Injury markers predict time to dementia in subjects with MCI and amyloid pathology

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ABSTRACT

Objectives: Alzheimer disease (AD) can now be diagnosed in subjects with mild cognitive impairment (MCI) using biomarkers. However, little is known about the rate of decline in those subjects. In this cohort study, we aimed to assess the conversion rate to dementia and identify prognostic markers in subjects with MCI and evidence of amyloid pathology.

Methods: We pooled subjects from the VU University Medical Center Alzheimer Center and the Development of Screening Guidelines and Criteria for Predementia Alzheimer's Disease (DESCRIPA) study. We included subjects with MCI, an abnormal level of β -amyloid₁₋₄₂ (A β ₁₋₄₂) in the CSF, and at least one diagnostic follow-up visit. We assessed the effect of APOE genotype, CSF total tau (t-tau) and tau phosphorylated at threonine 181 (p-tau) and hippocampal volume on time to AD-type dementia using Cox proportional hazards models and on decline on the Mini-Mental State Examination (MMSE) using linear mixed models.

Results: We included 110 subjects with MCI with abnormal CSF A β_{1-42} and a mean MMSE score of 26.3 ± 2.8. During a mean follow-up of 2.2 ± 1.0 (range 0.4–5.0) years, 63 subjects (57%) progressed to AD-type dementia. Abnormal CSF t-tau (hazard ratio [HR] 2.3, 95% confidence interval [CI] 1.1–4.6, p = 0.03) and CSF p-tau (HR 3.5, 95% CI 1.3–9.2, p = 0.01) concentration and hippocampal atrophy (HR 2.5, 95% CI 1.1–5.6, p = 0.02) predicted time to dementia. For subjects with both abnormal t-tau concentration and hippocampal atrophy, HR was 7.3 (95% CI 1.0–55.9, p = 0.06). Furthermore, abnormal CSF t-tau and p-tau concentrations and hippocampal atrophy predicted decline in MMSE score.

Conclusions: In subjects with MCI and evidence of amyloid pathology, the injury markers CSF t-tau and p-tau and hippocampal atrophy can predict further cognitive decline. *Neurology*[®] 2012; 79:1809-1816

GLOSSARY

 $A\beta_{1-42} = \beta$ -amyloid₁₋₄₂; AD = Alzheimer disease; CI = confidence interval; DESCRIPA = Development of Screening Guidelines and Criteria for Predementia Alzheimer's Disease; DSM-IV = Diagnostic and Statistical Manual of Mental Disorders, 4th edition; HR = hazard ratio; MCI = mild cognitive impairment; MMSE = Mini-Mental State Examination; p-tau = tau phosphorylated at threonine 181; t-tau = total tau; TMT = Trail Making Test; VUmc = VU University Medical Center.

Recently, 2 sets of research criteria^{1,2} were established, allowing a diagnosis of Alzheimer disease (AD) in subjects with mild cognitive impairment (MCI) and biomarker evidence of AD pathology. An international working group defined criteria for "prodromal AD" in 2007² and in 2011 the National Institute on Aging and the Alzheimer Association published criteria for "MCI due to AD."¹ However, at this moment, the prognosis of subjects fulfilling these criteria is largely unknown, which limits the use of the criteria in clinical practice. Prognostic markers

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for cognitive decline in subjects with MCI due to AD^1 or prodromal AD^2 are therefore urgently needed.

Subjects can be diagnosed with MCI due to AD¹ or prodromal AD² when they have a clinical diagnosis of MCI and biomarker evidence of either β -amyloid pathology, ADrelated neuronal injury, or both. Abnormal amyloid markers may already be present at the earliest stage of the disease and reach a plateau in a very early stage of the disease and can therefore be useful as an early diagnostic marker.^{3–5} Markers of the subsequent neuronal injury, on the other hand, such as CSF tau and hippocampal atrophy on MRI, may reflect more advanced pathology and might be useful as prognostic markers.^{3–5}

For the present study, we selected subjects with MCI and evidence of amyloid pathology, defined by an abnormal level of β -amyloid₁₋₄₂ ($A\beta_{1-42}$) in the CSF. We hypothesized that the injury markers total tau (t-tau) and tau phosphory-lated at threonine 181 (p-tau)⁶⁻⁸ in CSF and hippocampal atrophy on MRI^{9,10} would be associated with progression to AD-type dementia and cognitive decline.

METHODS Subjects. We selected subjects from the Development of Screening Guidelines and Criteria for Predementia Alzheimer's Disease (DESCRIPA) cohort and the memory clinic of the Alzheimer Center of the VU University Medical Center (VUmc). DESCRIPA is a European multicenter study performed in a memory clinic setting.11 The VUmc was one of the DESCRIPA partners and contributed an additional sample of subjects that were seen outside the DESCRIPA inclusion period. Inclusion criteria were a clinical diagnosis of MCI, an abnormal level of CSF A β_{1-42} , based on a clinically validated cutoff (\leq 550 pg/mL),12 and at least one follow-up diagnosis. Subjects with obvious causes for MCI other than AD, such as alcohol abuse or severe depression, were excluded. In 10 of the participating centers, CSF was collected. Of the subjects enrolled at these centers between 2003 and 2005, 64 subjects fulfilled the inclusion criteria. From the VUmc, 46 additional subjects were included.

Standard protocol approvals, registrations, and patient consents. The medical ethics committee at each center approved the study. All patients provided written informed consent.

Clinical assessment. Diagnosis of MCI was made according to the criteria of Petersen et al.¹³ Raw scores on neuropsychological tests were corrected for age, gender, and educational level in accordance with locally collected or published normative data and are expressed as *z* scores. MCI was defined as a *z* score less than -1.5 SD on any of the following tests: the learning measure or delayed recall of a verbal memory task, Trail Making Test (TMT) part A, TMT part B, verbal fluency, or Rey Figure Copy or equivalent test, as described in more detail previously.^{11,14} Follow-up assessment was performed annually up to 5 years. For subjects from the Alzheimer Center of the VUmc, follow-up was part of regular patient care. Diagnosis of AD-type dementia was made according to the *DSM-IV*¹⁵ and National Institute of Neurological and Communicative Disorders and Stroke–Alzheimer's Disease and Related Disorders Association criteria.¹⁶ Time to dementia was defined as the time between the baseline visit and the date AD-type dementia was diagnosed.

CSF analyses. CSF was collected by lumbar puncture, centrifuged, and stored at -80° C in polypropylene tubes. One sample was thawed twice, but analyses without this sample revealed similar results. CSF A β_{1-42} , t-tau, and p-tau were measured with an InnoTest sandwich ELISA (Innogenetics, Ghent, Belgium) in Gothenburg for the DESCRIPA cohort and in Amsterdam for the VUmc cohort. We corrected for interlaboratory ELISA differences by means of 33 samples that were analyzed at both laboratories and adjusted the VUmc values to those of DESCRIPA using the equating formula: Gothenborg = (SD Gothenborg/SD VUmc) × VUmc + average Gothenborg - [(SD Gothenborg/SD VUmc) × average VUmc].¹⁷

MRI analyses. For the DESCRIPA cohort, subjects were scanned according to the routine MRI protocol at each site. Scanning was performed at 1.0 or 1.5 T and included a 3-dimensional T1-weighted gradient echo sequence with near-isotropic voxels and a fast fluid-attenuated inversion recovery sequence.^{14,18}

Hippocampal volume was measured at the Department of Computing of Imperial College London, using LEAP, a segmentation technique based on atlas registration.¹⁹ We tested whether the MRI field strength influenced the LEAP scores in 348 subjects with MCI from the DESCRIPA cohort. Field strength did not affect the LEAP score (difference of 0.07%, p value = 0.8 after correction for age, gender, educational level, baseline Mini-Mental State Examination [MMSE] score, and follow-up diagnosis), and, therefore, we used data from both field strengths without correction.

MRI data were available in 35 of the 64 subjects (55%) from the DESCRIPA cohort and in 30 of the 46 subjects (65%) from VUmc. Subjects with and without MRI data available did not differ with respect to age, gender, educational level, *APOE* status, CSF markers, or score on the MMSE²⁰ at baseline.

APOE genotyping. DNA was isolated from 10 mL EDTAblood for *APOE* genotyping, using the light cycler *APOE* mutation detection kit (Roche Diagnostics GmbH, Mannheim, Germany).

APOE genotype was determined in 99 subjects (90%). Subjects in whom no APOE status was determined scored higher on the MMSE at baseline (27.7 vs 26.1, p = 0.005). There were no differences with respect to age, gender, educational level, medial temporal lobe atrophy, or CSF markers between subjects with and without APOE data available. Subjects were classified as APOE $\epsilon 4$ -positive when having 1 or 2 APOE $\epsilon 4$ alleles.

Statistical analyses. Analyses were performed with SPSS 18.0 for the Macintosh.

For group comparisons of subjects with and without ADtype dementia at follow-up we used χ^2 tests for categorical variables and Student's *t* tests for continuous variables. Data for the CSF markers were log-transformed to obtain an approximately normal distribution. For further analyses we used dichotomized values of the respective markers. We used clinically validated cutoff points for CSF t-tau (\geq 375 pg/mL) and p-tau (\geq 52 pg/ mL).¹² For hippocampal volume, we used a summed volume of the left and right hippocampus of 5.39 cm³ as the cutoff point.

Table 1	Baseline characteristics according to diagnosis at follow up				
		All subjects	No AD-type dementia at follow-up	AD-type dementia at follow-up	
No.		110	47	63	
Age, y, mean ± SD		70.8 ± 7.7	$\textbf{70.1} \pm \textbf{8.1}$	$\textbf{71.3} \pm \textbf{7.4}$	
Female, n (%)		51 (46)	20 (43)	31 (49)	
Education, y, mean ± SD		10.8 ± 3.5	10.5 ± 3.5	$\textbf{11.1}\pm\textbf{3.4}$	
Follow-up, y,	mean ± SD	$\textbf{2.2} \pm \textbf{1.0}$	$\textbf{2.3} \pm \textbf{1.1}$	2.0 ± 0.9	
APOE ∈4 pos	itive, n (%)ª	61 (62)	23 (54)	38 (68)	
$A\beta_{1-42}$, pg/m	L, mean \pm SD	382 ± 98	$\textbf{369} \pm \textbf{100}$	392 ± 97	
t-tau, pg/mL,	mean ± SD	564 ± 345	421 ± 252	670 ± 368^{b}	
t-tau, abnorn	nal, n (%) ^c	81 (74)	28 (60)	53 (84) ^b	
p-tau, pg/mL	p-tau, pg/mL, mean \pm SD		71 ± 35	$\rm 103\pm54^{b}$	
p-tau, abnorr	p-tau, abnormal, n (%)°		32 (68)	58 (92) ^b	
Hippocampa	Hippocampal volume, cm^3 , mean $\pm SD^d$		5.8 ± 0.8	5.2 ± 0.6^{b}	
Hippocampa	l atrophy, n (%) ^e	35 (54)	8 (31)	27 (69) ^b	
MMSE score	, mean ± SD	26.3 ± 2.8	26.8 ± 2.6	25.9 ± 2.8	
Verbal memo mean ± SD	ory, learning (z score),	-1.5 ± 1.0	-1.4 ± 1.1	-1.6 ± 0.9	
Verbal memo mean ± SD	ory, delayed recall (z score),	-1.6 ± 1.0	-1.3 ± 1.1	-1.9 ± 0.8^{b}	
Verbal fluend	cy (z score), mean ± SD	-0.8 ± 1.1	-0.7 ± 1.3	-1.0 ± 0.9	
TMT part A (2	score), mean \pm SD	-0.8 ± 1.8	-0.7 ± 1.6	-0.9 ± 2.0	
TMT part B (z	score), mean ± SD	-1.1 ± 1.6	-1.0 ± 1.6	-1.2 ± 1.6	
Visuoconstru	uction (z score), mean \pm SD	$\textbf{0.2} \pm \textbf{1.1}$	$\textbf{0.03} \pm \textbf{1.2}$	0.3 ± 1.0	
Abbreviations:	$A\beta_{1-42} = \beta \text{-amyloid}_{1-42}; \beta$	AD = Alzheimer	disease; MMSE	E = Mini-Menta	

Abbreviations: $A\beta_{1-42} = \beta$ -amyloid₁₋₄₂; AD = Alzheimer disease; MMSE = Mini-Mental State Examination; p-tau = tau phosphorylated at threonine 181; t-tau = total tau; TMT = Trail Making Test.

^a APOE genotype was determined in 99 subjects.

 $^{\mathrm{b}}\,p$ < 0.005 compared to no dementia at follow-up.

 $^{\rm c}$ Abnormal values were defined as ${\geq}375$ pg/mL for CSF t-tau and ${\geq}52$ pg/mL for CSF p-tau.

^d Hippocampal volume was determined in 65 subjects.

 $^{\rm e}$ Hippocampal atrophy was defined as a summed volume of left and right hippocampus of $<\!\!5.39\,{\rm cm}^3$

This cutoff point could best differentiate between healthy control subjects and subjects with AD-type dementia in the Alzheimer's Disease Neuroimaging Initiative cohort (S.J.B. Vos, I.A. van Rossum, F. Verhey, et al., unpublished data), based on the Youden index using R.^{21,22} This cutoff point was similar to the cutoff point of 5.34 cm³ that could best predict AD-type dementia in our own dataset.²³

We assessed the effect of *APOE* genotype, CSF levels of t-tau and p-tau, and hippocampal atrophy on time to dementia using Cox proportional hazards with correction for age, gender, education, and MMSE score at baseline. Analyses were performed for each variable alone and with all variables together using a stepforward model to select the variables that could best predict ADtype dementia.

We also assessed the association of CSF t-tau and p-tau and hippocampal volume with the decline in MMSE score. We performed mixed-model analyses with an unstructured covariance structure with correction for age, gender, educational level, and center.²⁴

RESULTS Baseline characteristics. We included 110 subjects with MCI and abnormal CSF $A\beta_{1-42}$. Sub-

jects were 70.8 \pm 7.7 years old (average \pm SD), 46% were female, and 62% had at least one *APOE* $\epsilon 4$ allele. Mean MMSE score was 26.3 \pm 2.8. Baseline characteristics of the subjects are shown in table 1. Two subjects progressed to other types of dementia (one subject with vascular dementia and one subject with Parkinson disease dementia). They were included in the group of subjects who did not progress to AD-type dementia. Excluding those 2 subjects from the analyses did not change the results (data not shown).

Predictors of progression to AD-type dementia. During a mean follow-up of 2.2 \pm 1.0 years (median 2.0 years, range 0.4–5.0 years), 63 subjects (57%) progressed to AD-type dementia. These subjects had higher levels of CSF t-tau (mean \pm SD, 670 \pm 368 vs 421 \pm 252 pg/mL, p < 0.001) and p-tau (103 \pm 54 vs 71 \pm 35 pg/mL, p < 0.001), a smaller hippocampal volume (5.2 \pm 0.6 vs 5.8 \pm 0.8 cm³, p = 0.002), and a lower score on the delayed recall of a verbal memory task (z score -1.9 ± 0.8 vs $-1.3 \pm$ 1.1, p = 0.004) than subjects who did not progress to AD-type dementia (table 1).

Predictors of time to AD-type dementia. Survival analyses using Cox proportional hazards models with correction for age, gender, and education showed that time to dementia was predicted by abnormal CSF t-tau (hazard ratio [HR] 2.3, 95% confidence interval [CI] 1.1-4.6, p = 0.03), abnormal CSF p-tau (HR 3.5, 95% CI 1.3-9.2, p = 0.01), and hippocampal atrophy (HR 2.5, 95% CI 1.1-5.6, p = 0.02) (figure 1, table e-1 on the *Neurology*[®] Web site at www.neurology.org). After correction for baseline MMSE score, results remained essentially the same, with an HR of 2.0 (95% CI 1.0-4.2, p =0.06) for CSF t-tau, 3.1 (95% CI 1.2–8.4, p = 0.03) for CSF p-tau, and 2.2 (1.0–5.0, p = 0.06) for hippocampal atrophy. Of the neuropsychological measures, only delayed recall predicted AD-type dementia (HR 2.1, 95% CI 1.0–4.3, p = 0.05) (table e-1). The APOE $\epsilon 4$ genotype, age, gender, and education did not predict time to dementia (table e-1). Cox multivariate analyses with forward-step selection and biomarkers entered as log-transformed continuous variables selected only CSF p-tau (β 1.2, HR 3.3, 95% CI 1.4–7.5, *p* = 0.005). In the multivariate analysis, we did not find a significant interaction between CSF p-tau or t-tau with hippocampal atrophy (p = 0.8).

MMSE slope analyses. Subjects with abnormal CSF t-tau declined more rapidly on the MMSE, with an annual decline of -1.1, compared with -0.4 for subjects with normal CSF t-tau (table 2). At baseline there were no differences in MMSE score between

subjects with normal and abnormal CSF t-tau (26.5 and 26.3, respectively). For CSF p-tau, results were similar (table 2). Subjects with hippocampal atrophy showed a more rapid decline in MMSE score compared with subjects without hippocampal atrophy (average annual decline -1.2 vs -0.5, p = 0.09) (table 2). At baseline, subjects with hippocampal atrophy had lower MMSE scores than subjects without hippocampal atrophy (25.6 vs 27.0, p = 0.02).

Biomarker subgroup analyses. To investigate the effect of the combination of abnormal CSF t-tau and hippocampal atrophy on progression to AD-type dementia and cognitive decline, we subdivided subjects with both CSF and MRI available (n = 65) into 3 groups, depending on their biomarker status at baseline (figure e-1): 1) normal CSF t-tau and no hippocampal atrophy (n = 9, of whom 1 progressed to AD-type dementia); 2) either abnormal CSF t-tau or hippocampal atrophy (n = 28, of whom 16 progressed to AD-type dementia); and 3) both abnormal CSF t-tau and hippocampal atrophy (n = 28, of whom 22 progressed to AD-type dementia). Compared with subjects with normal CSF t-tau and no hippocampal atrophy, subjects with either abnormal CSF t-tau or hippocampal atrophy had an HR of 5.2 (95% CI 0.7-40.3, p = 0.1) for progression to ADtype dementia. For subjects with both abnormal CSF t-tau and hippocampal atrophy, the HR was 7.3 (95% CI 1.0-55.9, p = 0.06) (table 3).

The annual decline in MMSE score was -0.1 (p value slope = 0.8) for subjects with normal CSF t-tau and no hippocampal atrophy, -0.8 (p = 0.001) for subjects with either abnormal CSF t-tau or hippocampal atrophy, and -1.1 (p < 0.001) for subjects with both abnormal CSF t-tau and hippocampal atrophy (table 3, figure 2). The slopes of decline of subjects with 1 or 2 abnormal markers differed from the slope of subjects with both markers normal, but not from each other. For subjects with only abnormal CSF t-tau (n = 21), the annual decline in MMSE score was -0.6 (-1.0 to -0.2, p = 0.006). For subjects with only hippocampal atrophy, no slope analyses could be performed, because of the small sample size (n = 7).

DISCUSSION In this prospective study of subjects who fulfilled the criteria for MCI due to AD^1 and prodromal AD^2 based on abnormal CSF $A\beta_{1-42}$, we found that during a mean follow-up of 2.2 years 63 subjects (57%) progressed to AD-type dementia. High CSF levels of t-tau and p-tau and hippocampal atrophy predicted progression to dementia and declines in MMSE score.

The overall annual conversion rate to dementia of approximately 20% in this study was higher than the



Red lines indicate the subjects with an abnormal value of each respective marker, defined as CSF total tau (t-tau) \geq 375 pg/mL (A), CSF tau phosphorylated at threonine 181 (p-tau) \geq 52 pg/mL (B), and hippocampal volume <5.39 cm³ (C). Blue lines indicate the subjects with normal values of each marker.

Table 2 Predictors for decline in MMSE score ^a								
	No.	Baseline MMSE	p Value ^b	Slope	p Value ^b			
CSF t-tau								
≥375 pg/mL	81	26.2 (25.1-27.4)	0.6	-1.1 (-1.4 to 0.8)	0.02			
<375 pg/mL	29	26.5 (25.3-27.9)		-0.4 (-0.9 to 0.2)				
CSF p-tau								
≥52 pg/mL	90	26.2 (25.1-27.3)	0.4	-1.1 (-1.3 to 0.8)	0.005			
<52 pg/mL	20	26.7 (25.3-28.2)		-0.04 (-0.7 to 0.6)				
Hippocampal volume								
$< 5.39 \text{cm}^3$	35	25.6 (23.8-27.5)	0.02	-1.2 (-1.5 to 0.8)	0.01			
≥5.39 cm ³	30	27.0 (25.2-28.8)		-0.5 (-0.9 to 0.1)				

Abbreviations: MMSE = Mini-Mental State Examination; p-tau = tau phosphorylated at threonine 181; t-tau = total tau.

^a Baseline MMSE scores and slope values of annual change in MMSE score were estimated using mixed models with correction for age, gender, educational level, and center. Values are estimated assuming that subjects are 50% female, are 70 years of age, and have 11 years of education. Data are means (95% confidence interval).

^b The p value of the difference between subjects with normal and abnormal values for each biomarker.

conversion rate typically observed in subjects with MCI unselected for biomarker status.²⁵ For comparison, subjects with MCI and a normal concentration of CSF $A\beta_{1-42}$ in our dataset had an annual conversion rate of less than 10% (data not shown). Still, a considerable percentage of our subjects did not develop AD-type dementia within the follow-up period. Because abnormal $A\beta$ is suggested to be an early marker for AD,³ higher progression rates to AD-type dementia might be expected with a longer follow-up period.

The rapid decline to dementia in subjects with high CSF levels of t-tau and p-tau and hippocampal atrophy could mean that these subjects either had a more aggressive course of the disease or were already in a more advanced stage when assessed at baseline. Slope analyses suggested that they had a more aggressive course of the disease because they showed a more rapid decline in MMSE score than subjects with normal values of these markers at baseline. This finding is in line with previous studies that showed a more rapid cognitive decline in subjects with AD-type dementia with high levels of CSF tau.^{26,27} Subjects with hippocampal atrophy may have also already been in a more advanced stage of the disease at baseline because they had lower MMSE scores at baseline than subjects without hippocampal atrophy. This result is consistent with the previously suggested order of events in the amyloid cascade,^{3,4} with hippocampal atrophy being a relatively late feature of AD pathology. In a previous study in subjects with MCI and biomarker evidence of A β pathology, hippocampal atrophy also predicted time to dementia.²⁸ In another study in subjects with MCI who all progressed to ADtype dementia, CSF t-tau, CSF p-tau, and hippocampal atrophy were also associated with rapid progression from MCI to AD-type dementia, whereas CSF A β_{1-42} was not.5 Our finding that the predictive value of the respective CSF and MRI markers for progression to AD-type dementia remained after correction for baseline MMSE score indicates that AD biomarkers can have prognostic value in addition to clinical measures alone.

The predictive accuracy of CSF t-tau and p-tau and hippocampal atrophy we observed in our MCI subjects with abnormal CSF $A\beta_{1-42}$ was lower than that reported in studies conducted in subjects with MCI regardless of amyloid biomarker status^{7-9,23} Most likely this is because in our analyses only the additional predictive effect relative to abnormal amyloid was tested, although differences could partly also be due to differences in setting and other study characteristics.

 Table 3
 Progression to AD-type dementia and rate of cognitive decline with respect to biomarker status at baseline^a

CSF t-tau and hippocampal volume ^b	No.	Dementia-free survival after 4 y, mean ± SE	Dementia, HR (95% CI)	Baseline MMSE, HR (95% Cl)⁰	Slope
Both normal	9	$\textbf{0.73} \pm \textbf{0.06}$	Reference	27.4 (25.2-29.6)	-0.1 (-0.9 to 0.7)
One abnormal	28	$\textbf{0.19} \pm \textbf{0.08}$	5.2 (0.7-40.3)	26.6 (24.7-28.5)	-0.8 (-1.2 to 0.4) ^d
Both abnormal	28	0.09 ± 0.03	7.3 (1.0-55.9)	25.5 (23.6-27.4)	-1.1 (-1.5 to 0.7) ^e

Abbreviations: AD = Alzheimer disease; CI = confidence interval; HR = hazard ratio; MMSE = Mini-Mental State Examination; p-tau = tau phosphorylated at threonine 181; t-tau = total tau.

^a Dementia-free survival and the HR were calculated using Cox regression analyses with correction for age, gender, and educational level. Baseline MMSE scores and slope values of annual change in MMSE score were estimated using mixed models with correction for age, gender, educational level, and center. Values are estimated assuming that subjects are 50% female, are 70 years of age, and have 11 years of education.

^b Abnormal CSF t-tau was defined as a value \geq 375 pg/mL; hippocampal atrophy was defined as a volume of both left and right hippocampus of <5.39 cm³.

^c Differences in baseline MMSE between the groups were not statistically significant.

^d The *p* value compared with both markers normal = 0.1.

^e The p value compared with both markers normal = 0.02.

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Slopes of decline in MMSE score in subjects with MCI and abnormal CSF β -amyloid_1-42 (A β_{1-42}) are shown. Subjects were classified according to their CSF t-tau levels and hippocampal volume at baseline. Abnormal values were defined as CSF tau \geq 375 pg/mL and hippocampal volume $<\!5.39\,cm^3$.

We found no differences in age, gender, and *APOE* status between subjects with and without dementia at follow-up, although age, gender, and *APOE* genotype are known risk factors for AD in the general population. A possible explanation for this finding is that advanced age and *APOE* $\epsilon 4$ genotype are risk factors for development of abnormal A β processing but do not influence clinical progression once abnormal A β processing is established.

We included subjects with MCI and abnormal amyloid. According to the criteria of the National

Institute on Aging and the Alzheimer Association,¹ these subjects would meet the criteria for "MCI due to AD-intermediate likelihood." Of the 65 subjects with both CSF and MRI data available, 9 subjects (14%) had both normal CSF t-tau and normal hippocampal volume and met the criteria for "MCI, biomarker evidence uninformative." The course of the disease in these subjects was relatively benign with a 27% conversion rate to AD-type dementia after 4 years, although the interpretation is limited by the small sample size. Twenty-eight subjects (43%) had both abnormal CSF t-tau and hippocampal atrophy and fulfilled the criteria for "MCI due to AD--high likelihood."1 Their prognosis was poor, with 91% progressing to AD-type dementia after 4 years. In 28 subjects (43%), the injury markers were conflicting, with either CSF t-tau abnormal or hippocampal volume abnormal. According to the National Institute on Aging and the Alzheimer Association criteria, it is not clear whether these subjects should be diagnosed as "MCI, biomarker evidence uninformative" or "MCI due to AD-high likelihood."1 Our data suggest that these subjects should be considered as "MCI due to AD-high likelihood" because the decline in MMSE score and progression rate to AD-type dementia (81%) was similar to that of subjects with both markers abnormal, whereas the rate of decline on the MMSE was worse than that of subjects with both markers normal, although group comparisons are hampered by the small sample size.

Two subjects included in the study progressed to other types of dementia, despite abnormal CSF $A\beta_{1-42}$ levels at baseline. One subject, aged 75 years, had extrapyramidal signs at baseline and was later diagnosed with Parkinson disease dementia. CSF $A\beta_{1-42}$ was 326 pg/mL, CSF t-tau and CSF p-tau were normal, and hippocampal volume was not available. Decreased CSF A β_{1-42} has been described before in subjects with alpha-synucleinopathies.²⁹ This finding highlights the importance of ruling out causes for the cognitive symptoms other than AD before the criteria for MCI due to AD can be applied.1 The other subject, aged 61 years, was diagnosed with vascular dementia at follow-up. She had a CSF A β_{1-42} concentration of 357 pg/mL and abnormal CSF t-tau and p-tau concentrations. On the MRI scan she had multiple vascular white matter lesions and parietal atrophy, in the absence of hippocampal atrophy. In retrospect, this subject may have had mixed dementia with both vascular and AD pathology.

A major limitation of our study is that we did not have MRI data available for all subjects, which limited the possibilities for multivariate analyses. Another limitation is the limited follow-up. Studies with longer clinical follow-up are needed to assess whether all subjects with MCI due to AD will indeed develop dementia eventually.

Our results indicate that markers of AD-related neuronal injury, such as CSF levels of t-tau and p-tau and hippocampal atrophy, could help to identify those subjects with MCI due to AD who will more rapidly progress to dementia. Subjects with both abnormal CSF $A\beta_{1-42}$ and abnormal injury markers, thereby fulfilling the criteria for "MCI due to ADhigh likelihood," showed the most rapid cognitive decline and a high progression rate to AD-type dementia, even within our limited follow-up period.

AUTHOR CONTRIBUTIONS

L. Burns, G. L'Italien, and P.J. Visser designed the study. I.A. van Rossum analyzed the data and wrote the manuscript with assistance from P.J. Visser. S.J.B. Vos, P. Scheltens, H. Soininen, L.-O. Wahlund, H. Hampel, M. Tsolaki, L. Minthon, W.M. van der Flier, K. Blennow, F. Barkhof, and F. Verhey were involved in data collection. C.E. Teunissen, K. Blennow, D. Rueckert, and R. Wolz performed biomarker analyses. D.L. Knol helped with statistical methods. All authors reviewed the manuscript and approved the final draft.

DISCLOSURE

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