability:** We used generalized estimating equations (GEEs) to test whether there were significant changes of MeDi score over time for a subset of subjects with more than one dietary assessment. These repeated dietary assessments were performed to obtain information on the temporal stability of dietary reporting. GEE takes into account the multiple visits per subject and that the characteristics of the same individual over time are likely to be correlated. The repeated measures for each subject are treated as a cluster. The GEE model included the MeDi score as the dependent variable and time (years) as predictor. A significant time effect would indicate a significant change of MeDi score over time. We performed separate exploratory analyses for subjects who experienced development of AD and for ones who remained nondemented.

**Survival Analyses:** We calculated Cox proportional hazards models with AD as the dichotomous outcome. Subjects who experienced development of other dementias \((n = 32)\) were excluded from these analyses. In exploratory analyses that censored (rather than excluded) these subjects, the results were unchanged (these analyses are not presented). The time-to-event variable was time from recording of baseline diet to first visit of dementia diagnosis; subjects who did not experience development of dementia were censored at the time of their last follow-up. The main predictor was MeDi score (from the baseline visit) as a continuous variable initially and in tertiles form subsequently (used for trend test calculation). We simultaneously adjusted for the following variables: cohort, age at intake in the study, sex, ethnicity, education, APOE, smoking, comorbidity index, and body mass index. Although caloric intake adjusted residuals were used in the MeDi score calculation, we also included caloric intake as a covariate in the models (as Willet and colleagues recommended). All predictors were used as time-constant covariates. We performed two additional sets of sensitivity analyses. First, we recalculated the models using as dementia diagnosis date the last date when subjects were considered to be cognitively nondemented. Second, we recalculated the models using as dementia date the midpoint between last nondementia diagnosis and first dementia diagnosis.
In other exploratory models, we repeated the analyses excluding subjects who were diagnosed as AD with stroke (ie, using only AD without stroke as the outcome). To increase our confidence that the degree of adherence to MeDi was not affected by early, subclinical dementia process, we performed additional analyses. First, we recomputed the Cox models excluding subjects with mild cognitive deficits at baseline (ie, CDR = 0.5). Second, we excluded both subjects with mild cognitive deficits at baseline (ie, CDR = 0.5) and subjects who were followed for less than 2 years.

To examine whether possible associations between MeDi and risk for AD were driven by associations of particular food categories, in further supplementary analyses, we included all nine individual components used to calculate the MeDi score in stepwise forward selection Cox models (entry criterion: \( p < 0.05 \); removal criterion: \( p > 0.05 \)). The Cox models fulfilled the proportionality assumption (martingale residuals method).

**RATES OF COGNITIVE DECLINE IN RELATION TO THE MEDITERRANEAN DIET**—The primary outcome in these analyses was rate of decline in cognition as assessed at each study visit. Using 12 neuropsychological tests from the administered battery,\(^{48-54}\) we calculated a composite cognitive measure. Details about this procedure have been published previously.\(^8\) In brief, to derive the composite measure, we first transformed each of the 12 individual cognitive tests into Z-scores. Means and standard deviations for each test were calculated from baseline scores of nondemented subjects, and Z-scores were averaged initially in cognitive domains, which were subsequently averaged again to produce the composite cognitive score. The Z-score of the composite cognitive measure at each evaluation was the primary outcome.

We used GEEs to test whether MeDi adherence was associated with differential rates of cognitive change. The repeated cognitive performance scores for each subject were treated as a cluster. The GEE models included the composite cognitive measure as the dependent variable and, as predictors, cohort, age, sex, education, ethnicity, baseline cognitive performance, MeDi at baseline, time (in years from baseline assessment), and a MeDi \( \times \) time interaction. A significant interaction term would indicate differential rates of change in cognitive function as a function of MeDi adherence.

**Results**

**Missing Data Analyses**

Compared with subjects with available dietary information (n = 527) (see Fig 1) had slightly lower education (9.1 vs 9.9 years; \( p = 0.001 \)). Subjects with missing dietary information also had higher proportions of dementia (17.5 vs 11%; \( p < 0.001 \)) and higher mortality (32 vs 18%; \( p < 0.001 \)), which may be related to that the dietary assessment was added after initiation of the study and was not available for subjects recruited earlier on. There were no significant differences between subjects with missing and those with available dietary information in age (76.7 vs 77; \( p = 0.30 \)), sex (33% male vs 33% male; \( p = 0.68 \)), ethnicity (White 25%, Black 31%, Hispanic 43%, Other 1% vs White 27%, Black 33%, Hispanic 39%, Other 1%; \( p = 0.24 \)), medical comorbidity index (2.1 vs 2.0; \( p = 0.27 \)), or APOE genotype (\( \epsilon 4 \) carriers 27 vs 28%; \( p = 0.79 \)).

Compared with subjects with available cognitive follow-up (n = 2,258), subjects with missing follow-up (n = 627) (see Fig 1) were slightly younger (76.4 vs 77.2 years; \( p = 0.01 \)), had lower education (9.4 vs 10.0 years; \( p = 0.005 \)), more medical comorbidities (2.2 vs 1.9; \( p = 0.002 \)), and higher mortality (30 vs 15%; \( p < 0.001 \)). There were no significant differences between subjects with missing and those with available follow-up in caloric intake (1,466 vs 1,428; \( p = 0.12 \)), sex (35% male vs 33% male; \( p = 0.42 \)), ethnicity (White
26%, Black 36%, Hispanic 38%, Other 1% vs White 28%, Black 33%, Hispanic 38%, Other 2%; p = 0.39), or APOE genotype (ε4 carriers 29 vs 28%; p = 0.67). Most important, there was no difference in MeDi score (4.3 vs 4.4; p = 0.21).

Examining evaluations for subjects who had available follow-up, 76 subjects were missing 1 evaluation, 11 were missing 1 evaluation, and 2 were missing 3 evaluations. Examining, in particular, the time period either between the dementia incidence evaluation and the evaluation preceding that (for AD subjects) or between the last evaluation and the evaluation preceding that (for nondemented subjects), only 23 subjects (7 AD and 16 nondemented) had 1 missing evaluation and only two subjects (1 AD and 1 nondemented) had 2 missing evaluations. Therefore, missing evaluations for subjects with available follow-up was a rare phenomenon in our study.

Stability of Mediterranean Diet Score

There were 390 subjects with multiple dietary assessments who did not experience development of AD or other dementia during follow-up. There were 2 dietary assessments available for 308 subjects, 3 for 71 subjects, and 4 for 11 subjects. The mean time interval between dietary assessments was 7.1 years (standard deviation, 2.36; range, 1–12.8). The reported MeDi score was stable (β = −0.01; p = 0.41).

There were 89 subjects with multiple dietary assessments who developed AD during follow-up; there were 2 assessments available for 78 participants, 3 for 8 participants, and 4 for 3 participants. The mean time interval between dietary assessments was 8.1 years (standard deviation, 1.9; range, 1.8–11.9). Again, the reported MeDi score did not change over time (β = −0.05; p = 0.09).

Clinical-Demographic-Dietary Characteristics

Subjects were followed (until AD incidence or last follow-up for subjects who remained without demented) for an average of 4.0 (±3.0; range, 0.2–13.9) years. Overall, 294 subjects experienced development of incident dementia; AD developed in 262 of these subjects (184/262 had AD without stroke).

Compared with subjects who remained without dementia, subjects in whom AD had developed at follow-up were older, less educated, and had a lower body mass index (Table 2). There was a higher proportion of Hispanics and a lower proportion of Whites (non-Hispanic) among subjects who experienced development of AD. Presence of ε4 allele was more frequent and caloric intake was higher for subjects in whom AD developed compared with subjects who remained without dementia, but the associations were not significant. Subjects who did and did not acquire AD did not differ in sex, medical comorbidity index, or smoking status. Compared with subjects who remained nondemented, subjects who experienced development of AD had lower MeDi scores.

There was no association between MeDi score and age, sex, education, APOE genotype, medical comorbidity index, or body mass index (Table 3). Hispanics adhered more and Blacks less to the MeDi pattern. Subjects adhering more to the MeDi tended to smoke less and had lower caloric intake.

Mediterranean Diet and Risk for Alzheimer’s Disease and Cognitive Decline

Higher adherence to the MeDi was associated with significantly lower risk for development of AD (Table 4 and Fig 2). The results were similar in adjusted and unadjusted models. Each additional unit of the MeDi score was associated with 9 to 10% less risk for development of AD. Compared with subjects in the lowest MeDi tertile (low adherence to the MeDi),
subjects in the middle MeDi score tertile had 15 to 21% less risk for development of AD, whereas those at the highest tertile (high adherence to the MeDi) had 39 to 40% less risk for development of AD, with a significant trend for a dose–response effect. In sensitivity analyses using alternative dementia diagnosis dates (either last date when subjects were deemed nondemented or the midpoint between last nondementia and first dementia diagnosis) the associations were unchanged.

The coefficients remained virtually unchanged and the associations significant in models excluding subjects with baseline evidence of mild cognitive impairment (CDR $p = 0.5$): 1,898 subjects at risk with 156 incident AD cases; hazard ratio, 0.88 (95% confidence interval, 0.80–0.97); $p = 0.007$, tertile analyses; $p$ for trend $= 0.018$. When both persons with CDR $= 0.5$ and those followed for less than 2 years were excluded, the coefficients remained essentially unchanged: 1,575 subjects at risk with 134 incident AD cases; hazard ratio: 0.89 (95% confidence interval, 0.80–0.98); $p = 0.020$, tertile analyses; $p$ for trend $= 0.027$. Results were similar in adjusted models.

When the same models were run with probable AD without stroke as the outcome (excluding AD with coexisting stroke, n = 78), the associations were unchanged: 2,144 subjects at risk with 184 incident probable AD without stroke; hazard ratio, 0.90 (95% confidence interval, 0.83–0.98); $p = 0.015$, tertile analyses; $p$ for trend $= 0.018$. Adjusted models produced similar results.

Regarding associations of other nutrition-related elements with risk for AD, and in agreement with previous publications, lower caloric intake (0.9997 for each additional kilocalorie; range, 0.9994–0.9999; $p = 0.007$) and higher body mass index (0.96; range, 0.93–0.98; $p = 0.002$) were related to lower risk for AD in the adjusted models.

In unadjusted Cox models that included the nine individual dietary components of the MeDi as predictors, mild-to-moderate alcohol consumption (0.61 [0.45–0.82]; $p = 0.001$) and higher vegetable consumption (0.76 [0.60–0.97]; $p = 0.030$) were associated with decreased risk for AD. In adjusted models, none of the individual components was a significant AD predictor.

In GEE analyses, there was a significant MeDi × time interaction $\beta = 0.003; p = 0.047$) indicating an association between higher adherence to MeDi and slower cognitive decline: each additional unit of adherence to the MeDi was associated with 0.3% of a standard deviation less decline per year.

**Discussion**

This study suggests that higher adherence to the MeDi is associated with a reduction in risk for AD and slower cognitive decline. The gradual reduction in AD risk for higher tertiles of MeDi adherence also suggests a possible dose–response effect. The associations between MeDi and incident AD remained unchanged and significant even when simultaneously adjusting for the most commonly considered potential confounders for AD, such as age, sex, ethnicity, education, APOE genotype, caloric intake, and body mass index. Higher adherence to MeDi reduced risk for probable AD either with or without coexisting stroke.

The association between high adherence to the MeDi and lower risk for AD may be mediated by the composite effect of some of its beneficial components, such as higher intake of fish, fruits, and vegetables rich in antioxidants such as vitamin C, vitamin E, and flavonoids and higher intake of unsaturated fatty acids. Some of the inconsistencies regarding some of the above dietary elements and risk for AD in the existing literature may be a result of failure to consider possible additive and interactive (antagonistic
or synergistic) effects among nutritional components, which may be captured in a composite dietary pattern such as the MeDi. For example, the effect of fish consumption in reducing blood pressure and blood lipids appears to be much more pronounced in subjects following a low-fat diet. When individual components used to derive the MeDi pattern were examined in our study, mild-to-moderate alcohol consumption and higher vegetable intake were associated with decreased risk for AD in unadjusted models. Nevertheless, none of the individual components was a significant AD predictor when other confounders were considered. These results strengthen even further our initial hypothesis that composite dietary patterns can capture dimensions of nutrition that may be missed by individual components, and that an overall dietary pattern is likely to have a greater effect on health than a single nutrient.

The MeDi may play a role in multiple potential mechanisms including oxidative stress and inflammation, which are both important in the pathogenesis of AD. Complex phenols and many other substances with important antioxidant properties such as vitamin C, vitamin E, and carotenoids are found in high concentrations in the typical components of the MeDi. In the Attica epidemiological study, participants on the highest tertile of the MeDi score had 20% lower CRP levels and 17% lower interleukin-6 serum levels. Higher adherence to the MeDi also has been associated with significant reduction in various other inflammatory and coagulation markers including white blood cell counts and fibrinogen levels.

Given the growing evidence for contribution of vascular risk factors in AD risk, vascular mechanisms are important to consider. There is strong evidence relating the MeDi to lower risk for vascular risk factors such as hypertension, dyslipidemia, and diabetes. Subjects in the highest tertile of MeDi adherence have also been reported to have 15% lower homocysteine levels. Thus, vascular variables are likely to be in the causal pathway between MeDi and AD and should be considered as possible mediators. Because mediators, as contrasted with confounders, should not be controlled for in the statistical analysis, and in accordance with previous MeDi-related analytical approaches, we did not include specific vascular variables in our models.

Although most studies of the MeDi have been conducted in Mediterranean populations, recent studies have indicated MeDi-related health benefits in other populations such as northern European, Indian, and Australian populations, suggesting that the advantages of the MeDi are transferable to other populations. Our study provided the opportunity to examine the effect of MeDi in a multiethnic community in the United States, and our results support the notion that the beneficial effects of the MeDi are generalizable to different populations.

This study has limitations. The use of an a priori distribution-derived MeDi score assumes underlying monotonic effects, does not address possible thresholds or the shape of the underlying curve, and weighs equally the underlying individual food categories, which, in turn, are composed of different numbers of food constituents. Frequencies of food intake are based on relatively few diet constituents, which may underestimate the overall quantity of food in each food category, and a common limitation of studies of diet and disease is misclassification of exposure due to limited accuracy. However, assuming that the measurement error was random, our results may actually underestimate the association between high MeDi adherence and lower AD risk.

Despite the use of standard criteria, the diagnostic expertise of our center, and the thorough workup, there is always the possibility of disease misclassification bias. We partially addressed this issue by conducting secondary analyses considering only probable AD without stroke as the outcome, and our results were virtually unchanged. Subjects who were
lost to follow-up were younger, less educated, and had more medical comorbidities, but MeDi was not related to age, education, or medical comorbidity burden. Also, the MeDi scores did not differ between subjects who remained in the study and those lost to follow-up. Therefore, it does not appear likely that our results could be explained by biases related to loss to follow-up.

It is possible that diet is related to socioeconomic status or to other habits or characteristics related to better health and a lower risk for AD. In our data, MeDi was not related to education or the overall burden of medical comorbidities, but was related to ethnicity and smoking. We addressed this by adjusting for years of education, ethnic group, medical comorbidity index, and smoking, but we cannot completely rule out residual confounding as an explanation for our findings. Another confounder usually considered in nutritional epidemiology is caloric intake, and higher MeDi adherence was related to lower caloric intake in our data. We used caloric intake adjusted residuals for MeDi calculation and also included caloric intake as a covariate (as Willet and Stampfer recommended). We found that higher MeDi adherence was related to a lower risk for AD over and above caloric intake.

Despite an average follow-up of 4 years, several studies have shown that subtle cognitive changes antedate the clinical diagnosis of AD by many years. Therefore, it is still possible that lower adherence to the MeDi could represent a consequence and not a cause of AD. We addressed this possibility in several ways. We found remarkable stability in MeDi scores over intervals greater than 7 years for subjects with multiple dietary assessments regardless of dementia status. In addition, the associations between MeDi score and incident AD did not change when subjects with mild cognitive deficits at baseline and subjects with less than 2 years of follow-up were excluded. Thus, bias due to preclinical disease does not appear to be a likely explanation for our findings.

Confidence in our findings is also strengthened by the following factors. Dietary data were collected with a previously validated instrument that was used widely in epidemiological studies. We used an a priori developed dietary pattern. Measures for multiple potential AD risk factors have been recorded carefully and adjusted for in the analyses. The diagnosis of AD took place in a university hospital with expertise in dementia and was based on comprehensive assessment and standard research criteria. The patients were followed prospectively at relatively short intervals. The study is community-based and the population is multiethnic, increasing the external validity of the findings.

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References


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Fig 1.
Fig 2.
Survival curves based on Cox analysis comparing cumulative Alzheimer’s disease (AD) incidence in subjects belonging to each Mediterranean diet (MeDi) tertile ($p$ for trend = 0.007). Low tertile (score 0–3; light gray lines) corresponds to lower adherence to MeDi, middle tertile (score 4–5; dark gray lines) to middle adherence, and high tertile (score 6–9; black lines) to higher adherence. Figure is derived from a model that uses all subjects and is adjusted for cohort, age, sex, ethnicity, education, apolipoprotein E genotype, caloric intake, smoking, comorbidity index, and body mass index. Duration of follow-up is truncated at 10 years.
### Table 1
Median Daily Intake for Individual Food Categories by Mediterranean Diet Score Tertiles and Overall

<table>
<thead>
<tr>
<th>Food Categories</th>
<th>Low Tertile (MeDi score 0–3)</th>
<th>Middle Tertile (MeDi score 4–5)</th>
<th>High Tertile (MeDi score 6–9)</th>
<th>All (25th, 75th percentiles)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dairy, gm/day</td>
<td>246</td>
<td>174</td>
<td>151</td>
<td>182 (128–292)</td>
</tr>
<tr>
<td>Meat, gm/day</td>
<td>101</td>
<td>86</td>
<td>65</td>
<td>85 (60–119)</td>
</tr>
<tr>
<td>Vegetable, gm/day</td>
<td>165</td>
<td>202</td>
<td>243</td>
<td>197 (153–250)</td>
</tr>
<tr>
<td>Fruit, gm/day</td>
<td>406</td>
<td>471</td>
<td>556</td>
<td>472 (372–582)</td>
</tr>
<tr>
<td>Legumes, gm/day</td>
<td>44</td>
<td>58</td>
<td>78</td>
<td>57 (38–90)</td>
</tr>
<tr>
<td>Cereal, gm/day</td>
<td>155</td>
<td>186</td>
<td>215</td>
<td>184 (140–233)</td>
</tr>
<tr>
<td>Fish, gm/day</td>
<td>15</td>
<td>21</td>
<td>47</td>
<td>20 (14–47)</td>
</tr>
<tr>
<td>MUFA/SFA ratio</td>
<td>0.57</td>
<td>0.82</td>
<td>0.97</td>
<td>0.80 (0.18–1.37)</td>
</tr>
<tr>
<td>Mild-to-moderate ethanol, %</td>
<td>21</td>
<td>33</td>
<td>45</td>
<td>32</td>
</tr>
</tbody>
</table>

Values represent calorie-adjusted residuals. For presentation purposes, a constant representing the predicted value for each category for 2,500 kcal for men and 2,000 kcal for women has been added. MeDi = Mediterranean diet; MUFA = monounsaturated fatty acids; SFA = saturated fatty acids.
Table 2
Demographic, Clinical, and Dietary Characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Nondemented (n = 1,964)</th>
<th>Incident AD (n = 262)</th>
<th>All (N = 2,226)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age, yr (SD)</td>
<td>76.5 (6.3)</td>
<td>81.8 (6.9)</td>
<td>77.2 (6.6)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>634 (32)</td>
<td>86 (33)</td>
<td>720 (32)</td>
<td>0.86</td>
</tr>
<tr>
<td>Mean education, yr (SD)</td>
<td>10.5 (4.6)</td>
<td>7.0 (4.5)</td>
<td>10.1 (4.7)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Ethnicity, n (%)</td>
<td></td>
<td></td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>White</td>
<td>592 (30)</td>
<td>28 (11)</td>
<td>620 (28)</td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>636 (32)</td>
<td>86 (33)</td>
<td>722 (32)</td>
<td></td>
</tr>
<tr>
<td>Hispanic</td>
<td>705 (36)</td>
<td>143 (55)</td>
<td>848 (38)</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>31 (2)</td>
<td>5 (2)</td>
<td>36 (2)</td>
<td></td>
</tr>
<tr>
<td>Presence of ε4 allele, n (%)</td>
<td>452 (27)</td>
<td>74 (32)</td>
<td>526 (27)</td>
<td>0.10</td>
</tr>
<tr>
<td>Smoking, n (%)</td>
<td>242 (12)</td>
<td>33 (13)</td>
<td>275 (12)</td>
<td>0.90</td>
</tr>
<tr>
<td>Mean comorbidity index (SD)</td>
<td>1.9 (1.4)</td>
<td>1.9 (1.5)</td>
<td>1.9 (1.4)</td>
<td>0.87</td>
</tr>
<tr>
<td>Mean energy, kcal (SD)</td>
<td>1422 (513)</td>
<td>1475 (560)</td>
<td>1428 (519)</td>
<td>0.15</td>
</tr>
<tr>
<td>Mean body mass index (SD)</td>
<td>27.5 (5.5)</td>
<td>26.5 (5.9)</td>
<td>27.4 (5.5)</td>
<td>0.005</td>
</tr>
<tr>
<td>Mean MeDi score (SD)</td>
<td>4.4 (1.7)</td>
<td>4.2 (1.7)</td>
<td>4.3 (1.7)</td>
<td>0.05</td>
</tr>
<tr>
<td>Low MeDi tertile, n (%)</td>
<td>326 (32)</td>
<td>96 (37)</td>
<td>722 (32)</td>
<td>0.23</td>
</tr>
<tr>
<td>Middle MeDi tertile, n (%)</td>
<td>823 (42)</td>
<td>107 (41)</td>
<td>930 (42)</td>
<td></td>
</tr>
<tr>
<td>High MeDi tertile, n (%)</td>
<td>515 (26)</td>
<td>59 (23)</td>
<td>574 (26)</td>
<td></td>
</tr>
</tbody>
</table>

The sum of nondemented and incident Alzheimer’s disease (AD) is smaller than all subjects combined (ie, n = 2,258; see Fig 1) because 32 subjects who experienced development of dementia other than AD are not included in this table. MeDi = Mediterranean diet; SD = standard deviation.
Table 3
Demographic and Clinical Characteristics by Mediterranean Diet Score Tertiles

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Low Tertile (MeDi score 0–3)</th>
<th>Middle Tertile (MeDi score 4–5)</th>
<th>High Tertile (MeDi score 6–9)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>WHICAP 1999 cohort, N (%)</td>
<td>395 (55)</td>
<td>508 (55)</td>
<td>312 (54)</td>
<td>0.99</td>
</tr>
<tr>
<td>Mean age, yr (SD)</td>
<td>77.3 (6.8)</td>
<td>77.0 (6.6)</td>
<td>77.3 (6.3)</td>
<td>0.50</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>218 (30)</td>
<td>303 (33)</td>
<td>199 (35)</td>
<td>0.23</td>
</tr>
<tr>
<td>Mean education, yr (SD)</td>
<td>10.2 (4.5)</td>
<td>9.9 (4.8)</td>
<td>10.1 (4.9)</td>
<td>0.45</td>
</tr>
<tr>
<td>Ethnicity, n (%)</td>
<td></td>
<td></td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>White</td>
<td>204 (28)</td>
<td>251 (27)</td>
<td>165 (29)</td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>280 (39)</td>
<td>292 (31)</td>
<td>150 (26)</td>
<td></td>
</tr>
<tr>
<td>Hispanic</td>
<td>225 (31)</td>
<td>374 (40)</td>
<td>249 (43)</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>13 (2)</td>
<td>13 (1)</td>
<td>10 (2)</td>
<td></td>
</tr>
<tr>
<td>Presence of ε4 allele, n (%)</td>
<td>164 (27)</td>
<td>214 (27)</td>
<td>148 (29)</td>
<td>0.68</td>
</tr>
<tr>
<td>Smoking, n (%)</td>
<td>111 (15)</td>
<td>119 (13)</td>
<td>45 (8)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Mean comorbidity index (SD)</td>
<td>1.9 (1.4)</td>
<td>1.9 (1.4)</td>
<td>2.0 (1.5)</td>
<td>0.95</td>
</tr>
<tr>
<td>Mean energy, kcal (SD)</td>
<td>1,498 (605)</td>
<td>1,399 (467)</td>
<td>1,387 (473)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Body mass index (SD)</td>
<td>27.5 (5.8)</td>
<td>27.5 (5.5)</td>
<td>27.2 (5.3)</td>
<td>0.55</td>
</tr>
</tbody>
</table>

WHICAP = Washington Heights-Inwood Columbia Aging Project; MeDi = Mediterranean diet; SD = standard deviation.
Table 4

Cox Proportional Hazard Ratios for Alzheimer’s Disease by Mediterranean Diet Score Either in Continuous or in Tertile Form

<table>
<thead>
<tr>
<th>Model</th>
<th>At Risk, n</th>
<th>MeDi Continuous</th>
<th>MeDi Tertiles</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>AD (%) HR (95% CI) p</td>
<td>HR (95% CI) p for Trend</td>
</tr>
<tr>
<td>1a</td>
<td>2,226</td>
<td>262 (12) 0.90 (0.83–0.96) 0.003</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 (reference)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Middle 0.79 (0.60–1.04)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>High 0.61 (0.44–0.85)</td>
<td></td>
</tr>
<tr>
<td>2b</td>
<td>1,759</td>
<td>219 (12) 0.91 (0.83–0.98) 0.015</td>
<td>0.007</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 (reference)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Middle 0.85 (0.63–1.16)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>High 0.60 (0.42–0.87)</td>
<td></td>
</tr>
</tbody>
</table>

The initial number at risk (n = 2226) is smaller than all subjects combined (ie, n = 2258; see Fig 1) because of excluding 32 subjects who experienced development of dementia other than Alzheimer’s disease (AD).

a Model 1 is unadjusted.

b Model 2 is adjusted for cohort, age, sex, ethnicity, education, apolipoprotein E genotype, caloric intake, smoking, comorbidity index, and body mass index.

MeDi = Mediterranean diet; HR = hazard ratio; CI = confidence interval.