Alzheimer’s disease after remote head injury: an incidence study

P W Schofield, M Tang, K Marder, K Bell, G Dooneief, M Chun, M Sano, Y Stern, R Mayeux

Abstract

Objective—To evaluate a history of remote head injury as a risk factor for subsequent dementia due to Alzheimer’s disease.

Methods—271 participants of a community based longitudinal study of aging in north Manhattan without evidence of significant cognitive impairment were interrogated for a history of head injury on two occasions at entry into the study. The examining physician sought a history of head injury with loss of consciousness. Independently, a risk factor interviewer inquired about a history of head injury with loss of consciousness or amnesia, the duration of any loss of consciousness, and the date of the head injury. Patients were followed up with standardised annual evaluations for up to five years to determine the first occurrence of dementia.

Results—Over the course of the study incident dementia due to probable or possible Alzheimer’s disease was diagnosed in 39 patients. Cox proportional hazards modelling showed that a history of head injury with loss of consciousness reported to the physician was associated with earlier onset of dementia due to Alzheimer’s disease (relative risk (RR) = 4:1, 95% confidence interval (95% CI) 1:3–12:7), head injury with loss of consciousness or amnesia reported to the risk factor interviewer was not significantly associated with earlier onset of Alzheimer’s disease overall (RR 2:0, 95% CI 0:7–6:2), but those who reported loss of consciousness exceeding five minutes were at significantly increased risk (RR 11:2, 95% CI 2:3–59:8). Incident Alzheimer’s disease was significantly associated with head injury which occurred within the preceding 30 years (RR 5:4, 95% CI 1:5–19:5).

Conclusion—The results of this cohort study are consistent with the findings of several case-control studies suggesting that head injury may be a risk factor for Alzheimer’s disease.

Methods

Data were obtained from a community based registry of conditions related to aging in north Manhattan. To create this registry, nursing homes, home healthcare agencies, private practitioners, and hospital admission and discharge lists were canvassed to identify service recipients aged 60 years or more who were invited to take a brief cognitive screening examination modified from the comprehensive assessment and referral interview. Almost all subjects entered the study between December 1989 and November 1991. All subjects who screened positive (score ≥ 2) and a randomly selected 26% of all subjects who screened negative (score < 2) were referred to a clinical evaluation team for comprehensive clinical assessments, which were repeated annually. Each annual assessment consisted of a clinical evaluation by a physician (comprising history and general medical and neurological assessments), and a battery of neuropsychological tests administered by a trained tester. The standardised neuropsychological battery consisted of tests of memory (the Buschke selective reminding test), orientation, abstract reasoning, language, and construction, and usually took about an hour to complete. Data from the medical and neuropsychological evaluations, as well as any other laboratory or neuroimaging data that might have been available, were reviewed at a consensus conference which was attended by neurologists directly involved with the clinical evaluation of subjects. At the consensus conference, clinical diagnoses were determined, and a clinical dementia rating score (CDR) was also assigned. Initial and follow up evaluations
were performed and reviewed according to the same procedures. Dementia was diagnosed by means of a strict algorithm which required that patients met threshold criteria on neuropsychological evaluation, had evidence of functional impairment either by history or examination, and that the cognitive problems could not be attributed to an acute confusional state. The neuropsychological criteria for dementia were performance below previously defined cut off scores in two out of three memory domains, and in at least two tests of other cognitive domains.10 Neuropsychological and functional criteria were also used to define a borderline category for subjects with clinically significant cognitive impairment who failed to meet our criteria for dementia. Most subjects in this category had a CDR of 0-5. Some subjects with positive screen scores were judged to be free of significant cognitive impairment on the basis of their neuropsychological test performance. NINCDS criteria were used for the diagnoses of probable and possible Alzheimer’s disease.24

HISTORY OF HEAD INJURY
A history of head injury was sought from subjects on two separate occasions. The physician probed medical history according to a standardised format which included the question: “have you ever had a head injury with loss of consciousness?”. Because subjects were seen in their own homes, clarification and supportive history could be sought from spouses or family members to obtain the best possible information. Independently, a one time risk factor questionnaire was conducted with subjects at entry into the study, by trained technicians usually in person. The risk factor questionnaire included the question: “have you ever had a head injury with loss of consciousness, or amnesia?” Subjects who endorsed this question were asked when the head injury had occurred, and the duration of any loss of consciousness, which was recorded as either: < 5 minutes, 5–29 minutes, 29–59 minutes, 1–24 hours, or > 24 hours. A report of head injury with amnesia but no loss of consciousness, would lead to classification as head injury with loss of consciousness < 5 minutes by the risk factor interviewer, but that same history would be recorded as no head injury with loss of consciousness by the physician. Because the two independently obtained histories provided different information, each with specific advantages, we performed parallel analyses using both categories of history of head injury. A history of head injury with loss of consciousness reported to the physician we refer to as head injury+PHYS, head injury reported to the risk factor interviewer we refer to as head injury+RF. Physicians also routinely inquired about memory complaints (recorded yes or no).

HISTORY OF ALCOHOL
Alcohol misuse increases the risk of head injury.22 Details of past and present alcohol consumption were obtained at the risk factor interview. We defined a history of problem drinking if any of the following had occurred due to alcohol: the subject had argued, blacked out, been violent, been arrested, been charged with drink driving, lost his job, or missed work. The risk factor questionnaire has been found to be reliable on repeated interviews for history of head injury ($k = 0.89$), and for history of alcohol use ($k = 0.54$).5 Apolipoprotein E (APOE) genotype was also known for a subsample of subjects.

ENTRY CRITERIA
We included all registry subjects who had completed the risk factor interview, and who were neither demented at the initial evaluation, nor in the border zone category of cognitive impairment referred to earlier.

STATISTICAL ANALYSES
We used $\chi^2$ or $t$ tests to evaluate the association between a history of head injury and the cognitive screen score, performance on the Buschke selective reminding test at entry to the study, age, education, sex, frequency of memory complaints, or a history of problem drinking. We performed survival analyses using the Cox proportional hazards model13 to assess the age at onset of incident dementia in subjects with and without a history of head injury. The end point chosen for these analyses was the first diagnosis of dementia, or the last visit for those who remained non-demented. Similar Cox analyses were performed to assess the importance of severity of head injury for risk of Alzheimer’s disease. In these analyses subjects were grouped according to severity of head injury, and the group specific risk estimates for incident Alzheimer’s disease were obtained relative to the referent group comprising all subjects without head injury. We assessed the effect of latency of head injury in similar fashion, in which latency was defined as the time between head injury and diagnosis of dementia, or between head injury and final visit for subjects who remained non-demented. Previous studies have found latencies < 10 years,14 or < 30 years15 to be associated with increased risk for Alzheimer’s disease. We therefore classified subjects with previous head injury into groups based on these latencies (< 10 years, ≥ 10 years, and also < 30 years, ≥ 30 years) and undertook Cox analyses to obtain risk estimates for these groups compared with the referent group of subjects without head injury. In all Cox analyses we adjusted for sex and education and stratified by age at entry into the study (age < 70, 70–80, 80+).

Results
Two hundred and seventy one subjects satisfied entry criteria. Subjects were followed up from 0–5 years, with a mean duration of follow up of 20–5 months. Twenty seven subjects reported head injury+RF and 19 subjects reported head injury+PHYS. Sixty one (26%) subjects without any history of head injury were lost to follow up after the initial evaluation as were four (21%) head injury+PHYS
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Table 1 Comparison of risk factor interviewer and physician history of head injury

<table>
<thead>
<tr>
<th>Physician history</th>
<th>History of head injury with LOC or amnesia?</th>
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<tbody>
<tr>
<td>Yes (n)</td>
<td>14 (21)</td>
</tr>
<tr>
<td>No (n)</td>
<td>13 (20)</td>
</tr>
</tbody>
</table>

*Includes three subjects who denied head injury at risk factor interview for whom no response was recorded to head injury question in physicians’ history.

LOC = loss of consciousness; AD = probable or possible Alzheimer’s disease.

and six (22%) head injury+RF subjects. These subjects did not differ for age, education, or baseline total recall compared with those with follow up. Data on head injury were missing from the physician interview for three subjects, all of whom denied head injury at the risk factor interview. To obtain the most conservative estimates of the association between head injury+PHYS and Alzheimer’s disease we assumed all three lacked a history of head injury, and included them in all analyses. A history of head injury was obtained by both the physician and risk factor interviewer from 14 subjects, by the physician only from five subjects, and by the risk factor interviewer only from 13 subjects (table 1). We attempted to contact all 18 subjects for whom the risk factor and physician interviews were incongruent. We succeeded in reinterviewing only six subjects, none of whom had been demented when last formally assessed. A history of head injury with loss of consciousness was confirmed in three, was denied in two, and remained uncertain in one instance. Reclassification of the two subjects who denied head injury strengthened the association between head injury and Alzheimer’s disease and we therefore included all subjects in our analyses according to their initial classification. Table 2 presents subject characteristics. Subjects with head injury+PHYS were younger at entry to the study, but there were no other significant differences between subjects with and without head injury, whether recorded by the physician or the risk factor interviewer.

Table 2 Subject characteristics: 14 subjects are included in both HI+RF and HI+PHYS categories

<table>
<thead>
<tr>
<th>Sex</th>
<th>HI (n = 239)</th>
<th>HI+RF (n = 27)</th>
<th>HI+PHYS (n = 19)</th>
<th>Total (n = 271)</th>
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<tbody>
<tr>
<td>Male</td>
<td>177 (74)</td>
<td>16 (60)</td>
<td>11 (58)</td>
<td>197 (73)</td>
</tr>
<tr>
<td>Female</td>
<td>17 (26)</td>
<td>1 (21)</td>
<td>8 (39)</td>
<td>24 (25)</td>
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<table>
<thead>
<tr>
<th>Age at entry (y) (mean SD)</th>
<th>HI (n = 239)</th>
<th>HI+RF (n = 27)</th>
<th>HI+PHYS (n = 19)</th>
<th>Total (n = 271)</th>
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<tbody>
<tr>
<td>75.6 (7.4)</td>
<td>73.9 (7.2)</td>
<td>71.2 (6.5)*</td>
<td>75.3 (7.3)</td>
<td></td>
</tr>
<tr>
<td>80.0 (4.1)</td>
<td>82.3 (3.8)</td>
<td>88.3 (3.4)</td>
<td>80.4 (4.1)</td>
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<tr>
<td>36.9 (8.3)</td>
<td>37.9 (7.8)</td>
<td>38.0 (8.5)</td>
<td>37.0 (8.3)</td>
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<tr>
<td>2.6 (2.5)</td>
<td>2.5 (2.3)</td>
<td>3.0 (3.0)</td>
<td>2.6 (2.6)</td>
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<tr>
<td>84 (35)</td>
<td>11 (41)</td>
<td>7 (37)</td>
<td>96 (35)</td>
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<tr>
<td>10 (4)</td>
<td>2 (7)</td>
<td>2 (10)</td>
<td>14 (5)</td>
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<tr>
<td>20.7 (15.7)</td>
<td>20.2 (13.8)</td>
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<th>Duration of follow up (months) (mean SD)</th>
<th>HI (n = 239)</th>
<th>HI+RF (n = 27)</th>
<th>HI+PHYS (n = 19)</th>
<th>Total (n = 271)</th>
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<tbody>
<tr>
<td>9 (35)</td>
<td>9 (35)</td>
<td>7 (37)</td>
<td>96 (35)</td>
<td></td>
</tr>
<tr>
<td>10 (4)</td>
<td>2 (7)</td>
<td>2 (10)</td>
<td>14 (5)</td>
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<tr>
<td>20.7 (15.7)</td>
<td>20.2 (13.8)</td>
<td>22.8 (15.5)</td>
<td>20.5 (15.5)</td>
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<th>Any dementia at follow up (n%)</th>
<th>HI (n = 239)</th>
<th>HI+RF (n = 27)</th>
<th>HI+PHYS (n = 19)</th>
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<tr>
<td>34 (14)</td>
<td>4 (15)</td>
<td>4 (21)</td>
<td>39 (14)</td>
<td></td>
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<tr>
<td>82.8 (7.9)</td>
<td>80.9 (9.8)</td>
<td>75.0 (10.0)</td>
<td>82.0 (8.2)</td>
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<thead>
<tr>
<th>Age at AD diagnosis (y)</th>
<th>HI (n = 239)</th>
<th>HI+RF (n = 27)</th>
<th>HI+PHYS (n = 19)</th>
<th>Total (n = 271)</th>
</tr>
</thead>
<tbody>
<tr>
<td>82.8 (7.9)</td>
<td>80.9 (9.8)</td>
<td>75.0 (10.0)</td>
<td>82.0 (8.2)</td>
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INCIDENT DEMENTIA
Incident dementia was found in 48 subjects. Aetiological diagnoses included probable and possible Alzheimer’s disease (n = 39), dementia associated with Parkinson’s disease (n = 2), and other dementias (n = 7). Alzheimer’s disease was diagnosed in three subjects with head injury by both histories, in one subject with head injury by physician’s history alone, and in one subject with head injury by risk factor interview alone (table 1).

In Cox’s proportional hazards model analyses, the RR for incident dementia due to any cause associated with head injury+PHYS was 3.2 (95% CI 1.2-8.6), with head injury+RF risk factor the RR was 2.1 (95% CI 0.8-5.3).

To investigate the association between head injury and Alzheimer’s disease, we excluded the nine subjects with non-Alzheimer’s disease dementia, and repeated the analyses. Head injury+PHYS was associated with significant risk for dementia (RR 4.1, 95% CI 1.3-12.7), but the risk associated with head injury + RF was not significant (RR 2.0, 95% CI 0.7-6.2). To determine if risk differed within strata defined by cognitive screen score, we conducted separate Cox analyses for subjects with cognitive screen score ≤ 2 (low score), and for those who scored > 2 (high score). Similar point estimates of risk associated with head injury+PHYS were obtained within the low score (OR 6.0) and high score (OR 3.9) groups. Risk estimates for Alzheimer’s disease associated with head injury+RF were also similar in the low score (OR 3.6) and high score groups (OR 2.5). Furthermore, Cox models which included the cognitive screen score group (high or low) as a dichotomous independent variable did not alter the risk estimates associated with history of head injury. In another Cox model, covariates included sex; education; the total recall score of the Buscke selective reminding test at baseline as a continuous variable to adjust for initial cognitive performance; memory complaints as a dichotomous variable (yes or no); in addition to history of head injury. In these models the risk estimates associated with head injury+PHYS (RR 3.6, 95% CI 1.2-11.2) or head injury+RF (RR 1.9, 95% CI 0.6-5.8) were similar to previous estimates. Finally, we assessed head injury in this last Cox model redefined as a positive history of head injury to both physician and risk factor interviewer. Head injury defined in this way was also associated with increased risk for incident Alzheimer’s disease, although it missed the level of significance (RR 3.1, 95% CI 0.9-10.8).

SEVERITY OF HEAD INJURY AND LATENCY TO DEMENTIA
Eighteen subjects reported the duration of their loss of consciousness to the risk factor interviewer and this was used as a dichotomised index of severity of head injury: loss of consciousness < five minutes (n = 12), and loss of consciousness ≥ five minutes (n = 6). In Cox analyses, loss of consciousness ≥ five minutes was associated with significantly
increased risk for Alzheimer’s disease (RR 11-2, 95% CI 2.3–59.8); whereas loss of consciousness < five minutes was not (RR 1-7, 95% CI 0.4–7.5).

The mean latency in four subjects with incident Alzheimer’s disease was 14.5 (SD 15.3) years, and in the 14 who reported their age at head injury and remained non-demented, the mean latency was 35.5 (SD 26.2) years. Head injury with latency < 30 years was significantly associated with Alzheimer’s disease (RR 5.4, 95% CI 1.5–19.5), whereas more remote head injury was not (RR 1.7, 95% CI 0.2–14.4). In a separate analysis, head injury with latency < 10 years (RR 4.5, 95% CI 1.0–21.0) and with ≥ 10 years (RR 3.0, 95% CI 0.7–14.2) were both associated with point estimates of increased risk.

We repeated our analyses, adjusting for APOE status, known for only 125 subjects, including 11 head injury+PHYS and 14 head injury+RF subjects. In these analyses, the point estimates of risk associated with head injury+PHYS (OR 3.1, 95% CI 0.8–12.5), and head injury+RF (OR 2.3, 95% CI 0.5–10.0) remained raised. Only one head injury+PHYS and two head injury+RF subjects were APOE ε4+, none of whom became demented.

Discussion

Our results indicate that head injury may be associated with an earlier age at onset of Alzheimer’s disease, and are consistent with the results of several previous case-control studies,14 which suggest that a history of head injury with loss of consciousness may be a significant risk factor for Alzheimer’s disease.

The discordance in the history of head injury obtained by the physician and that obtained by the risk factor interviewer has several possible explanations. Because the risk factor question was more inclusive and led to more reports of head injury, more mild head injury may have been recorded by the risk factor interviewers. In addition, some subjects may have had variable recall of head injury, although the high consistency we previously demonstrated for response to the risk factor for head injury question suggested that it probably occurred very infrequently.5 No independent source of the history of head injury was sought, such as hospital or doctors’ records, because of the associated practical difficulties and the potential bias introduced by excluding injuries which did not have medical attention.

Errors of recall or misclassification of history of head injury would only have been critical for our study if they arose differentially. Although we obtained the history of head injury from subjects who were all initially cognitively intact to avoid the obvious potential for recall bias inherent in case-control studies of Alzheimer’s disease, some potential for recall bias still existed. Subjects with memory complaints—typically for recent rather than distant events and perhaps reflecting the earliest manifestations of preclinical Alzheimer’s disease—might have been more likely to recall a past head injury in an effort to account for their symptoms. However, we found no association between memory complaints and history of head injury, and the association between head injury and subsequent Alzheimer’s disease was not significantly modified by the addition of memory complaints to the Cox model analyses. Thus although there may have been some errors with respect to the history of head injury, we have no evidence to suspect a bias in the frequency of errors that may have been related to the outcome.

In the study by Williams et al,8 821 subjects without evidence of the target illnesses before their head injury, who were older than 40 years, if most recently evaluated, were followed up by medical linkage retrieval over 15,000 person-years for the development of dementia and other degenerative diseases. The standardised morbidity for dementia in this study population, due to dementia and to Alzheimer’s disease specifically, was no different from that in a control population. Latency from head injury to onset of dementia was not significant; nor was there any significant effect of severity of head injury for the risk of subsequent dementia. In a prospective cohort study, Katzman et al followed up 434 volunteers who at intake were “ambulatory, functional, presumably nondemented, and between 75 and 85 years of age.”9 Fifty six subjects became demented, and head injury did not seem to be a risk factor. A prior history of head injury was reported in 31% and 20% of subjects with postintake diagnoses of Alzheimer’s disease and multi-infarct dementia/mixed dementia, and in 10% of those who remained without cognitive impairment at the end of the study. Neither severity nor latency effects were evaluated in that study. There are several methodological differences between these studies and ours, which may in part account for the different results. In the study by Williams et al, dementia was diagnosed by the subjects’ own physicians; subjects with onset of dementia as young as 40 were eligible for inclusion, and standardisation of estimates of dementia onset may have been difficult. By contrast, our subjects were older than 60 at their initial evaluation, (almost all were older than 65). Because they were evaluated annually we were able to establish the age at onset of dementia using uniform diagnostic criteria, based on detailed neuropsychological testing. In the study by Katzman et al, subjects were on average older than our subjects. In addition they were volunteers, by contrast with most of our subjects who were approached with a request to participate in the study.

There are several possible mechanisms by which head injury might increase the risk for Alzheimer’s disease. Head injury might cause cerebral damage and thereby lower the reserve against the cognitive consequences of subsequent, entirely unrelated cerebral pathology.
including Alzheimer’s disease.\textsuperscript{23} There are few human pathological data concerning the cerebral consequences of mild to moderate head injury—necropsy studies are limited to a few reports of individuals who died from causes not directly related to the head injury.\textsuperscript{24, 25} On the other hand, several large follow-up studies of victims with mild head injury indicate that neuropsychological abnormalities may persist for weeks or even months after injury\textsuperscript{26–29} and recovery from the cognitive effects of head injury seems even more delayed in those with a history of previous head injury.\textsuperscript{30} Evoked potential abnormalities are detected in a high proportion of victims of head injury, persisting in some cases for more than six weeks.\textsuperscript{31} These results are consistent with the possibility that mild to moderate head injury causes structural brain changes in some subjects. At least two predictions follow from the “loss of reserve” theory, as far as population studies of head injury and subsequent Alzheimer’s disease are concerned. Head injury should be associated with a younger age at onset of dementia; and more severe head injury should be associated with greater risk for Alzheimer’s disease, as it is likely to be associated with more brain damage. The findings in our study were consistent with both these predictions. Gedye et al also reported that head injury was associated with a younger age of onset of Alzheimer’s disease.\textsuperscript{32} A positive association between increasing severity of head injury and increasing risk for Alzheimer’s disease has been found in a previous study.\textsuperscript{5} A second possible mechanism is that head injury might be particularly damaging when it occurs during the presymptomatic phase of Alzheimer’s disease. The duration of the presymptomatic period in Alzheimer’s disease is unknown, but pathological studies of subjects with Down’s syndrome—in whom Alzheimer’s disease-like changes develop with some predictability—suggest that it might well exceed 10 years.\textsuperscript{33–35} and a recent clinical study indicated that subtle cognitive changes may precede clinical dementia by at least seven years.\textsuperscript{36} The brains of normal animals subjected to mild head injury show characteristic cytoskeletal abnormalities very soon after injury, but there is subsequent repair.\textsuperscript{37–40} If the capacity for repair was diminished in presymptomatic Alzheimer’s disease, even mild head injury might add to the neuropathological burden leading to earlier diagnosis of the underlying Alzheimer’s disease. Conversely, if head injury accelerated the progression of Alzheimer’s disease pathology in those with presymptomatic disease, earlier clinical onset would also arise, consistent with the findings of this study. The APOE ε4 allele may be associated with less effective CNS repair after injury.\textsuperscript{41} The results of a recent case-control study in which head injury was a risk factor for Alzheimer’s disease only among subjects with APOE ε4 would be consistent with the hypothetical mechanisms considered above.\textsuperscript{4} In the current study, too few subjects with APOE ε4 reported head injury for us to seek an interaction between APOE ε4 and head injury, but our results did suggest that the presence of APOE ε4 may not be necessary for the association between head injury and earlier onset of Alzheimer’s disease. A third possible mechanism for the association between head injury and Alzheimer’s disease is that head injury might function as a catalyst, to trigger or promote some critical early event in the pathogenesis of Alzheimer’s disease. Several reported cases of early onset Alzheimer’s disease which presented after head injury suggest such a possibility.\textsuperscript{42–44} Recent studies by Roberts et al indicate that, with severe head injury at least, upregulation of amyloid precursor protein processing may occur.\textsuperscript{45} As aberrant amyloid processing or excessive amyloid production are proposed by some to be central early events in Alzheimer’s disease pathogenesis,\textsuperscript{46} this finding suggests a mechanism by which head injury might play a part in causation of Alzheimer’s disease. There were some limitations to our study. Laboratory evaluations were not routinely available for all subjects, and this may have led to diagnostic errors with respect to the aetiology of the dementia in some cases. Relatively few subjects gave a history of head injury, and the 95% CIs for the association between head injury and Alzheimer’s disease were wide, particularly when we stratified analyses to assess the importance of severity and latency of head injury. Larger prospective studies with greater statistical power will clearly be important to re-evaluate this question. Our study also has strengths. The cohort design significantly reduced the likelihood of recall bias with respect to history of head injury. A history of head injury was sought twice, and we identified incident cases of dementia by uniform criteria derived mainly from the results of comprehensive neuropsychological evaluations.\textsuperscript{1, 2} In our study, estimates of severity of head injury were available, by contrast with many previous case control studies. In summary, our findings offer further support for an association between head injury and Alzheimer’s disease. This association could arise if head injury shortened the long preclinical period of Alzheimer’s disease, or if head injury had a direct role in initiating the disease.
a link between divergent hypotheses? Neurology 1995;45:
7 Mortimer JA, Van Duijn CM, Chandra V, et al. Head trauma as a risk factor for Alzheimer's disease: a collabo-
7 78 tive re-analysis of case-control studies. Int J Epidemiol 
8 Williams DH, Anagnostou JF, Kokmen E, O'Brien PC, 
Furland LT. Brain injury and neurologic sequelae: a 
cohort study of dementia, parkinsonism, and amy-
dementing illnesses in an 80-year-old volunteer cohort. 
neuropsychological paradigm-based diagnosis of dementia and quantified correction for the effects of education. 
12 Gurka B, Wilder D. The "CARE" interview revisited: 
development of an efficient, systematic, clinical assess-
13 Busche H, Fuld PA. Evaluating storage, retention, and 
retrieval in disorders memory and learning. Neurology 
1974;24:1019-25.
14 Benton AL. The visual retention test. New York: The 
Psychological Corporation, 1955.
15 Wechsler D. Wechsler adult intelligence scale-revised. 
16 Mattis S. Mental Status examination for organic mental 
syndrome in the elderly patient. In: Bellak L, Karasu TB, 
ed. Geriatric psychiatry, New York: Grune and Stratton, 
1976.
17 Kaplan E, Goodglass H, Weintraub S. Boston naming test. 
Philadelphia: Lea and Febiger, 1983
18 Ebert H, Kaplan D. The assessment of aphasia and 
related disorders. 2nd ed. Philadelphia: Lea and Febiger, 
1981.
19 Rome W. The Roven drawing test. Bronx, New York: 
Veterans Administration Medical Center, 1981.
20 Burke WJ, Miller JP, Ruben EH, et al. The reliability of the 
Washington University clinical dementia rating. Arch 
21 Ghebremeskel G, Drachman D, Foltstein M, Katzman R, Price 
D, Stadlan EM. Clinical diagnosis of Alzheimer's dis-
ease: report of the NINCDS-ADRDA Work Group 
under the auspices of Department of Health and Human 
Services Task Force on Alzheimer's disease. Neurology 
1984;34:279-44.
22 Galbraith S, Murray WR, Patel AR, et al. The relationship 
between alcohol and head injury and its effect on the 
23 Lawless JF. Statistical model and methods for lifetime data. 
24 Mortimer JA, Pirezzolo JF. Remote effects of head trauma. 
25 Oppenheim DR. Microscopic lesions in the brain follow-
ing head injury. J Neurol Neurosurg Psychiatry 1968;31: 
299-306.
26 Hume Adams J, Doyle D, et al. Diffuse axonal injury in 
head injury: definition, diagnosis and grading. 
27 Gronwall D, Wrightson P. Delayed recovery of intellectual 
generation of small minor head injury. Arch Neurol 
28 Bohnen N, Jolles J, Twijnstra A. Neuropsychological 
deficits in patients with persistent symptoms six months 
29 Barth JT, Maciocio SN, Giordani B, Rimel R, Jane JA, 
Boil TJ. A neuropsychological sequelae of minor head 
30 Gronwall D, Wrightson P. Cumulative effect of concus-
31 Montgomery EA, Fenton GW, Meggitt RJ, MacFynn 
G, Rutherford WH. The psychobiology of minor head 
32 Gudie A, Beattie BL, Tuokko H, Horton A, Korsarek E. 
Severe head injury hastens age of onset of Alzheimer's 
33 Whalley LJ. The dementia of Down's syndrome and its 
relation to sociological studies of Alzheimer's disease. Ann 
34 Chase ME, Tigner R, Smuelle DJ, Liu L. Age-related neu-
ropsychological deficits in Down's syndrome. Biol 
35 Lai F, Williams RS. A prospective study of Alzheimer's 
disease in Down syndrome. Arch Neurol 1989;46: 
849-53.
36 Linn RT, Wolf PA, Bachman DL, et al. The "preclinical 
37 Povlishock JT, Becker DP, Cheng CL, Vaughan GW. 
Azoal change in minor head injury. J Neuropathol Exp 
38 Povlishock JT, Ebr DE, Astruc J. Axonal response to 
traumatic brain injury: reactive axonal change, de-
fermentation, neuroplasticity. J Neurotrauma 1992; 
9:189-200.
39 Povlishock JT. Traumatically induced axonal injury: patho-
genic and pathobiological implications. Brain Pathol 
40 Ebr DE, Povlishock JT. Neuroplasticity following tra-
matic brain injury: a study of GABAergic terminal loss 
and recovery in the cat dorsal lateral vestibular nucleus. 
41 Poisner J. Apolipoprotein E in animal models of CNS 
injury and in Alzheimer's disease. Trends Neurosci 
42 Rudelli R, Strom JO, Welsh PT, Ambler MW. Post-
traumatic premature Alzheimer's disease: neuropatho-
logic findings and pathogenetic considerations. Arch 
Neurol 1982;39:570-5.
43 Cornelli JAN, Brie EJ. Observations on the pathology of 
insidious dementia following head injury. J Ment Sc 
44 Clague H, Coia J. Démence pré-sénile post-traumatique: 
après fracture du crâne: considérations médico-légales. 
45 Roberts GW, Gentleman SM, Lynch A, Murray L, Landon 
M, Graham DJ. Beta amyloid protein deposition in the 
brain after severe head injury: implications for the patho-
genesis of Alzheimer's disease. J Neurol Neurosurg 
Psychiatry 1994;57:419-25.
46 Selkoe DJ. Physiological production of the Beta-amyloid 
protein and the mechanism of Alzheimer's disease. 