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Neurodegenerative Biomarkers in Healthy Elderly - with special reference to the preclinical pattern of biological and cognitive markers for Alzheimer's disease

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2009

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Citation for published version (APA):

Stomrud, E. (2009). *Neurodegenerative Biomarkers in Healthy Elderly - with special reference to the preclinical pattern of biological and cognitive markers for Alzheimer's disease*. [Doctoral Thesis (compilation), Clinical Memory Research]. Clinical Memory Research Unit, Lund University.

Total number of authors:

1

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Neurodegenerative Biomarkers in Healthy Elderly

– with special reference to the preclinical pattern of
biological and cognitive markers for Alzheimer's disease

Erik Stomrud, MD

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LUND UNIVERSITY

Doctoral Thesis

With due permission of the Faculty of Medicine at Lund University to be
publicly defended on December 12, 2009 at 9.00 am, in the Main Lecture
Hall at the Clinical Research Centre (CRC), Malmö University Hospital,
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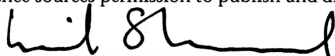
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Malmö 2009

Organization LUND UNIVERSITY Faculty of Medicine Department of Clinical Sciences, Malmö Clinical Memory Research Unit	Document name DOCTORAL DISSERTATION	
	Date of issue December 12th, 2009	
	Sponsoring organization	
Author(s) Erik Stomrud		
Title and subtitle Neurodegenerative biomarkers in healthy elderly - with special reference to the preclinical pattern of biological and cognitive markers for Alzheimer's disease		
Abstract <p>Background: Alzheimer's disease (AD) is a neurodegenerative disorder characterized by tau and amyloid brain pathology. With the gradual degeneration of neurons, cognitive symptoms will arise and the affected individual will eventually develop AD dementia. The neuropathologic hallmarks of AD have been observed also in cognitively healthy individuals, which has led to the assumption that the disorder has a long preclinical phase. Several biological markers for detecting and predicting AD have been validated over the years, where CSF biomarkers are one of the most recent and accurate markers. The generally perceived notion today is that these biological markers will be altered also in the preclinical phase. An additional aspect is that a review made in this thesis of the control samples in CSF articles, suggests that controls have been selected without efforts to minimize the misclassification of preclinical AD individuals. Therefore the aim of this study was to investigate biological and cognitive markers for AD in a group of cognitively healthy elderly individuals who were used as clinical control subjects in research studies.</p> <p>Setting: The study sample consisted of 62 cognitively healthy elderly individuals from a clinical control group. They were followed for 4.5 years at three occasions and underwent assessments of EEG activity and regional cerebral blood flow (rCBF) as well as repeated assessments of cognitive function and CSF biomarker levels. The CSF biomarkers were Aβ42, total tau (t-tau) and hyperphosphorylated tau (p-tau). The cognitive testing included among others the MMSE, the ADAS-cog, cognitive speed (AQT), and subjective memory impairment.</p> <p>Results: In the sample there were individuals with clinically pathological assessments on each separate biological marker. CSF Aβ42 levels predicted development of subjective memory impairment affecting quality of life at the 3 years follow-up and correlated with delayed word recall and cognitive speed at the 4.5 years follow-up. Additionally, the individuals with decreasing CSF Aβ42 levels during the follow-up performed cognitively worse than those with stable levels at the 4.5 years follow-up. CSF tau levels on the other hand correlated with an increase of the low-frequency theta activity on EEG and showed covariance with rCBF in the right medial frontal lobe and the left fronto-parieto-temporal area. In each case the correlations were stronger for p-tau levels compared to t-tau levels. Increase in theta activity was also correlated with slower cognitive speed.</p> <p>Discussion: In this group of cognitively healthy elderly individuals there were individuals with deteriorated cognitive and biological markers associated with AD. These markers further correlated to one another in specific patterns, where it was the known AD-associated changes of the markers (i.e. low CSF Aβ42, high CSF t-tau and p-tau, decreased EEG rhythm, and decreased rCBF) that were primarily related. The findings could imply that the biomarkers might indicate early neurodegenerative changes of the brain and that these changes could be detectable before extensive cognitive impairment. The findings could also suggest that preclinical AD might be present in this "healthy" study sample. Hence, pathological processes prevailing in AD might bridge the clinically created arbitrary division of normal and non-normal aging of the brain.</p>		
Key words: Alzheimer's disease, dementia, MCI, preclinical Alzheimer's disease, biomarkers, cerebrospinal fluid, amyloid beta protein, tau protein, phosphorylated tau protein, EEG, SPECT, cerebral blood flow, cognition, memory, early diagnosis, control groups.		
Classification system and/or index terms (if any):		
Supplementary bibliographical information:		Language English
ISSN and key title: 1652-8220, Doctoral Dissertation Series 2009:116		ISBN 978-91-86443-05-4
Recipient's notes	Number of pages 180	Price
	Security classification	

Distribution by (name and address)

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Malmö 2009

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ISSN 1652-8220

ISBN 978-91-86443-05-4

Lund University, Faculty of Medicine Doctoral Dissertation Series 2009:116

Layout and printed in Malmö, Sweden by Medicinsk Informationsteknik, 2009

“The important thing in science is not so much to obtain new facts as to discover new ways of thinking about them”

Sir William Bragg

ABSTRACT

ENGLISH

Neurodegenerative Biomarkers in Healthy Elderly

– with special reference to the preclinical pattern of biological and cognitive markers for Alzheimer’s disease

Erik Stomrud, MD

Background: Alzheimer’s disease (AD) is a neurodegenerative disorder characterized by tau and amyloid brain pathology. With the gradual degeneration of neurons, cognitive symptoms will arise and the affected individual will eventually develop AD dementia. The neuropathologic hallmarks of AD have been observed also in cognitively healthy individuals, which has led to the assumption that the disorder has a long preclinical phase. Several biological markers for detecting and predicting AD have been validated over the years, where CSF biomarkers are one of the most recent and accurate markers. The generally perceived notion today is that these biological markers will be altered also in the preclinical phase. An additional aspect is that a review made in this thesis of the control samples in CSF articles, suggests that controls have been selected without efforts to minimize the misclassification of preclinical AD individuals. Therefore the aim of this study was to investigate biological and cognitive markers for AD in a group of cognitively healthy elderly individuals who were used as clinical control subjects in research studies.

Setting: The study sample consisted of 62 cognitively healthy elderly individuals from a clinical control group. They were followed for 4.5 years at three occasions and underwent assessments of EEG activity and regional cerebral blood flow (rCBF) as well as repeated assessments of cognitive function and CSF biomarker levels. The CSF biomarkers were A β 42, total tau (t-tau) and hyperphosphorylated tau (p-tau). The cognitive testing included among others the MMSE, the ADAS-cog, cognitive speed (AQT), and subjective memory impairment.

Results: In the sample there were individuals with clinically pathological assessments on each separate biological marker. CSF A β 42 levels predicted development of subjective memory impairment affecting quality of life at the 3-year follow-up and correlated with delayed word recall and cognitive speed at the 4.5-year follow-up. Additionally, the individuals with decreasing CSF A β 42 levels during the follow-up performed cognitively worse than those with stable levels at the 4.5-year follow-up. CSF tau levels on the other hand correlated with an increase of the low-frequency theta activity on EEG and showed covariance

with rCBF in the right medial frontal lobe and the left fronto-parieto-temporal area. In each case the correlations were stronger for p-tau levels compared to t-tau levels. Increase in theta activity was also correlated with slower cognitive speed.

Discussion: In this group of cognitively healthy elderly individuals there were individuals with deteriorated cognitive and biological markers associated with AD. These markers further correlated to one another in specific patterns, where it was the known AD-associated changes of the markers (i.e. low CSF A β 42, high CSF t-tau and p-tau, decreased EEG rhythm, and decreased rCBF) that were primarily related. The findings could imply that the biomarkers might indicate early neurodegenerative changes of the brain and that these changes could be detectable before extensive cognitive impairment. The findings could also suggest that preclinical AD might be present in this “healthy” study sample. Hence, pathological processes prevailing in AD might bridge the clinically created arbitrary division of normal and non-normal aging of the brain.

Neurodegenerativa biomarkörer hos friska äldre

– med fokus på tidiga förändringar av biologiska och kognitiva markörer vid Alzheimers sjukdom.

Erik Stomrud

Alzheimers sjukdom är den vanligaste demenssjukdomen och cirka 14 miljoner personer världen över lever med Alzheimers demens. En person som insjuknat i en demenssjukdom får problem med minnet och/eller andra intellektuella funktioner som leder till svårigheter att klara av vardagen. Vid Alzheimers sjukdom beror detta på att de två proteinerna amyloid och tau omsätts felaktigt och inlagras i hjärnan i form av senila plack och tangles, vilka kan ses vid mikroskopisk undersökning av hjärnvävnaden. Denna felaktiga omsättning leder till att nervcellerna i hjärnan fungerar sämre och att de till slut bryts ned med förlust av hjärnvävnad som följd. Mycket talar dessutom för att dessa processer vid Alzheimers sjukdom startar flera decennier innan den drabbade personen får problem med minnet.

Det finns idag flera undersökningar och tester för att påvisa dessa sjukdomsprocesser. En av de markörer som har visat sig vara bäst på att urskilja personer med Alzheimers demens från friska individer är analys av ryggvätska. Här mäts mängden av en speciell typ av proteinet amyloid (A β 42) samt totala mängden av proteinet tau (t-tau) och mängden hyperfosforylerat tau (p-tau). Utöver detta kan vid Alzheimers sjukdom förändringar ses även av den elektriska aktiviteten i hjärnan (EEG-undersökning) och av blodflödet i olika delar av hjärnan (SPECT-undersökning). Eftersom sjukdomsprocesserna vid Alzheimers sjukdom tros föregå försämrat minne, har det spekulerats i att även dessa undersökningar skulle kunna påvisa sjukdomstecken innan minnet påverkas.

I avhandlingen redovisas en sammanställning av urvalsprocessen för de friska kontroller som har använts inom forskning om ryggvätskemarkörer. Sammanställningen visar att individerna väljs enbart utifrån minnestester och att i en tredjedel av fallen har individerna haft subjektiva problem med minnet. Eftersom sjukdomsprocesserna troligen börjar långt innan minnesproblemen visar sig finns det därmed en stor risk att individer med Alzheimers sjukdom i dessa studier bedömts som friska äldre, vilket kan påverka resultaten i studierna. Därför valde jag i denna avhandling att i en grupp intellektuellt friska äldre individer studera biologiska och intellektuella markörer som i tidigare studier visat samband med Alzheimers sjukdom.

I studien följdes 62 stycken intellektuellt friska äldre individer med olika tester under 4,5 år. De genomgick upprepade minnestester, ryggvätskeprov vid två tillfällen, en EEG-undersökning och en SPECT-undersökning. Flera intellektuella funktioner testades, däribland närminne, intellektuell snabbhet och subjektiva minnesproblem. Ryggvätskeprovet innebar att ryggvätska tappades ut från ländryggen och att nivåerna av proteinerna A β 42, t-tau och p-tau mättes.

I studien sågs att det för varje specifikt test av de biologiska markörerna fanns individer med värden som i en klinisk vardag skulle ha bedömts som avvikande. Vidare sågs i gruppen att de undersökta markörerna var relaterade till varandra i specifika mönster. Lägre nivåer av proteinet A β 42 i ryggvätskan kunde förutsäga sannolikheten att utveckla subjektiva minnesproblem efter 3 år men hade också ett samband med sämre resultat på minnestester avseende närminne och intellektuell snabbhet vid uppföljningen efter 4,5 år. Höga nivåer av t-tau och p-tau hade däremot ett samband med förlängsamrad elektrisk aktivitet framför allt i bakre delen av hjärnan men även med sänkt blodflöde i den främre högra hjärnhalvan och ökat blodflöde i den bakre vänstra hjärnhalvan. Förlängsamrad elektrisk aktivitet var också relaterat till sänkt intellektuell snabbhet.

Studien visade således samband mellan de undersökta markörerna i en grupp intellektuella friska äldre individer. Vid en närmare genomgång kunde detta samband ses just mellan de förändringar som i tidigare studier visat samband med Alzheimers sjukdom. Resultaten skulle kunna tala för att sjukdomsprocesserna vid Alzheimers sjukdom även finns hos friska äldre samt att de i studien undersökta markörerna möjligen skulle kunna påvisa dessa processer. För att säkert kunna fastställa detta behöver emellertid resultaten upprepas i större studier med längre uppföljningstid. Sammanfattningsvis talar resultaten i denna studie för att sjukdomsprocesserna vid Alzheimers sjukdom överskrider den tidigare symtombaserade, kliniska uppdelningen i vad som är ett friskt respektive ett avvikande åldrande av hjärnan.

LIST OF ORIGINAL PUBLICATIONS

I

Stomrud E, Hansson O, Blennow K, Minthon L, Londos E.
Cerebrospinal Fluid Biomarkers Predict Decline in Subjective Cognitive Function over 3 Years in Healthy Elderly.
Dementia and Geriatric Cognitive Disorders 2007; 24: 118–124

II

Stomrud E, Hansson O, Minthon L, Blennow K, Rosén I, Londos E.
Slowing of EEG Correlates with CSF Biomarkers and Reduced Cognitive Speed in Elderly with Normal Cognition over 4 Years.
Neurobiology of Aging xxx (2008) xxx–xxx;
doi:10.1016/j.neurobiolaging.2008.03.025

III

Stomrud E, Hansson O, Zetterberg H, Blennow K, Minthon L, Londos E.
Longitudinal CSF Biomarkers Correlate with Cognitive Decline in Healthy Elderly.
Archives of Neurology 2009; accepted

IV

Stomrud E, Forsberg A, Hägerström D, Ryding E, Blennow K, Zetterberg H, Minthon L, Hansson O, Londos E
CSF Biomarkers Correlate with Regional Cerebral Blood Flow on SPECT in Healthy Elderly.
Submitted for publication

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ABBREVIATIONS AND DEFINITIONS

ABBREVIATIONS

A β	Amyloid beta
A β 42	Amyloid beta 1-42
AD	Alzheimer's disease
ADAS-cog	Alzheimer's Disease Assessment Scale - cognitive subscale
ADL	Activities of daily living
APOE	Gene coding for apolipoprotein E
APOE- ϵ 4 allele	Allele coding for isoform apolipoprotein E4
APP	Amyloid precursor protein
AQT	A Quick Test on cognitive speed
CDR	Clinical dementia rating
CDT	Clock drawing test
CERAD	The Consortium to Establish a Registry for Alzheimer's Disease
CSF	Cerebrospinal fluid
CT	Computed tomography
DLB	Dementia with Lewy bodies
DSM-IV	Diagnostic and Statistical Manual of Mental Disorders, 4 th edition
EC	Entorhinal cortex
FTD	Frontotemporal dementia
HMPAO	Hexamethylpropylene amine oxime
ICD-10	International Classification of Diseases, 10 th revision, by the World Health Organisation
ICF	International Classification of Functioning, Disability and Health, by the World Health Organisation
MCI	Mild cognitive impairment
MemQoL	Quality of life assessment subscale memory

MMSE	Mini Mental State Examination
MRI	Magnetic resonance imaging
MTL	Medial temporal lobe
NFT	Neurofibrillary tangles
NIA-Reagan	National Institute on Aging and the Reagan Institute
NINCDS-ADRDA	National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer's Disease and Related Disorders Association
PET	Positron emission tomography
Pre-MCI	Stage preceding mild cognitive impairment
P-tau	Hyperphosphorylated tau protein
rCBF	Regional cerebral blood flow
SBU	The Swedish Council on Technology Assessment in Health Care
SPECT	Single photon emission computed tomography
SPM	Statistical Parametric Mapping, software program by The Wellcome Trust Centre for Neuroimaging, London, UK
T-tau	Total tau protein
^{99m}Tc	Metastable nuclear isomer of technetium-99
qEEG	Quantitative electroencephalography
WHO	The World Health Association

DEFINITIONS

Alzheimer's disease	Presence of neuropathology changes associated with post-mortem definition of the disease.
AD dementia	Fulfilment of criteria for dementia with concurrent explanatory presence of AD neuropathology changes or fulfilment of clinical criteria of AD
Prodromal AD	Presence of substantial AD neuropathology changes and cognitive symptoms however not enough to yield a dementia diagnosis
Preclinical AD	From the first AD neuropathology changes up to the first appearance of measurable cognitive symptoms

THESIS AT A GLANCE

	Hypothesis	Setting	Results	Conclusion
I-IV	In a group of cognitively healthy elderly individuals there will exist individuals with deteriorated biologic markers for Alzheimer's disease, and these will be associated with proxy markers for future dementia, such as worse cognitive performance.	Repeated cognitive function, repeated CSF biomarkers, EEG activity, and regional cerebral blood flow were assessed in 62 cognitively healthy elderly individuals over a period of 4.5 years.	CSF A β 42 levels correlated with cognitive performance whereas CSF tau levels correlated with EEG activity and rCBF change. The combination of CSF A β 42 and p-tau levels predicted cognitive decline and correlated with EEG activity and cognitive speed.	In a group of cognitively healthy elderly individuals there existed individuals with deteriorated levels of AD-associated biological markers and these could in part be associated with worse cognitive performance. <i>Hypothesis: true & false</i>
I	AD-associated changes in the CSF biomarkers A β 42, t-tau and p-tau correlate with and precede cognitive decline in a group of cognitively healthy elderly individuals.	Baseline CSF biomarker level analysis in 54 cognitively healthy elderly with cognitive follow-up after 3 years. <i>Study design: prospective, longitudinal observation study</i>	Baseline CSF A β 42 levels and the combination of high CSF p-tau and low CSF A β 42 levels could predict development of subjective memory impairment affecting quality of life after 3 years with reasonable accuracy.	CSF biomarkers are related to future cognitive impairment before its onset and imply that AD-related biological signs might be detectable already in preclinical disease stages. <i>Hypothesis: true</i>

	Hypothesis	Setting	Results	Conclusion
II	AD-associated changes in the CSF biomarkers A β 42, t-tau and p-tau are related to a concurrent change in the brain electric activity in a group of cognitively healthy elderly individuals.	Assessment of CSF biomarkers, quantitative EEG activity and cognitive performance in 33 cognitively healthy elderly individuals.	Increased CSF tau levels and tau/A β 42 ratio correlated with increased relative theta activity furthestmost in the right posterior part of the brain. These changes in CSF biomarkers and theta activity was further associated with slowing of cognitive speed.	Neuropathologic processes represented by CSF biomarkers correlate with cerebral function visualized by EEG rhythm and cognitive speed in cognitively unimpaired individuals. The biomarkers in this study might indicate early abnormal degenerative changes in the brain. <i>Hypothesis: true</i>
III	Individuals with a progressive deterioration towards an AD pattern in the CSF biomarkers A β 42, t-tau and p-tau levels will perform cognitively worse than those with stable levels.	Change in CSF biomarker levels over 4.5 years in relation to cognitive performance at 4.5 years follow-up in 37 cognitively healthy elderly individuals. <i>Study design: cross-sectional</i>	Both low CSF A β 42 levels and substantial longitudinal decrease in CSF A β 42 levels were associated with worse delayed episodic memory and cognitive speed. Substantial longitudinal increase in CSF p-tau levels was additionally associated with worse cognitive speed.	In these cognitively unimpaired individuals a concurrent decrease in CSF A β 42 levels and cognitive performance might be observed, which would implicate that CSF biomarker levels might reflect very early neurodegenerative processes in the brain. <i>Hypothesis: true & false</i>
IV	AD-associated changes in the CSF biomarkers A β 42, t-tau and p-tau are related to a concurrent change in regional cerebral blood flow (rCBF) in a group of cognitively healthy elderly individuals.	Assessment of CSF biomarkers, rCBF, and cognitive performance in 32 cognitively healthy elderly individuals <i>Study design: longitudinal</i>	Increased CSF p-tau and t-tau levels were associated with decreased rCBF in the right medial frontal lobe and increased rCBF in the left fronto-parieto-temporal area.	In cognitively unimpaired individuals AD-related biochemical processes represented by CSF biomarkers correlates with AD-related cerebral function visualized by rCBF. Hence, preclinical degenerative changes in the brain might be present as indicated by these biomarkers. <i>Hypothesis: true & false</i>

INTRODUCTION 1

BACKGROUND 1|1

Imagine a train out of control rushing towards you down a railway track. This is how I have chosen to visualize development of Alzheimer's disease (AD) throughout my doctoral-studies. Everything starts at the point when the stationary train is set in motion. In this metaphor this event represents the onset of the first pathologic changes in the brain. At first no one notices that the train is moving with increasing speed. Not until it reaches the place where you are standing do you become aware of it. The time that has elapsed up until this moment represents the preclinical stage of the disease where the pathology can be extensive but cognitive symptoms limited. Later, as the symptoms progress, they ultimately reach a threshold where they become evident. To continue the train journey, you try to board the train but the speed is very high and you have to run along beside it for quite a while. Finally, you manage to throw yourself on board only to realize that the end of the track is in view and a crash will soon be unavoidable. You look for the emergency brake, but find that the train is not equipped with one. The only thing left to do is to jump off the train and stand on the side to see it crash. Returning to AD, health care providers try to investigate and diagnose the disease while the pathology continues to progress in the individual. Once we "boarded" the disease and diagnosed it we discover that we can neither cure it nor slow it down. The only thing we can do is to give proper care and symptomatic treatment as we see the individual deteriorate until the time of a premature death.

In order to change the series of events for the rushing train several actions could be taken. The first would be to make it impossible for the train to be set in motion to begin with. The second would be to install an emergency brake and the third would be to notice the rushing train earlier, especially if there was an emergency brake on board. Again, translated to AD the two first actions would mean the development of a preventive and disease-modifying or -arresting treatment. This is currently ongoing with several potential substances being tested. The third possible action would be to diagnose the disease earlier, already in the preclinical stages, before the point where extensive irreversible brain damage has occurred. Only then preventive treatment will be of value. Hence, increased knowledge of preclinical AD will be imperative if the new treatment strategies prove to be effective. I hope that this study will contribute to the acquirement of new knowledge and a better understanding of this preclinical stage in AD development.

RESEARCH APPROACH 1|2

PRE-EXISTING UNDERSTANDING 1|2|1

A central point of departure has been the opinion that the symptomatic, clinical definition of dementia and AD dementia in particular^{1,2} is not enough to understand the disorders as they present in the everyday contact with patients. This sense of inadequacy has further increased as disease-modifying or -arresting treatments are under development that will alter the underlying neuropathologic changes in the brain. Consequently, focus needs to shift from treating AD solely as a symptomatic condition towards treating it as a biological condition. This is further reinforced with the revised NINCDS-ADRDA research criteria where markers of biological changes have strengthened their position as proxy markers for the post-mortem histologic definition of the disease that is suggested as the gold standard.³ In this biologically focused perspective, it is presumed that a biological sign that differs by more than two standard deviations from a validated, arbitrary reference value is most likely an indication of pathology and disease. It is this pre-existing understanding from which the scientific questions, hypotheses, aims and study design in this thesis have been formulated and hence determine the conclusions this study can provide.

ISSUES THE STUDY CAN ANSWER 1|2|2

22 There are methodological restrictions to the scientific questions that can be addressed in this study. The study is designed in such a way that it may indicate associations between the investigated variables, which could then be discussed in the light of neurodegenerative disease. However, the study is relatively underpowered, which could lead to type-II bias. Hence, absence of associations between the investigated variables in the study does not conclude that the variables are not related. Therefore, the study is limited in its ability to extract conclusions from negative findings and this will therefore be avoided in the thesis. If the restrictions are interpreted for this specific study it would suggest that: associations between markers for AD could indicate shared underlying pathology. However, claims whether the inter-individual spread of these markers is due to natural variation or disease cannot be made. Similarly, the outcome measurements available in the study prohibit attempts to estimate diagnostic validity of the markers for detecting incipient AD.

The sample in this study consists of cognitively healthy elderly individuals. Thus, the variance in cognitive performance and the incidence of longitudinal cognitive decline is limited. In order to overcome the limitation in available outcome measurements, only markers for AD have been investigated, which have been extensively validated for the manifest disease. If both experimental biomarkers and outcome measurements with little variance would have been used, then the ability to find true associations would be unacceptably low and attempts to draw conclusions would be practically impossible. Furthermore, the study has not been designed in a way that enables it to make statements regarding other dementia disorders. This could be a confounding factor since other causes of dementia might influence the cognitive performance variables.

One of the reasons behind the limited sample size, leading to absence in power, has been that invasive investigations have been performed without clinical indication. However, in dementia research these investigations are seldom performed on cognitively healthy controls, which will be shown in the CSF article analysis below. An even more unique element in dementia research is invasive investigations performed repeatedly on the same healthy individual. Nevertheless, one of the strengths of the current study is the sample size, albeit the power limitations. The fact that invasive and brain imaging radiating investigations have been performed on the same individuals additionally strengthens the novelty of this study.

Instead of providing the “true” preclinical pattern of the investigated markers for AD, the purpose of this study has been to explore the preclinical pattern of these AD-associated biological markers in a group of cognitively healthy elderly individuals. The attempts to explore this essentially unknown area of research must be regarded as initial steps that will need to be reproduced with larger and better powered studies. However, the current study could highlight those preclinical relationships that merit further investigation.

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CONTRIBUTION OF THE AUTHOR 1|2|3

As a physician I am inclined to adopt an interest in exploring the biological perspective of AD. Additionally, I have a special interest in the clinical situation where the patient meets the professionals in the health care setting, which also contribute to the direction of this study. Therefore, the scientific questions of this study originate from the clinical questions that arise in these meetings. Hence, this study does not explore the actual techniques for investigating AD markers or the mechanisms behind the changes of the markers. Instead, the

point of departure is the desire to achieve a deeper understanding for how to interpret these markers preclinically and how to balance between them if they are in conflict. This is the same considerations that are made in the patient – physician relationship in order to identify the cause of the patient’s problem.

THEORETICAL POSITIONS 1|3

The scientific questions in this study can never be treated as isolated phenomena, separate from the rest of the AD research field. In fact there are adjacent scientific issues within which the study acts, that the researcher and the readers should be made aware of in order to appreciate the relevance of the study and its findings.

PATHOLOGY VERSUS MANIFESTATION 1|3|1

24 The first issue is whether to treat AD as a disorder of biological changes or of cognitive manifestation. This question is part of the more general discussion of symptoms versus aetiology. The first aspect of this issue is the WHO classification of functioning, disability and health (IFC),⁴ in which a distinction can be made between body structure and function, impairment and disability (Figure 1). Body structure and function is, in this case, a biological term of physiological or anatomical properties of the body. Impairments refer to a deviation from a generally accepted standard in the individual capacity. It is not the same as the underlying pathology but instead a manifestation of this pathology. Hence, limping could be the effect of both congenital damage and of an adult-life injury. Finally, disability could be defined as the individual’s interaction with the surrounding world. Impairment hereby does not have to lead to a disability since both the individual’s behaviour as well as adaptation of the surrounding can compensate for the impairment. In the case of AD neuropathologic features and reduced cerebral blood flow are examples of body structure and function, whereas decline in cognitive performance is an example of impairment and finally, problems with ADL is an example of disability (Figure 1). This discussion is the core issue in the discussion of whether dementia is a disorder defined by the cognitive symptoms (impairments) or neuropathologic changes (body structure and function). Even though the relationships between the positions are not always conclusive, they are by definition not in opposition to one another. Instead they have different importance in different situations, which can be very evident in clinical practice. Therefore, the need to differentiate between symptoms and body changes, manifestations and pathology, clinical description and

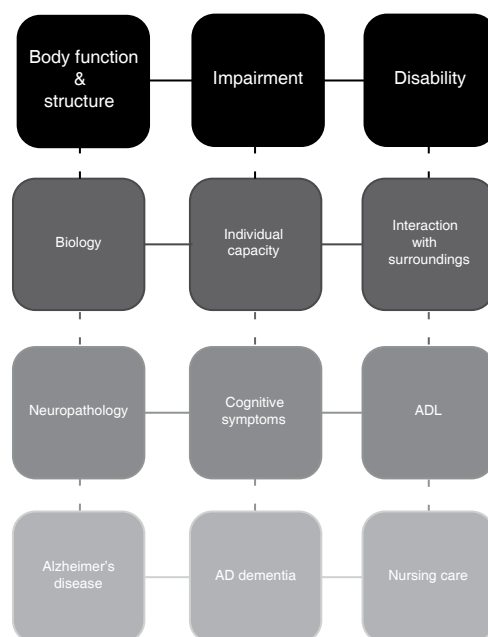


Figure 1. The International Classification of Functioning, Disability and Health (ICF) by WHO (top line) and how it can be interpreted for disease in general (second line), dementia disorders (third line), and Alzheimer's disease in particular (fourth line).

aetiology, in dementia diagnostics has been stressed by many for a long time.^{3, 5-11} The direct impact this has on the current study is that I refer to the biological body changes as Alzheimer's disease (AD) and to the clinical descriptive, manifestation of pathology as AD dementia. In summary, the terms dementia and AD imply more than solely if an individual has a disease or not. The linguistic and declarative reality is quite simply more complex and it is in this reality that the current study tries to navigate.

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SEQUENCE OF CHANGE 1|3|2

The second issue, which is derived from the prior one, is the importance of determining in which order changes in markers occur. In general, changes in body structure and function (pathology) can be assumed to precede impairments (manifestation of pathology). In AD it has further been suggested that change in functions precede change in structure.¹² Hence, the neuron will malfunction before it deteriorates and contributes to the progressive cell loss. With a logical step backward it would be reasonable to assume that the intracellular pathological process in its turn precedes malfunction of the neuron. Consequently, a logi-

cally derived rationale can be proposed for the probable internal order in which different investigations and biological markers for AD could be suspected to be sensitive in reflecting the disease.

NORMALITY 1|3|3

The third and final issue concerns the concept of normality. An exhaustive discussion covering the entire spectra of normality will not be possible in this thesis, since it spreads over several fields of sciences. However, in the light of the current study a number of aspects should be emphasized. Firstly, the word “normal” has different meanings depending on the circumstance in which it is used. For instance, in a biological setting, normal would be used to represent normal distribution, i.e. the Gaussian distribution, which is the probability distribution of a variable that tends to cluster around an average. In clinical practice, in the health care setting, normal would instead represent the values of a variable that lies within an arbitrary reference range (often set to cover 95 % of expected values in a representative population). In a philosophical setting it could be speculated that normal represents a preset or unspoken reference state, which is regarded as the desirable state and which the majority of individuals strive to attain. In the Dorland’s Illustrated Medical Dictionary the word normal is defined as “*agreeing with the regular and established type*” and in the Swedish Academy Glossary it is defined in part as “*an example that constitutes guiding principle or pattern*”.^{13,14}

26 This discussion leads to the second aspect, which is relevant in medical research. Controls are used as “types” and “patterns” to which others are compared and they are often referred to as “normal” individuals. Is, however, the value of a variable in a control individual by definition normal? In dementia research I would most often have to say no. The reason is that different scientific questions have different requirements for the controls. Sometimes the controls need to represent a general public and sometimes they need to be *free of disease*. Since selection of controls in dementia research is based on cognitive manifestations instead of a probable pre-existing, underlying pathology, the controls can seldom fulfil the requirement *free of disease* when needed. Thus, it is reasonable to speculate that deviant values in control individuals could be regarded as pathologic as well as non-pathologic. Hence, this issue influences research on biological markers for AD by shifting the way of thinking from the statement “it is normal since it occurs in the controls!” to the question “is it normal merely because it occurs in controls?”

STUDY DESIGN 1|4

The design of this study is based on a quantitative research setting. It is a prospective observational study and the collected longitudinal data allows both cross-sectional and longitudinal analyses, which will be specified in each paper.

The development of the current study and its aims can be traced back to the need for a control group at the memory clinic at Malmö University Hospital that could be used in research studies with different patient samples. As time went by and more research started to include biological markers, concerns arose in the research directory group that the controls were possibly not *free from disease* though they were cognitively healthy. The division of study samples based solely on cognition appeared very problematic due to the suspected long preclinical phase of AD. It was therefore decided that a clinical follow-up of the individuals was necessary, and I was recruited to do this follow-up. The first preliminary results from this follow-up indicated that perhaps these healthy controls were not free from disease, which was later published as the first paper of this thesis. These findings led to the arrangement of an additional, more extensive follow-up, which also included investigations of biological markers. Hence, longitudinal data was extended to include more than only cognitive performance.

The current study is an attempt to widen the knowledge of the preclinical pattern of biological markers, to explore possible mutual associations between the biological markers and to investigate possible relationships with cognitive impairment. However, this biological perspective of the disease will not be contradictory to the descriptive definition of the disease based on cognitive manifestations. As mentioned above, it is instead a separate, co-existing aspect of the disease. Consequently, the aim of this study to increase the knowledge in one of these aspects does by definition not restrict the knowledge or validity of the other. 27

THE CONTRIBUTION TO EXISTING RESEARCH 1|5

In the research of biological markers for AD the investigations of controls have been limited to creating reference values or as representing the “normal”. This has provided a certain preclinical knowledge, which is reviewed in the following chapter, but more research is needed. At the start of this thesis only one article about CSF biomarkers with focus on cognitively healthy elderly individuals had previously been published. Research on biological markers for AD needs to un-

dergo the same development as research on AD neuropsychology and neuropathology where published longitudinal research on cognitively healthy individuals can be found. At the same time it is pertinent to ask the question: How has the preclinical knowledge, that already exists, been adopted by the AD research community? This question gives rise to two new questions: “What is the rationale for control selection?” and “What kinds of controls have been used in the previous research of biological markers if represented by the CSF biomarkers?” In the following two sections these questions will be investigated.

CONTROL SELECTION 1|5|1

Selection of controls is a difficult process if the case-control samples are to be comparable. Different selection principles have been proposed but it is often impossible to satisfy them all and conflicts between them inevitably arise. Wacholder et al. have proposed four principles for control selection: 1) study base, individuals should be enrolled from the same study base to reduce selection bias; 2) deconfounding, known confounders should be controlled for and possible confounders should have as little variability as possible to avoid confounding bias; 3) comparable accuracy, errors in measurements or obtaining information should be equal between groups to reduce information bias; 4) efficiency, constraining the other principles by limiting the time and resources available.¹⁵ The principles however do not demand equality, merely comparability.¹⁵ The validity of study outcomes in fact depends on the careful consideration in control selection.¹⁶ Chui H has further suggested that study validity is decided by: 1) appropriate comparison groups, at least one of which is free of the targeted disease; 2) a clear description of the spectrum of patients and controls; 3) an independent and blind comparison of the test result with an appropriate reference (“gold”) standard.¹⁷

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In the text above there are two phrases that become relevant for this study. Firstly ... *known confounders should be controlled for* ... and secondly ... *at least one of which (groups) is free of the targeted disease*. A problem that arises in dementia research is that normative studies and control groups in the elderly traditionally do not exclude those who have preclinical dementia.¹⁸ It is well known that the sensitivity of cognitive tests is low for detecting future dementia and that the biomarkers studied often change earlier than clinical cognitive symptoms do when neuropathology is present.^{3,19,20} Hence, a situation may arise in which preclinical cases are misclassified and in the end may lead to overestimation of normative age-related changes and delay in detecting pathological impairments.¹⁸

In summary, a careful control selection is crucial in the process to determine the normative data, validity and clinical applicability of a diagnostic biomarker. Misclassification of individuals with the targeted pathology should be avoided. The threshold for controls should basically be disease or not (pathology) instead of diagnosis or not (manifestation), since clinical diagnosis emerges far too late in AD.²¹ The question that arises then, is how has this been applied to real life research settings up until now?

ANALYSIS OF CSF ARTICLES 1|5|2

To investigate how carefully the selection of controls in CSF studies has been made over the last years, a systematic evaluation was performed of the controls used in published articles on CSF biomarkers in AD. The detailed method for and the flow chart of the article selection are described in Box 1 and Figure 2.

Box 1. Method for the CSF article selection

Inclusion criteria

- 1) PubMed search on June 3rd, 2006
- 2) Combination of MESH-keywords
 - a. Alzheimer Disease
 - b. Biological Marker / cerebrospinal fluid
- 3) Published from 2003 to June 3rd, 2006
- 4) Original study using a normal material
- 5) English

Exclusion criteria

- 1) Non-AD dementia
- 2) Technique assessment independent of specific neuropathologic disease

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Evaluated variables

- 1) Impact factor of publishing journal
 - 2) Study design
 - 3) Control selection process
 - 4) Control examination
-

A PubMed search was conducted according to the set criteria and limits above. The identified articles in the set time period were reviewed and those articles that fulfilled the inclusions criteria and did not fulfil any exclusion criteria were selected. The remaining articles underwent an in-depth review where the information that was presented for the controls was abstracted. The impact factor of the publishing journal was determined by the Journal Citation Reports® in 2006, with one exception (see matrix in the supplements). An overview of the article selection process is presented in Figure 2.

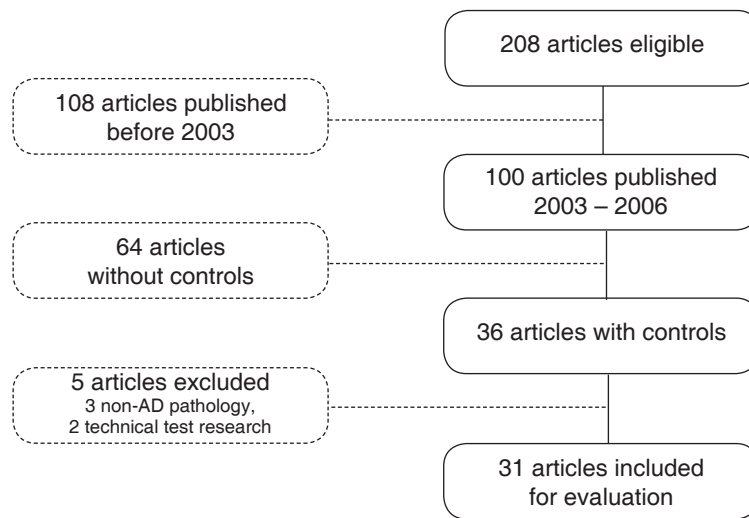


Figure 2. Flowchart of the article selection in the systematic evaluation of the controls used in CSF research on Alzheimer’s disease.

Between 1985 and June 2006, 208 articles on CSF biomarkers and AD were published. With the exception of 1998 a substantial increase in the annual number of published articles was evident during the beginning of this decade (Figure 3). Of the 31 evaluated articles almost one third (10 articles) have used control material consisting of individuals with subjective memory impairment who do not fulfil the dementia criteria (Figure 4). Moreover, over two thirds (21 articles) have used other patient groups, such as patients with neurological complaints/disorders, psychiatric disorders, spinal anaesthesia before surgical intervention or subjective cognitive impairment. Whether these individuals are truly cognitively unaffected, i.e. healthy controls, is evidently uncertain. Only 3 articles have used either population based randomization or volunteering as control selection. Furthermore, 6 articles do not provide information on whether or not cognitive assessment was made on the controls. Examination for medical conditions that could affect cognition has been stated in 20 articles and examination for psychiatric conditions in 16 articles. Thus, the number of articles examining for these possible confounding factors does not exceed the number that has used possibly unreliable samples. As a reader I can only assume that the controls in the remaining articles had been examined at selection. In alignment with the findings of this review, previous systematic reviews on diagnostic tools have reported sufficient diagnostic evaluation of the control group in only one third of available studies.²²

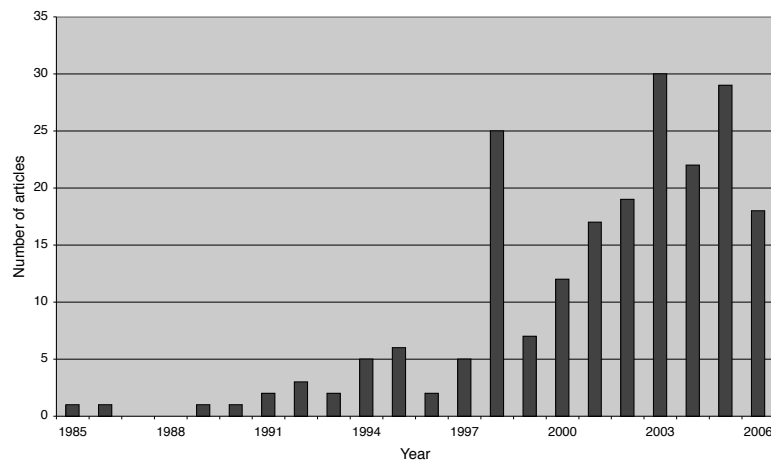


Figure 3. The number of published articles on CSF in AD before 2006.

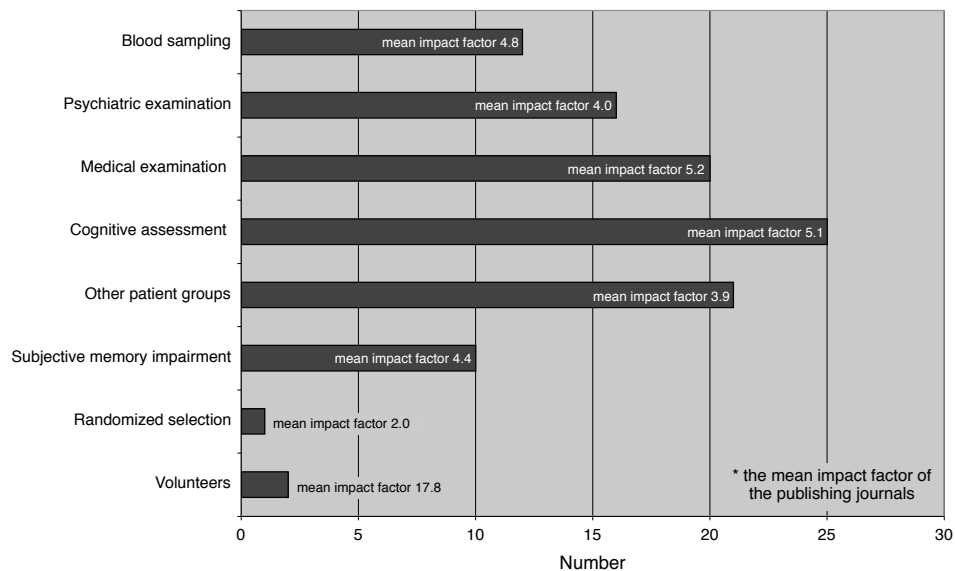


Figure 4. The number of CSF articles that described their control selection divided into different important points of information. The mean impact factor of the publishing journals was calculated for each point of information.

Moreover, one could hypothesize that high impact journals would set higher requirements for the control selection. However, the impact factor of the journal is not related to whether the controls include individuals with subjective cognitive impairment or other conditions. The only exception is the two very high impact factor journals of 12 and higher, which contain the only articles that used healthy volunteers. For the complete evaluation of the CSF articles please go to the matrix in the supplements (Table 4).

In summary, the selection of controls in the evaluated articles could increase the susceptibility of misclassification of incipient AD and thus could be in conflict with the deconfounding principle.¹⁵ Several post-mortem studies have also shown that the neuropathological burden is greater in individuals with cognitive impairment than those cognitively unaffected.²³ Consequently, the frequent use of individuals with subjective memory impairment as “healthy” controls is questionable. An additional interesting aspect is that the controls had been investigated for physical disorders in 7 of the 10 articles that used cognitively impaired individuals as controls, which are conditions that would less likely affect CSF results compared to subjective cognitive impairment of unknown origin. Hence, the knowledge of avoiding confounding factors is there, but the correlation between subjective memory impairment and possible future dementia disorder is apparently overlooked. Fortunately, it appears as if an increasing number of studies after 2006 have been published with cognitively non-impaired individuals as controls. This could perhaps be due to a decrease in the reluctance of performing lumbar puncture without clinical indication, but this is merely speculation.

LOOKING FORWARD 1|5|3

32 The answer to the general question of how the current knowledge has been adopted in previous research can be summarized as questionable. If the focus is instead turned to the future, a few circumstances emerge that require closer attention in future studies of biological markers for AD. The first of these circumstances is the need to identify the requirements of the control selection in the specific study. The second is to gather the knowledge that currently exists for the investigated marker’s preclinical pattern. The third and last is to adopt the outcomes of the former questions to the specific study. With this thesis I wish to contribute to increasing the knowledge in the second of these circumstances. In the introduction, I have therefore tried to show that there is both a need and a possibility of accomplishing this. In the case of the subsequent need to adopt the outcomes in actual study design, this is the future responsibility and duty of each individual researcher.

STRUCTURE OF THE THESIS 1|6

In the introduction, the intention is to give the background to why the scientific questions and design of the study are valid and important to make. The chosen disposition and topics of the introduction are intended as an effort to present the scientific considerations that occur in medical science but not always declared. In the following chapter I have chosen to quite extensively review the preclinical pattern of neuropathology, cognitive manifestations and relevant biological markers for this study. This is followed by the aim, design and results of the study and its separate papers. Finally the entire study is discussed and commented with the intention to take this discussion to a higher level than is possible for the articles on their own.

CURRENT KNOWLEDGE 2

DEMENTIA DISORDERS 2|1

The diagnosis of dementia is a clinical diagnosis determined by the fulfilment of criteria defined by either DSM IV or ICD-10.^{1,2} It is an arbitrarily set threshold based on the degree of decline in social and occupational activities due to impairments in different cognitive functions (Box 2). The term dementia says nothing about the underlying biological events and hence the condition can be caused by several different disorders. Some of the most common forms of dementia are: AD, vascular dementia, dementia with Lewy bodies (DLB) and frontotemporal lobe dementia (FTD).^{11,20} These diagnoses are all defined by post-mortem neuropathologic changes. In clinical practice, however, there is a need for a diagnosis when the patient is still alive. Therefore, clinical classification criteria have become available to determine the type of possible dementia disorder.^{1,2,24-28} The diagnosis of AD, however, is made primarily through excluding other dementia disorders (Box 3).^{20,24}

Current knowledge

The focus of the last decade has shifted more and more towards the underlying neuropathologies of dementia disorders and the biomarkers reflecting them. In alignment with this shift, the revised NINCDS-ADRDA research criteria for diagnosis of AD have set the cognitive requirements lower than needed for a dementia diagnosis. This makes it possible for an individual to be regarded as having AD without having dementia.³ The underlying disease (AD diagnosis) is hereby separated from the symptoms and their effect on ADL functions (dementia

Box 2. Dementia short facts.

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DEMENTIA^{1, 2, 342}

Criteria

- A. Memory disturbance.
- B. Disturbances in other cognitive functions (DSM IV: aphasia, apraxia, agnosia, or executive function) (ICD-10: thinking, judgement, language, orientation, comprehension, calculation, or learning capacity).
- C. Progressive disturbances with decline from previous higher levels of function. (ICD-10 = requires at least 6 months)
- D. Impairments in social functioning and ADL functions.
- E. Deficits also in the absence of delirium.

Prevalence

From 1 % (60–64 years of age) to 24–33 % (85 years of age or older) in developed countries. Worldwide 24.3 million people in year 2001 and 81.1 millions in year 2040.

Box 3. Alzheimer's disease short facts.

ALZHEIMER'S DISEASE ^{20, 124, 343–345}

Neuropathology:

Amyloid plaques, neurofibrillary tangles, synapse degeneration and neuron cell loss.

Risk factors:

High age. Genetic (APOE-ε4 allele and first line relative with AD). Somatic (middle-age hypertension). Intellectual (Low educational level and reduced mental activity in late life).

Cognitive symptoms at diagnosis:

Progressive episodic-memory impairment, aphasia, apraxia, and agnosia.

Prevalence:

From 0.6 % (60–64 years of age) to 14–20 % (85 years of age or older).
Worldwide 14 million people in year 2001 and 49 millions in year 2040. (AD dementia approximately 60 % of dementia cases)

Treatment – symptomatic:

Cholinesterase inhibitors (rivastigmine, donepezil and galantamine) and NMDA-receptor antagonist (memantine)

Treatment – disease-modifying:

None clinically available. Future possible agents: secretase modulators, Aβ immunotherapy, Aβ fibrillisation inhibitors, and anti-tau drugs.

Prognosis:

Life expectancy shortened and is in mean 3.5 years from diagnosis.

diagnosis). This is possible due to the fact that AD is a neurodegenerative disease in which the onset of neuropathologic changes precedes symptoms by decades.²⁹⁻³²

A coherent terminology of AD development has been suggested in the revised NINCDS-ADRDA research criteria³, in which AD dementia refers to the phase in which symptoms is severe enough to meet both dementia and AD diagnostic criteria. Prodromal AD refers to the phase in which symptoms of AD are present though not severe enough to yield a dementia diagnosis. The prodromal AD phase includes the mild cognitive impairment (MCI) condition. Preclinical AD refers to the long asymptomatic phase from the first neuropathologic changes to the first appearance of measurable symptoms (Figure 5).

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The MCI classification is, as with the diagnosis of dementia, a clinical, criteria-based definition (Box 4).³³⁻³⁷ It is an arbitrarily set threshold primarily dependant on the relative absence of decline in social and occupational activities despite presence of impairments in different cognitive functions. MCI can have the same underlying diseases as the dementia condition and a majority of the individuals with MCI progress to a dementia diagnosis before their death. There are however individuals who remain cognitively stable over time or even regain their earlier cognitive function.^{35, 37, 38} The DSM IV has a similar definition called mild neurocognitive disorder, but it is stricter and requires measurable deficits in at least two cognitive functions.¹ It has further been suggested that a pre-MCI

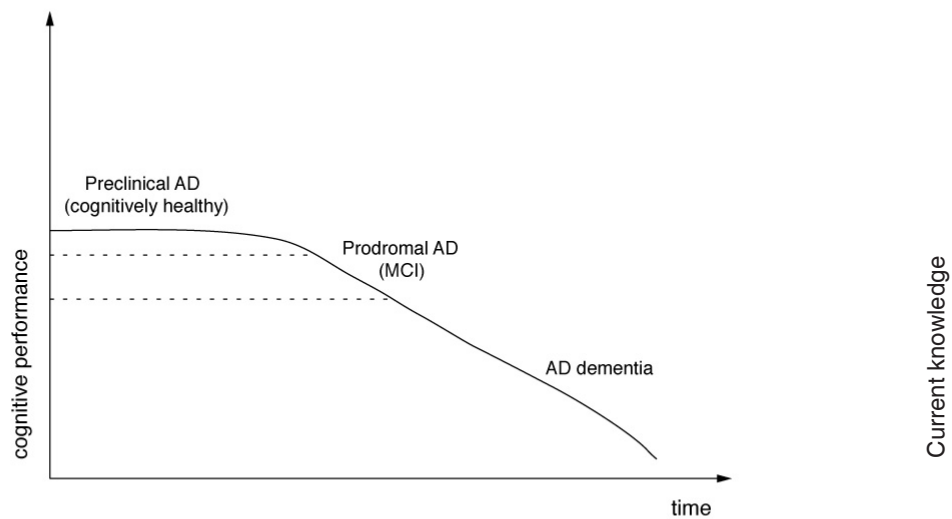


Figure 5. The natural development of a neurodegenerative disorder such as Alzheimer's disease with its preceding stages.

Box 4. Mild cognitive impairment short facts.

MCI ^{33, 35-37}

Criteria:

- A. Cognitive complaint (subjective or through informant)
- B. Objective cognitive impairment
- C. Preserved general cognitive function
- D. Largely intact ADL functions
- E. Not clinically demented.

Prevalence:

Between 3 % and 19 % in an older population. Yearly incidence rate estimated to 8 58/1000 individuals.

Prognosis:

Progression-rate to dementia is 5–15 % per year and up to 80 % have progressed after 6 years.

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phase exists, with non-measurable subjective memory impairment up to 15 years prior to developing MCI. Individuals with this subjective memory impairment have been observed to have a five-fold increase in the risk of developing MCI compared to individuals without subjective complaints.^{39, 40} However, this definition is theoretical by nature since it would be impossible to apply it to clinical practice due to a large inter-individual overlap and heterogeneous underlying pathogenesis.

In summary, dementia disorders are classified according to their symptoms and effect on an individual's ADL functions. The classification ranges from asymptomatic via MCI to dementia. Meanwhile, the individual can concurrently be classified according to the underlying neuropathology (AD, DLB, FTD, etc.) irrespective of the symptom classification (Figure 6).

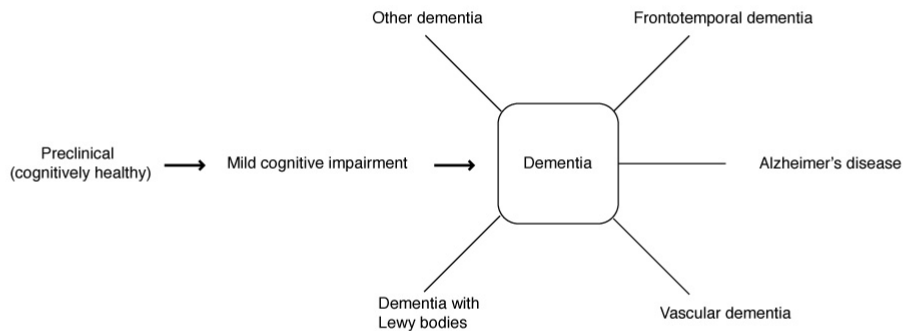


Figure 6. The development of dementia and different dementia subtypes.

AD AND ITS PRECEDING STAGES 2|2

NEUROPATHOLOGY 2|2|1

GENERAL (CLASSIFICATION CRITERIA)

38 Amyloid plaques (senile and neuritic) and neurofibrillary tangles (NFT) are hallmarks of the neuropathologic diagnosis of AD.^{20, 24, 41-43} The extracellular plaques are composed of amyloid β depositions aggregated with different cerebral cells, whereas the intracellular NFT are composed of hyperphosphorylated tau protein aggregated in the cytoplasm (Figure 7).^{20, 43-46} In AD, these histologic features are further accompanied by synapse degeneration, neuron loss and disturbance of acetylcholine transmitter activity.^{10, 20, 41, 42, 47} Several neuropathologic classification criteria have been proposed based on these changes but with different requirements for type, amount and localisation of the changes. Some of the most common criteria are the Khachaturian consensus criteria,⁴¹ the Braak neurofibrillary staging system,⁴² the NIA-Reagan criteria⁴⁸ and the CERAD⁴⁹.

Neither the generating force behind AD pathogenesis nor the precise sequence of neuropathogenic events are fully understood but some central hypotheses have been suggested. One of these is the amyloid cascade hypothesis, which argues that the accumulation of amyloid β in the brain is the trigger mechanism.⁴⁵ Increased production and oligomerization of amyloid β_{1-42} occur from an alternative cleavage pathway of the precursor protein (APP) and the subsequent ac-

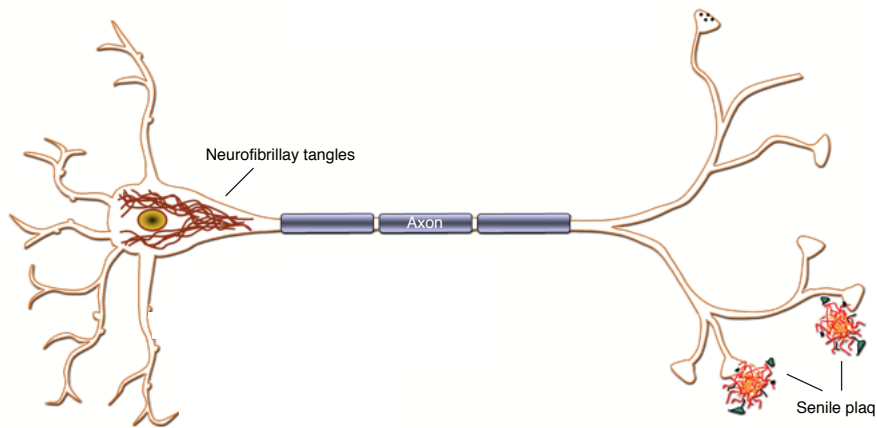


Figure 7. A neuron with AD pathology, i.e. intracellular neurofibrillary tangles (NFT) in the cytoplasm and extracellular deposits of senile plaques close to the synapses.

Modification of drawings by Professor K. Blennow. With permission.

cumulation into plaques leads to synapse and neuron damage. This in turn alters the phosphorylation of tau protein thus causing neuronal cell dysfunction and cell death. In support of this theory there are reports of neuronal toxicity of human $A\beta$ oligomers⁵⁰ as well as of an abundance of soluble $A\beta_{42}$ and subsequent plaques in individuals with Down's syndrome or hereditary AD due to an overexpression of APP. Hence, these individuals have a greatly increased risk of developing AD despite a later ensuing tau pathology.⁵¹⁻⁵⁵ Others argue, however, that tau abnormalities are the triggering mechanism.⁴⁴ It has been shown that pathologic hyperphosphorylation of tau protein leads to soluble tau proteins in the cells, which aggregate into neuropil threads and non-degradable NFT in the cytoplasm. The neuron can survive for many years with these deposits but becomes increasingly dysfunctional and will eventually deteriorate leading to cell loss. Beside these two hypotheses several other theories have been launched, suggesting that inflammatory processes, oxidative stress, or mitochondrial dysfunction are triggering mechanisms.²⁰

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In the dispute between the tau hypothesis and the amyloid cascade hypothesis, both pathologies are argued to occur exclusively from one another.⁴³ However, there is evidence from both sides of situations where the other hypothesis cannot fully explain post-mortem findings. As an alternative, Price and Morris therefore have suggested a combination of the two pathologies where both are crucial for the development of AD, as opposed to one being superior to the other.³¹ They propose that NFT alone slowly increase ubiquitously with age,^{29,}

^{56, 57} whereas plaques do not accumulate with age.^{57, 58} Only when plaques are present, the NFT increase in an accelerated AD pattern. If the neurons manage to survive during these pathologic processes the individual remains clinically asymptomatic but because of the loss of neurons, due to progressive cell death, cognitive impairment emerges.⁵⁹⁻⁶⁴

In summary, it is the general notion that the neuropathologic features in clinically diagnosed AD dementia include NFT, senile and neuritic plaques, synapse destruction, neuron cell loss and regional brain tissue loss.

AD

In early AD, NFT accumulate in the hippocampus, entorhinal cortex (EC) and subiculum whereas plaques are first observed in the EC and neocortex.^{20, 29, 43, 44, 57, 65-69} It is, moreover, in these areas that the subsequent synapse dysfunction, neuron loss and brain atrophy is first observed.^{3, 59-64, 70} In general, it appears as if the areas of the human brain that have evolved last, are the ones to be first affected by AD neuropathology.⁴⁴ Later, at the time of the clinical AD dementia diagnosis, the neuron loss is substantial compared to non-demented individuals and the location reflects the cognitive impairments present.^{20, 61-64} By now the entire temporal lobes, and especially their medial parts, are affected most intensely, followed by progression to the parietal lobes and eventually the frontal lobes.^{44, 71, 72}

40 It should be added that in individuals with clinical AD dementia the presence of multiple pathologies is frequent (20-50 %).^{73, 74} Multiple brain pathologies are suggested to increase the risk for an individual to develop dementia with an odds ratio of 20.7. These individuals also perform worse on cognitive tests and require only a mean of 1.9 NFT to develop dementia compared to a mean of 15.7 in individuals without multiple pathologies.^{73, 75} AD pathology and cerebrovascular lesions is the most commonly observed combination.^{73, 74}

In summary, neuron loss and multiple pathologies appears to be the primary events directly related to the cognitive impairment seen in AD dementia. These findings could hereby support the theories of vulnerability and reserve capacities or in other words why only some individuals develop clinical AD dementia in the presence of a certain amount of NFT and plaques.⁷⁵⁻⁷⁷

MCI

Although MCI is a heterogeneous condition, it is associated with an increased risk of developing AD dementia. This is also evident in the neuropathologic features of MCI individuals (CDR 0.5), where a substantial amount of NFT and plaques are often observed.^{29, 60, 68, 78-82} The frequency of AD neuropathology in these individuals is reported to lie between 80-100 %, ^{78, 82} with the highest frequencies in groups with amnesic MCI. One study has reported intermediate numbers of neuritic plaques in neocortical areas and NFT in the medial temporal lobe (MTL) in MCI individuals compared to mild AD dementia and controls.⁸¹ Moreover, memory function was most closely related to the number of NFT in the MTL, which is in alignment with other studies.^{66, 81}

Neuron loss in MTL and especially in layer II of the EC is substantial in MCI individuals. The degree of this neuron loss is intermediate to that seen in AD dementia and cognitive healthy individuals or equivalent to that seen in mild AD dementia.^{61, 64} The degree of layer atrophy and neuron loss is further related to the level of cognitive impairment.⁶⁴ Multiple pathologies are another histologic feature of AD dementia that can be seen on intermediate levels in MCI individuals. It has also been suggested that amnesic MCI, as compared to non-amnesic MCI, more closely resembles mild AD dementia and has a higher frequency of multiple pathologies.⁷³

In summary, the histological features of AD precede dementia diagnosis and are almost fully developed before or concurrent with the first objective cognitive impairment.

PRECLINICAL

Neuropathologic investigations of older individuals without any measurable cognitive impairment (CDR 0) almost exclusively report the presence of NFT and plaques in the brain in an appreciable number of individuals.^{23, 29, 30, 56-58, 60, 65-67, 73, 79, 80, 82-87} The frequency of AD neuropathology varies between different studies, which could be explained by different classification criteria being used but also by difficulties of standardizing a definition of a non-demented individual. In general frequencies of 10–40 % are reported,^{23, 30, 56, 57, 80, 84, 86} however frequencies as high as 90 % have been reported as well as frequencies below 10 % according to the most strict criteria.^{82, 84}

The first areas affected by these changes in non-demented individuals are primarily regions related to AD neuropathology, thus the MTL for the NFT and the temporal neocortex for the plaques.^{29, 43, 57, 65-68} In contrast to MCI and AD

dementia there is no substantial neuron loss in individuals without cognitive impairment regardless of whether or not there is an abundance of NFT or plaques.⁵⁹⁻⁶⁴ In addition, the occurrence of multiple pathologies is much more scarce (19 %) compared to cognitively impaired individuals (56%) although single pathologies are common (48 %).⁷³ Furthermore, some studies have reported correlations between the presence of AD neuropathology and poorer episodic memory,^{23, 30, 56, 57, 66, 80} although other studies have not been able to reproduce these findings.^{84, 85} This lack of coherence in the findings might be due to the same causes as discussed above for neuropathology frequencies, i.e. different classification criteria and the inexact definition of a non-demented individual.

In general, the consensus is shifting more and more towards regarding non-demented individuals with abundant AD-associated neuropathology as pre-clinical AD.^{23, 29, 56, 57, 68, 80, 85} However, it still has to be emphasized that a majority of non-demented older individuals only have a limited amount of AD neuropathology if any at all.⁸⁴ It appears to be the quantity and distribution rather than the quality of the changes that determines whether neuron loss and subsequent cognitive impairment is present.^{60, 68, 79} Meanwhile, if the histological changes represent a preclinical AD state, the individuals would not be healthy. Then the generally perceived notion that AD neuropathology is abundant in cognitively healthy elderly control subjects would be false and instead indicate presence of disease.^{56, 68}

In conclusion, all histologic features associated with AD can in fact be observed in brains of cognitively healthy older individuals. Neuropathologic findings are however of little use in the clinical every-day life since we cannot perform autopsies for neuropathologic investigation when the individual is still alive. Hence proxy markers are needed to identify the neuropathologic changes and the possible preclinical AD individuals.

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COGNITIVE FEATURES 2|2|2

AD

Cognition in AD dementia can be regarded in several ways. It is the most important symptom of the disease and disturbances in cognition are required for diagnosis.³ It is in fact what defines the clinical disease.^{1,2} Meanwhile, it is also an important marker of disease severity and is increasingly being used as a risk and prediction marker for future development of AD dementia for example in MCI.

At the time of dementia diagnosis, episodic memory and foremost delayed recall are the cognitive functions most closely related to AD.^{3, 20, 88-91} In the revised NINCDS-ADRDA criteria it has even been suggested that solitary epi-

sodic memory impairment is sufficient for a diagnosis of AD.³ Furthermore, large community cohort studies have reported that episodic memory, semantic memory, visuospatial functions, executive function and perceptual speed are the most precise in differentiating very mild AD dementia from healthy controls.⁹²⁻⁹⁵ In general the classical initial cognitive impairment triad in AD dementia beside episodic memory deficits is apraxia (praxis), agnosia (visuospatial function) and aphasia (language).^{3, 20, 88} With the progression of the disease these cognitive functions become more severely reduced and are followed by impairments in other cognitive functions. In the later stages of the disease frontal lobe cognitive impairments (i.e. personality change, fluctuating mood with aggression, apathy, etc) can be seen. In the end, almost all cognitive functions are severely impaired and the individual loses much of the ability to communicate with the surrounding world.⁸⁸

In addition, non-cognitive disturbances such as depression are also common in AD at the time of clinical diagnosis.⁸⁸ In general, the sequence of impaired cognitive functions follows the locations of the neuropathologic changes of the disease.^{44, 96} It is important to emphasize that episodic memory deficits are not pathognomonic for AD dementia and when the deficits are present other causes such as depression, somatic disease, non-AD dementia, etc have to be considered.³

MCI

Deficits in cognitive functions are the definition of the clinical condition MCI.^{35, 38, 97, 98} Due to its heterogeneous pathogenesis, the combination of impaired cognitive function will differ between MCI individuals. Episodic memory is often thought to be impaired in the individuals who progress to AD dementia and has the strongest predictive value for this conversion.^{19, 35, 90, 96, 99-102} Other cognitive impairments seen in individuals with MCI that progress to AD dementia include executive function, perceptual speed and new learning.^{35, 90, 94, 96, 101, 103} It has further been reported that MCI with impairments in multiple cognitive functions have a significantly increased likelihood of converting to dementia.^{89, 94, 103-105} However, differences in the prognosis between different impaired cognitive functions in MCI have not been observed and AD dementia and vascular dementia are similar in the extent of their cognitive deficits in the MCI stage.^{19, 35, 82, 90} There is also a considerable overlap in cognitive performance in MCI individuals converting to AD dementia and those who remain stable.⁹⁴ Hence, the pattern of cognitive impairment is not sufficient to separately predict conversion from MCI to AD dementia.

PRECLINICAL

The term preclinical in the discussion of cognitive impairment is rather contradictory since the period by definition precedes the first cognitive impairments. It is therefore inevitable that the information in this section overlaps with that in the MCI section above. However, variance in cognitive performance between individuals does not necessarily mean that someone has to be impaired.

Many large longitudinal epidemiological community cohort studies of non-demented elderly have investigated the preclinical cognitive traits of incipient AD. The longest follow-up period exceeds 20 years in one of the studies.¹⁰⁶ In all of these studies, impairment in episodic memory is a characteristic cognitive feature of AD dementia and one of the first and most severely affected in the development of the disease.^{89-91, 94, 99, 100, 102, 103, 106-113} In addition, other memory functions are also reported to be affected in these early stages,¹¹²⁻¹¹⁷ as well as other cognitive functions such as cognitive speed, executive functions, abstract reasoning, new learning and attention.^{89, 94, 103, 104, 106, 113, 115-117} The length of time from the first observed impairment to the dementia diagnosis differs between studies, depending on the length of follow-up and the cognitive function tested for. In general, onset of cognitive impairment six to nine years before AD dementia diagnosis is reported.^{90, 100, 106-108, 110-112, 114, 115, 118, 119}

Overall, it appears to be delayed memory that is the main and best validated predictive cognitive feature of future AD dementia.^{89-91, 99, 100, 106, 108, 110-112} However, despite the stringency in the findings of these large community sample studies there are a couple of aspects that need to be taken into consideration. As mentioned previously, these cognitive deficits are not pathognomonic for AD. The normal aging process probably causes deficits in the same cognitive functions as preclinical AD, i.e. episodic memory, perceptual speed and executive functions.^{19, 94} Nevertheless, this age association impairment could be due to the presence of preclinical AD in the control samples, leading to overestimation of age affect that has been discussed in the introduction.¹⁸ It has been argued that when it comes to cognition, preclinical AD should be treated as both quantitatively and qualitatively different from age-associated cognitive decline.⁹¹ Finally, it was reported in one study that one third of individuals who develop AD dementia do not present any cognitive deficits 3 years prior to diagnosis and only 38 % had both subjective memory complaints and domain-specific cognitive impairments.¹⁰⁷ Hence, some individuals tend not to progress from normal aging to clinical AD dementia through a long period with cognitive impairment.

In summary, impairments in cognitive performance are crucial for and define AD dementia but do not alone have sufficient ability to act as a proxy marker for future AD dementia.

DIAGNOSTIC BIOMARKERS 2|2|3

In AD, neuropathologic changes have previously been regarded as the diagnostic gold standard and clinical criteria as the major proxy marker for probable AD.^{3, 24} However, biomarkers within molecular science and brain imaging have in the last decades increasingly positioned themselves within the research field. The general definition of a biomarker is that it should detect a fundamental and preferably unique feature of neuropathology of the specific disorder.^{120, 121} Its qualities should be confirmed by at least two independent, well powered studies with specified control subjects.¹²⁰ Furthermore, a biomarker can have several purposes, for example as a diagnostic marker, as a marker for monitoring a disease or preceding conditions and as a prognostic or predictive marker for future disease development.¹²² Fox and Growdon have further suggested a division of biomarkers into marker of trait, state or rate. A trait marker predicts the likelihood or susceptibility of developing a certain disease, a state marker is a diagnostic marker and a rate marker detects progression of the pathophysiology.¹²¹ Sometimes the synonym “stage” is used instead of “rate”.¹²³

Current knowledge

In an attempt to define the requirements of a valid biomarker a working group at NIA-Reagan have proposed that: “*The ideal biomarker for AD ... should have a sensitivity of 80 % for detecting AD and a specificity of 80 % for distinguishing other dementias; it should be reliable, reproducible, non-invasive, simple to perform, and inexpensive.*”¹²⁰ However, the requirements of non-invasive, simple to perform and inexpensive will inevitably be relative. First, these requirements cannot be mandatory if superior and compelling diagnostic data is presented for the biomarker.¹²² Second, no clear definition can be made of the requirements and what is expensive and complicated for one individual might be regarded as inexpensive and simple by another.

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According to recent large systemic reviews/technical assessments from NINCDS-ADRDA and SBU in Sweden, the biomarkers that are currently best validated for early diagnosis of AD are CSF biomarkers, structural imaging with MRI and functional imaging with PET (Table 5 in the supplements).^{3, 124} This was also indicated in the older American Academy of Neurology practice parameters from 2001¹²⁵ and is supported by several other published reviews.^{20, 35,}

In the CSF it is amyloid β_{1-42} , total tau-protein, hyperphosphorylated tau protein and combinations of the three that have been best validated with sensitivities of 85–94 % and specificities of 83–100 %.^{3, 123, 128} In MRI it is age-corrected MTL atrophy that best distinguish and predict AD dementia with sensitivities and specificities greater than 85 %.^{3, 20, 127, 129} In assessments of atrophy, CT is inferior to MRI, however due to cost and access in clinical practice CT is useful to structurally identify treatable differential causes of cognitive impairment.^{124, 127} The PET investigation can measure blood flow, glucose metabolism and protein aggregates of amyloid and tau, and the pooled evidence for discriminative and predictive sensitivities and specificities have been reported to be greater than 80%.^{3, 127, 130} SPECT, in similarity with PET, measures blood flow but has generally produced sensitivities and specificities between 70–80 %, which is below the recommended requirements of a diagnostic marker as described above.^{3, 131, 132} Hence, SPECT is inferior to PET in accuracy although currently often less expensive and more accessible in clinical practice.¹²⁷ The EEG activity alters with the development of AD, however the diagnostic accuracy is substantially inferior to the other biomarkers and the clinical utility is concluded to be insufficient to suggest EEG as a biomarker for initial evaluation.^{124, 133} APOE- $\epsilon 4$ allele is already one of the biological trait markers for AD and increases the susceptibility of AD development. However, it has no determining position in the diagnostic evaluation of AD dementia.^{20, 124} Additional biomarkers that could be of value in the future are plasma/serum/blood markers, urine markers and functional MRI.^{122, 126, 127} The combination of different biomarkers has furthermore been reported to add to the diagnostic accuracy.^{129, 131, 134-137}

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As mentioned above all these biomarkers reflect different processes and hence will be in the transition from healthy to pathologic levels at different times of the disease development. Almost exclusively, these transitions should occur in pre-clinical stages since they reflect neuropathologic processes leading to decline in cognitive function. It can be noted that a few studies report that glucose metabolism in the EC in PET, the hippocampal atrophy rate, theta activity on qEEG and CSF biomarkers have predicted development of MCI in healthy elderly.^{134, 138-142} As discussed in the introduction, it can be speculated that trait markers occur first (APOE- $\epsilon 4$ allele), followed by state markers reflecting pathogenesis (CSF biomarkers) and later state markers reflecting cerebral and neuronal function (rCBF PET and EEG activity). Finally, state markers reflecting structure would change (MTL atrophy on MRI). The current level of evidence in preclinical AD for the biomarkers investigated in this thesis will be summarized separately in the “Biomarkers” section below.

COGNITIVE TESTS 2|3

MMSE 2|3|1

The MMSE is one of the most used cognitive screening tests in the world. Attention, verbal recall, expressive language, visual construction, orientation and calculations are the main cognitive functions tested.¹⁴³ The normative values have been derived from several large community-based population studies and are suggested to lie between 24 – 28 points.¹⁴⁴⁻¹⁴⁷ In comparison, Folstein et al. suggested a normative value of 27 in the original MMSE study.¹⁴³ The reliability of the test is reported high (0.83–0.89) and the sensitivity and specificity are suggested to lie between 69–91 % and 87–99 % respectively.¹⁴⁸ However, in a systematic review by SBU in 2006, these numbers could not be calculated since no study with the required quality standard had been published.¹²⁴ The MMSE is known to be of little value in early stages of cognitive decline and 25 % in a population-based elder control sample score 29 or 30 points out of 30 points.¹⁴⁵ Confounding factors are reported to be age (3 points decline over 20 years), educational level (1 point lower with low educational level) and possibly gender.^{145, 146, 148} However in a large population study, the MMSE score correlated with age but did not change over time in the same individual. This could suggest that the tested cognitive functions are not affected by increasing age but instead by underlying preclinical neuropathology that increases in a population with increasing age.¹⁴⁹

Current knowledge

ADAS-cog 2|3|2

The ADAS-cog is similar to other cognitive screening tests such as the MMSE but with the exception that some cognitive functions are tested somewhat more thoroughly. The cognitive functions tested are, among others, attention, verbal recall, expressive language, visual construction and orientation. The more cognitively impaired the higher results on the test, which spans from 0 to a maximum of 70 points (without delayed recall) or 85 points (with delayed recall).¹⁵⁰ Normative values are estimated to around 5 points (without delayed recall).¹⁵¹⁻¹⁵⁴ Reliability of the ADAS-cog has been reported to be high and the sensitivity and specificity are 90–100 % between healthy controls and AD dementia.^{150, 153, 155} Small confounding effects have been reported by age (primarily on delayed recall), educational level and gender.¹⁵¹

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AQT 2|3|3

AQT is a test where 40 figures are to be named according to their colour, form and combination colour and form as quickly as possible (figure 14). It is a test of cognitive speed and is dependent on visual recognition and verbal fluency, naming and automaticity.¹⁵⁶ The naming of the dual colour-form task is the most cognitively challenging. CBF measurements during the test session have reported activation of parietal lobes and deactivation of frontal and fronto-temporal lobes compared to resting levels. This is a similar pattern to those seen in tests of working memory with visual input.¹⁵⁶ Normative values are set to < 60 sec in the naming of both colour and form, whereas a result > 70 sec is suggested to be pathologic. Results between 60 – 70 sec are said to be possibly pathologic.¹⁵⁷ The sensitivity and specificity have been reported to be 97 % in one study, compared to a sensitivity of 100 % and a specificity of 84 % for the MMSE in the same sample.¹⁵⁸ The reliability of AQT was estimated to 0.88–0.96 with the same objections as for the sensitivity and specificity.¹⁵⁷ Increasing age has been the only observed confounding factor and is associated with a slower performance.^{156, 157}

CLOCK DRAWING TEST 2|3|4

In the clock drawing test (CDT) the test subject is instructed to draw a clock and put the hands to show 10 minutes past 11 (Figure 8). This can be qualitatively as well as quantitatively evaluated and a large number of assessment scales have been proposed.¹⁵⁹ Cognitive functions involved in the test are visuospatial function, constructional praxis, comprehension and semantic input with the re-

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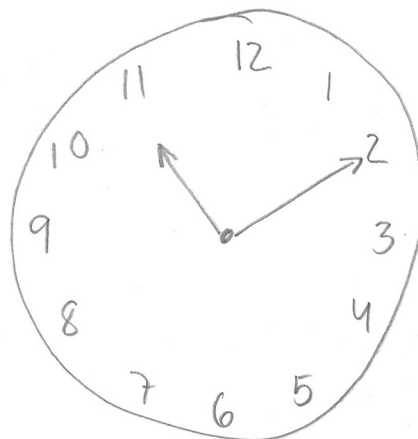


Figure 8. Correctly drawn hours and hands on the clock drawing test.

quirement of abstract thinking.¹⁵⁹ Good reliability measures have been reported however not specified and is not applicable between the different assessment scales.¹⁶⁰⁻¹⁶² The normative value is estimated to 8.5 points in the 10 point maximum score assessment scale used in the current study.¹⁶⁰ Sensitivity and specificity are suggested to be around 85 % in systematic reviews and thus valuable for cognitive screening.^{124, 159} However, others have reported much poorer numbers, and with great differences between the assessment scales.¹⁶¹ In spite of this, several studies have suggested that the CDT is more sensitive for detecting early cognitive impairments compared to the MMSE.^{159, 163} The CDT, however, has not been able to predict conversion to AD dementia in MCI samples, which limits its use.^{164, 165} Educational level has been reported to be a confounder but the test has the advantage of being useful even for illiterates.^{159, 161} Both age and gender are suggested confounders.¹⁶¹

CUBE COPYING 2|3|5

Cube copying is included in other cognitive screening tests such as the ADAS-cog. The test subject is instructed to make a copy of a transparent, three-dimensional figure in the shape of a cube (Figure 9).^{150, 166} It can be both qualitatively and quantitatively evaluated, and Maeshima et al. have suggested that the numbers of correct vertexes and plane-drawing lines should be calculated.¹⁶⁷ Cube copying tests several cognitive functions including visuospatial function and constructional praxis. The three-dimensional aspect of the test is proposed to make it more sensitive to early changes in cognition than other copying tests.¹⁶⁸ Healthy elderly perform 18 – 20 points out of a maximum of 20 points. The test has been shown to discriminate healthy controls from AD dementia and to

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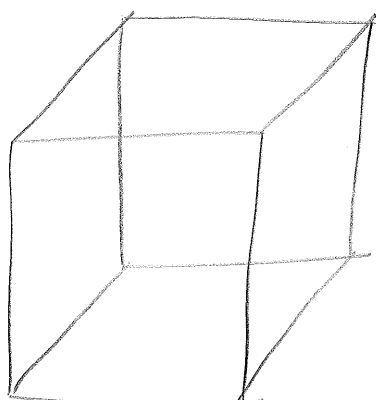


Figure 9. A correctly drawn cube on the cube copying test.

a certain extent predict conversion to AD dementia in individuals with MCI.¹⁶⁵
¹⁶⁶ Low educational level has been observed to be a confounder, whereas the reported confounding effect of age and gender is not as clear.^{169, 170}

BIOMARKERS 2|4

CSF 2|4|1

At present, the best validated biomarkers for AD in the CSF are A β 42, t-tau and p-tau.^{128, 129, 136, 171, 172} A β 42 constitutes a crucial component in the core of senile and neuritic plaques that are seen in AD. It is a proteolytic product from the trans-membrane APP in the neuron cell wall. The APP is encoded on chromosome 21 and can be metabolized in two pathways where one leads to the extracellular release of A β 42 by cleavage of the β - and γ -secretase (Figure 10).^{171, 173} Tau is instead a major component in the NFT that are seen in AD. It is an intracellular protein found in the neuronal axons and contributes to the neuron's stability and axonal transportation. If hyperphosphorylation of the tau occurs it loses these abilities and aggregates into intracellular paired helical filaments, which make up the NFT in the cytoplasm of the neuron. With neuron degeneration, tau is released into the CSF. T-tau reflects all tau released whereas p-tau reflects the hyperphosphorylated tau released. Hence, p-tau is suggested to reflect whether a pathologic phosphorylation state was present in the degenerated neurons (Figure 11).^{171, 173, 174}

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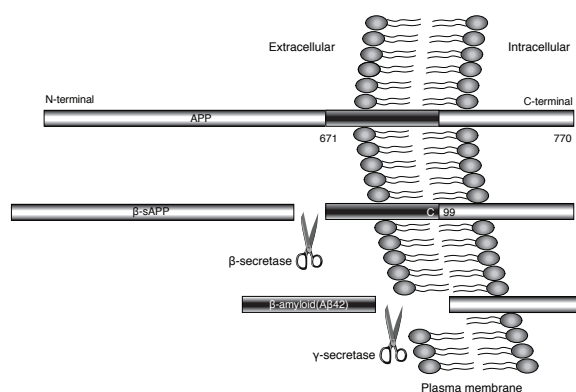


Figure 10. Metabolism of the amyloid precursor protein (APP) with amyloid β 1–42 generation.

Modification of drawings by Professor K. Blennow. With permission.

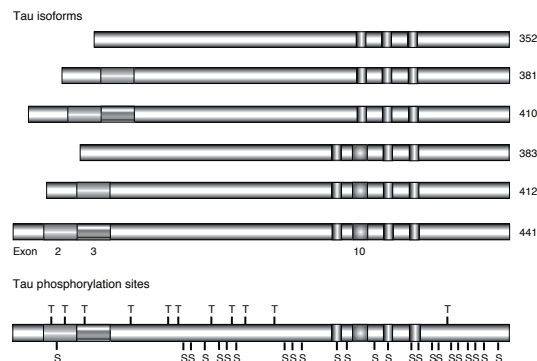


Figure 11. Tau protein isoforms and below possible phosphorylation sites of tau, either the amino acids threonine (T) or serine (S).

Modification of drawings by Professor K. Blennow. With permission.

DIAGNOSTIC ABILITY

In AD a decrease in CSF A β 42 levels and an increase in CSF t-tau and p-tau levels can be found.^{3, 123, 128, 136, 171, 172, 175} The magnitude of these changes has been assessed to a 50 % decrease in CSF A β 42 and a 300 % increase in CSF t-tau.^{136, 172} Numerous studies have reported these changes and estimated the sensitivities and specificities to well over 80 % in the discrimination between AD dementia and healthy controls. Progression in individuals with MCI has further been predicted with equally good accuracy.^{123, 128, 176-179} The diagnostic accuracy has been additionally improved by using combinations or ratios of the three.^{136, 171, 178-181} There have been reports suggesting that the main predictive ability for conversion in MCI is in the CSF t-tau and p-tau levels,¹⁸² whereas the ability to differentiate AD from other causes of cognitive impairment has instead been attributed to the CSF A β 42 and p-tau levels.^{123, 171} In addition, AD-associated changes in CSF biomarkers (low CSF A β 42 and high CSF t- and p-tau) have by some also been associated with a more advanced cognitive decline in early stages of cognitive impairment.¹⁸³

The separate changes in the CSF biomarkers are not specific to AD.^{171, 173} A decrease in CSF A β 42 levels is also seen in conditions such as DLB,¹⁸⁴ amyotrophic lateral sclerosis¹⁸⁵ and multiple system atrophy.¹⁸⁶ An increase in CSF t-tau levels is seen in conditions with massive neuron loss such as Creutzfeldt-Jakob disease (CJD) and transiently after acute stroke.^{187, 188} Mild increases of CSF t-tau levels can also be found in FTD.¹⁸⁹ In contrast, CSF p-tau is not especially increased in CJD or after stroke and hence is suggested to be a more specific biomarker for AD.^{171, 173} In some studies old age has been associated with a gradual increase of tau in the CSF^{173, 190-193} but this has been negated in other studies.¹⁹⁴⁻¹⁹⁸ In con-

trast, most studies have been consistent in showing no correlation between age and CSF A β 42 levels.^{193, 194, 198}

PRECLINICAL

In contrast to other biomarkers, the predictive value of CSF biomarkers has been investigated already in preclinical stages. Four such studies have, to our knowledge, been published until the middle of 2009 (not including the manuscripts of this doctoral thesis). The first study reported that CSF A β 42 levels were lower in non-demented, healthy older individuals who developed dementia within 3 years.¹³⁹ The second reported that high CSF t-tau:A β 42 and p-tau:A β 42 ratios predicted conversion from CDR 0 (cognitively healthy) to CDR greater than 0 (MCI or dementia) in a community-based sample of elderly over 5 years.¹³⁸ The third reported that cognitively healthy controls with high CSF tau:A β 42 ratio had a significantly increased risk of converting to MCI within 4 years.¹⁴⁰ The fourth and last study reported that low CSF A β 42 levels were the sole predictor for cognitive decline tested with MMSE or development of dementia over an 8 years period in a community-based older cohort.¹⁴¹ In summary, these studies suggest that AD-associated changes in CSF biomarkers could possibly be seen in preclinical stages and could be related to poorer cognitive performance. In addition to these studies, decreased CSF A β 42 levels have been correlated to decreased whole brain volume in healthy controls.¹⁹⁹ An interesting aspect in that study was that in mildly demented individuals the correlation was instead observed with increased CSF t-tau and p-tau levels. Nevertheless, other studies have denied such correlations between CSF biomarkers and whole-brain volume in samples with manifest AD dementia.²⁰⁰

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LONGITUDINAL STABILITY

Longitudinal changes in CSF biomarker levels have also been previously investigated. With few exceptions,^{191, 198, 201} CSF t-tau and p-tau levels have been reported to remain stable regardless if investigated in AD dementia, MCI or healthy controls.^{182, 192, 194-196, 202-208} CSF A β 42 levels on the other hand have both been reported stable,^{191, 194, 198, 205, 207-210} as well as declining over time.^{203, 204} An explanation to the difference in results is that the studies with stable levels have had an insufficient length of follow-up.²⁰⁴ Another possible explanation could be the time point in the AD development at which the investigation was performed, since one study reported longitudinal decrease in CSF A β 42 levels only in prodromal AD.¹⁹¹

CSF VERSUS OTHER BIOMARKERS

Correlations with other biomarkers for AD are not uniform. One explanation to this could be the disease stage at which the markers have been investigated. For example, the CSF biomarkers have been correlated with neuropathologic features of AD,²¹¹⁻²¹⁴ but in one study of manifest AD dementia no such correlation was seen.²⁰⁰ The CSF biomarkers can also be claimed to correlate with cognitive performance since they differentiate groups of healthy elderly, MCI and dementia from one another. However, the CSF biomarkers and cognitive performance do seldom correlate within each group,^{174, 196, 197, 215} with some exceptions.^{176, 191} In most studies APOE- ϵ 4 allele carriers have lower CSF A β 42 levels^{140, 179, 216-220} with some exceptions,^{190, 200, 210} whereas studies report inconclusive results for CSF t-tau levels.^{140, 174, 179, 192, 200, 209, 212, 218-222} Finally, hippocampal volume loss measured by MRI has in some studies been associated with AD-associated CSF changes.^{134, 205, 223}

CAUSES AND UNDERLYING MECHANISMS

The underlying mechanism behind why APP is increasingly cleaved in the A β 42 pathway and why tau becomes hyperphosphorylated is not fully known. Similarly, the mechanisms behind the AD-associated decrease in measurable CSF A β 42 levels are not either completely known. An additional disturbing fact is that AD development in Down's syndrome and familial AD, due to APP gene mutations, is characterized by an increase in APP and hence A β 42 production in young ages.⁵³ However, at the time of dementia diagnosis even individuals with Down's syndrome will display a decrease in A β 42.⁵⁴ Nevertheless, some of the proposed causes behind the decrease in CSF A β 42 levels have been the actual aggregation into plaques, decreased A β 42 production with the neuron loss, increased clearance from the CSF and deficit drainage of CSF. In contrast, the increase in CSF tau levels is thought to depend on the degeneration of neurons.¹⁷³

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In summary, changes in CSF biomarker levels are as yet among the best for the discrimination and prediction of AD dementia. In addition, they also appear promising in the preclinical stages, even though the findings need to be reproduced.

EEG 2|4|2

EEG is the recording of the brain's spontaneous electric activity due to the synchronous firing of large numbers of neurons with similar orientation. The EEG recording is assessed by the rhythmic activity, which can be divided into and quantified (qEEG) within different frequency bands. The most common are the delta (0-4 Hz), theta (4-7 Hz), alpha (8-12) and beta (< 30Hz) frequencies (Figure 16).

DIAGNOSTIC ABILITY

The characteristic qEEG changes in AD are a diffuse, low-frequency activity (low mean frequency) due to a decrease in high frequencies (alpha and beta) and an increase in low frequencies (delta and theta). In addition, a decrease can be seen in fast-wave coherence (the synchronicity of EEG rhythms between two locations).²²⁴⁻²³² The degree of these changes correlates with the disease stage.^{224, 225, 227}

54 A proposed sequence of the AD-associated changes in EEG activity has been extrapolated from data on MCI samples. First, an increase in relative theta activity is seen, followed by a decrease in relative beta activity, a decrease in alpha activity and at last an increase in delta activity.^{227, 230, 233} The increase in relative theta activity has also been reported to differentiate early stages of AD dementia from healthy controls and to have intermediate characteristics in MCI.^{142, 224, 227, 230, 233-240} Already in individuals with subjective memory impairment, the increase in relative theta activity has been reported to predict cognitive decline.¹⁴² Concurrent changes in other frequency bands have been reported but the affected frequencies differ between the studies.^{224, 230, 235, 236, 239, 241} In contrast, some studies have instead reported solitary changes restricted to relative alpha frequency to be associated to AD.^{224, 228, 235} All these AD-associated changes in EEG activity are further supported by studies other than longitudinal case-control studies. For example, administration of anti-cholinergic substances (scopolamine) to older individuals has been found to enhance the AD-associated changes in EEG activity, whereas administration of pro-cholinergic substances (acetylcholinesterase inhibitor) reduces the same.²⁴²⁻²⁴⁶

EEG RHYTHM VERSUS OTHER BIOMARKERS

The low-frequency EEG activity that is characteristic of an AD brain has further been correlated to other AD-associated changes such as hippocampal atrophy

on MRI,²⁴⁷ increased CSF tau levels,²⁴⁸ decreased rCBF,²⁴⁹⁻²⁵² regional hypometabolism,²⁵³ autopsy-confirmed neuron cell loss,²⁵⁴ presence of the APOE-ε4 allele,²⁵⁵⁻²⁵⁷ decrease in ADL-functions^{226, 258} and reduced memory performance.^{227, 228, 236, 259} Moreover, the low-frequency EEG activity has been suggested to correlate strongest with AD when located in the posterior, temporo-parietal areas.^{227, 228, 249, 260} Finally, it should be stated that there are some contradictory studies that report no alterations in EEG rhythm in MCI or mild AD dementia compared to healthy controls.^{261, 262} Furthermore, slowing of EEG rhythm and increase in relative theta activity has been associated with normal aging as well as other encephalopathies such as encephalitis, acute stroke and head trauma.^{256, 260}

CAUSES AND UNDERLYING MECHANISMS

The mechanisms behind slowing of the EEG rhythm and increase in the relative theta activity in AD are not fully elucidated. One suggested mechanism is cholinergic dysfunction.^{230, 243, 256, 263-266} The enhancement and reversibility of AD-associated changes in EEG activity in individuals when subjected to anti-cholinergic and pro-cholinergic substances, supports this theory.²⁴²⁻²⁴⁶ Alterations in hippocampal theta activity by cholinergic transmissions have further been observed in animal models.²⁶⁷⁻²⁶⁹ Another suggested mechanism is cortical hypoperfusion, since a relationship between theta activity and decreased rCBF has been observed.²⁴⁹⁻²⁵² Both mechanisms are suggested to involve hippocampal neurons, which would make the hippocampus a potentially important area of theta activity regulation.^{247, 268}

Overall, individuals with AD dementia have significantly altered EEG rhythm compared to healthy individuals. There are reports that these changes are present and measurable already when the first cognitive symptoms appear. The diagnostic and predictive accuracy has, however, not been sufficient to make qEEG an initial investigational marker in clinical practice.¹³³ The strength of EEG lies instead in the fact that it is a non-invasive, non-radiating and relatively inexpensive investigation with high-density spatial mapping and good replicable ability.²³⁹ It could therefore be of use as a second line investigational tool.¹³³

SPECT 2|4|3

DIAGNOSTIC ABILITY

Regional CBF can be measured with SPECT by obtaining images with a gamma camera after injection of a radionuclide. In AD, the change in resting rCBF occurs in certain, mainly bilateral patterns of reduction. These reductions are initially seen in the EC, posterior cingulate, precuneus and hippocampus.²⁷⁰⁻²⁷² From the start of cognitive symptoms up to moderate AD dementia a reduction is seen in the medial temporal lobes and later in parietal and posterior temporal lobes. In the more severe stages of the disease the frontal lobes become affected.^{132, 270, 273-279} And though the frontal lobe reductions are suggested to occur late there have been observations of local frontal engagement concurrent with the early posterior reductions.^{274, 276-278, 280-282}

REGIONAL CBF VERSUS OTHER BIOMARKERS

These suggested early posterior changes in rCBF best predicted conversion to AD dementia in MCI individuals^{270, 271, 274, 277, 281, 283-287} as well as the speed of cognitive decline.²⁸⁵ Moreover, the structural distribution of neuropathology according to Braak and the impairment in cognitive functions in AD dementia have been temporally correlated to the sequence of change in rCBF.^{44, 271, 273, 288} However, discordance in the location of the hypoperfusion, histologic neuropathology and atrophy can be found in the earliest stages of the disease.²⁸⁸⁻²⁹⁰ Increase in CSF tau levels have both been suggested and rejected to correlate to posterior hypoperfusion in MCI and manifest AD dementia.^{272, 278, 285} In contrast to histologic and structural features of AD, the hypoperfusion does not appear to progress with the deterioration of the disease.²⁷³ Finally, the early changes in rCBF have been observed to be reversible if pro-cholinerg substances are administered.^{132, 276} Nevertheless, compared to functional imaging with other modalities, rCBF measured by SPECT is inferior in sensitivity and specificity and has even had problems reaching a diagnostic accuracy of 80 %.^{3, 120, 131, 132, 241} According to one study 20 % of individuals with AD dementia are indistinguishable to healthy controls when hippocampal blood flow is measured with SPECT.²⁷⁹

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CAUSES AND UNDERLYING MECHANISMS

The mechanisms behind the hypoperfusion in AD are not fully understood, especially in the early stages. One proposed mechanism is a deafferentation of the areas with hypoperfusion due to remote neuronal loss in other areas.^{132, 273,}

^{276, 279, 283, 290, 291} This could explain why hypoperfusion is first seen in the posterior cingulate gyrus, whereas neuron loss and structural changes are first seen in the hippocampus and the EC.^{44, 288, 289} The non-progressiveness and reversibility of the hypoperfusion in AD could on the other hand suggest that the underlying mechanisms precede the progressive neuron loss.²⁷³ Other suggested underlying mechanisms are active oxidative stress in the posterior association cortex and a cholinergic deficit.^{292, 293} Hence, it is still uncertain whether hypoperfusion is the cause or the effect of the neuronal dysfunction associated with AD.

In summary, AD-associated changes in rCBF can be observed and when used in combination with other markers, such as MTL atrophy or CSF biomarkers, the diagnostic and predictive accuracy can be valuably improved.^{132, 272, 285, 294, 295} Changes in rCBF can also possibly contribute to the differentiation of AD from other causes of cognitive impairment.^{132, 276, 294-297}

AIM, OBJECTIVES AND HYPOTHESES 3

The neuropathologic changes in AD development precede the first measurable cognitive symptoms by decades. In spite of this, healthy clinical control samples are often selected solely due to their unaffected cognitive performance. In addition, these selection processes have not been fully controlled for possible cognitive impairment as described in the introduction. Hence, these “healthy” control samples could inevitably include individuals with AD neuropathology. Several of the recent biological markers for AD could be useful already in early, preclinical stages. Studies supporting these theories, however, are scarce. Therefore, the aim of this doctoral thesis was:

- To investigate biological and cognitive markers for AD in a group of cognitively healthy elderly individuals who have been used as clinical control subjects in research studies.

CSF biomarkers, qEEG recordings, rCBF measurements and several cognitive screening tests were the validated markers for AD available for collection in this study. The incidence of conversion to dementia in this study is expected to be low, due to the sample size and the length of follow-up, thus excluding conversion as an outcome measurement. Instead, measurable objective cognitive decline was used as an outcome measurement. In light of these prerequisites the aim of this thesis was specified in three main objectives:

1. To study the distribution of the biological and cognitive marker levels in the group. (Paper I – IV)
2. To study the cross-sectional and longitudinal relationships between the biological markers and cognitive performance on an individual level. (Paper I and III)
3. To study the relationship between the different biological markers on an individual level. (Paper II and IV)

From the current level of evidence for preclinical AD and the preclinical pattern of markers for AD, hypotheses were set for the thesis as well as for each paper.

Main thesis hypothesis:

- In a group of cognitively healthy elderly individuals there will be individuals with deteriorated biologic markers for AD such as CSF biomarkers, rCBF and electric activity and these proposed deteriorations in biological markers will be associated with proxy markers for future dementia, such as declining cognitive performance.

Individual paper hypotheses:

1. AD-associated changes in the CSF biomarkers $A\beta_{42}$, t-tau and p-tau correlate with and precede cognitive decline in a group of cognitively healthy elderly individuals. (Paper I)
2. AD-associated changes in the CSF biomarkers $A\beta_{42}$, t-tau and p-tau are related to a concurrent change in the brain electric activity in a group of cognitively healthy elderly individuals. (Paper II)
3. Individuals with a progressive deterioration towards an AD pattern in the CSF biomarkers $A\beta_{42}$, t-tau and p-tau levels will perform cognitively worse than those with stable levels. (Paper III)
4. AD-associated changes in the CSF biomarkers $A\beta_{42}$, t-tau and p-tau are related to a concurrent change in rCBF in a group of cognitively healthy elderly individuals. (Paper IV)

METHODS AND MATERIAL 4

STUDY DESIGN 4|1

The papers of this doctoral thesis are derived from a longitudinal study of clinical healthy controls at the Clinic of Neuropsychiatry at Malmö University Hospital, Malmö, Sweden. Recruitment was done through advertisements posted at locations frequented by the elderly, as well as through information directed towards non-blood relatives to patients visiting the clinic. At the initial evaluation the participants underwent a thorough examination, including medical history, physical and psychiatric examinations, and cognitive testing. In addition, they underwent a CT scan according to routine clinical practice, in order to exclude structural brain damages. Inclusion criteria at baseline were: intact ADL functions, no complaints of memory loss and cognitive test results within expected normal range. Exclusion criteria at baseline were: active physical or mental disease that could affect cognitive status, including advanced pathology on the CT scan, fulfilment of criteria for AD²⁴ or other dementia types and fulfilment of criteria for MCI.^{34, 35} When indicated, additional inclusion and exclusion criteria were added and specified in the paper concerned.

After inclusion the participants underwent a number of physical and cognitive investigations with two clinical follow-up visits over a period of 4.5 years. The physical investigations in the study were all well documented and reliable methods used to distinguish AD patients from healthy controls. They included repeated CSF analysis, APOE genotyping, EEG and SPECT-CT. The temporal order of investigations is specified in Figure 12. It should be added that the baseline CSF was collected within 6 months after inclusion. Moreover, the EEG recording was performed approximately 6 months after the cognitive assessment at the 3-year follow-up and less than one year before the cognitive assessment and CSF collection at the 4.5-year follow-up.

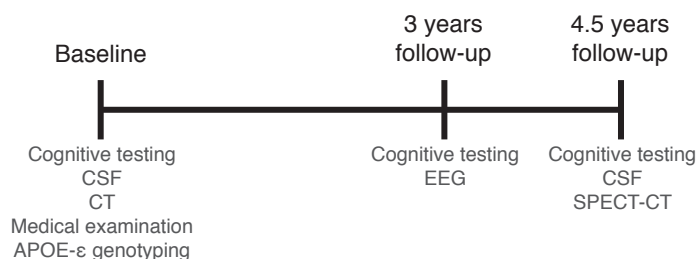


Figure 12. The investigations and assessments performed at baseline and each follow-up.

The incidence of AD dementia in a group of cognitively healthy individuals followed for 4.5 years is too low to allow these diagnoses to serve as outcome measurements in this study.²⁹⁸ Neuropathologic post-mortem examinations are similarly not feasible with the current follow-up period. Instead, subjective or objective cognitive decline without the need to fulfil any diagnosis criteria has been set as a clinical outcome in the longitudinal analyses. The first paper was written prior to the 4.5-year follow-up which limited the available cognitive outcome measurements in this paper.

The initial baseline evaluations were performed by a number of experienced physicians at the Clinic of Neuropsychiatry at the Malmö University Hospital, Malmö, Sweden. The locations and administrators of the other investigations are specified in each section below.

STUDY POPULATION 4|2

The participants in this study were initially recruited to constitute a healthy elderly control sample for studies of clinical dementia disorder. However, as specified in the introduction, the interest to separately study these presumed healthy individuals is increasing. Due to the original purpose of the study population it was important to cognitively optimize the group in order to minimize the presence of neurodegenerative pathology. The flowchart of the participants in the study is presented in Figure 13. One individual summoned to the initial evalu-

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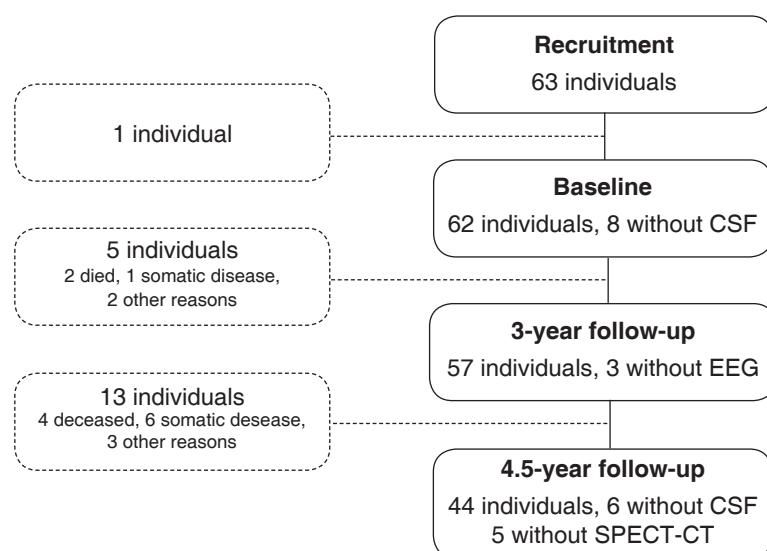


Figure 13. Participant flow chart for the entire study period including reasons for drop-out.

ation did not meet the inclusion criteria of unaffected cognitive function. Of the 62 participants at baseline 57 remained at the 3-year follow-up and 44 at the 4.5-year follow-up. The number of participants that successfully underwent each investigation is also stated in Table 1. Since different participants failed to complete different investigations the specific number of participants in each study might differ from that of the follow-ups in general.

The baseline characteristics are presented in Table 2. The mean age and sex distribution is similar to that seen in many AD studies. The frequency of APOE- ϵ 4 allele was similar to that reported in a previous control sample and lower compared to the AD sample in the same study.²⁹⁹ A pathologic albumin ratio could

Table 1. The investigations that were performed at each assessment occasion including the number of participants that completed each of these investigations.

Investigations (n)	0 – 6 months (baseline)	3 – 3.5 years (first follow-up)	4.5 years (second follow-up)
Cognitive assessments	● (62)	● (57)	● (44)
Biologic markers			
CSF collection	● (54)		● (38)
EEG		● (54)	
SPECT-CT			● (39)
CT	● (62)		
APOE genotyping	● (62)		
Medical examination	● (62)		

Table 2. Baseline characteristics of the participants including demographics, medical history, and cognitive assessments.

Participant characteristics (baseline)	Number	Percent
Sex (F/M)	41 / 21	
Age (mean \pm SD)	72.7 \pm 8.0	
APOE- ϵ 4 (homozygote)	18 (1)	29 % (2 %)
Pathologic albuminratio	5	9.6 %
Hypertension	9	15 %
Diabetes	1	1.7 %
Thyroid-disease history	7	11.7 %
Depression history	2	3.2 %
Antidepressant therapy	3	5 %
Smoking, active	4	6 %
Smoking, previous	28	45 %
MMSE score (mean \pm SD)	29.2 \pm 0.9	

be a sign of cerebrovascular disease²⁹³ and is in the study seen in a minority of the participants. Of the somatic disease history, all disease frequencies are below the numbers reported in adult population-based studies and systematic reviews.³⁰⁰⁻³⁰³ The very high mean MMSE score and low SD value reflect the effort to cognitively optimize the study population and it should be noted that the 25th to 75th percentile lies between 29 points and the maximum 30 points of the test. This can be compared to only 25 % in previous large population-based samples.¹⁴⁵ Unfortunately, educational level and ethnicity have not been registered for the participants. It can be presumed, however, that they have at least 7 years education in the publicly financed mandatory elementary school system, which existed in Sweden during the first 6 decades of the 20th century. Furthermore, there is no indication that any of the participants grew up in a country that did not have a similar mandatory school system.

The participants who dropped out of the study before the 4.5-year follow-up performed almost 1.5 points lower on the delayed word recall task on the ADAS-cog ($p < 0.001$) compared to the remaining participants. However, no difference in the crude MMSE was observed. Since declining episodic memory could be a sign of neurodegenerative disease, this could lead to fewer preclinical cases in the remaining group. If so, a possible presence of preclinical AD in the study sample would be harder to detect. No differences in age, sex, APOE genotype, or baseline CSF biomarker levels were observed between the groups. At the 4.5-year follow-up, no remaining participants had developed AD dementia. There were 3 participants whose cognitive performance bordered on an MCI diagnosis. They were re-evaluated one year later, however, without any of them progressing to a definite MCI. It was not possible to evaluate for the presence of possible MCI, AD dementia or other dementia in the dropouts.

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COGNITIVE ASSESSMENTS 4|3

The participants performed all cognitive tests consecutively during one visit on each assessment occasion. The cognitive tests used were the MMSE¹⁴³, the ADAS-cog¹⁵⁰, AQT¹⁵⁶, CDT¹⁶⁰ and cube copying¹⁶⁶. Additionally, the participants were assessed for presence of subjective memory impairment affecting quality of life (memQoL) as well as their living situation at the 3-year follow-up. Unfortunately, all cognitive tests were not performed at each assessment occasion (Table 3). At the 4.5-year follow-up the cognitive testing was done in conjunction to the subsequent CSF collection. At the other occasions, the cognitive testing was performed separate from the other investigations. All cognitive assessments

Table 3. The cognitive tests that were performed at each assessment occasion.

Cognitive tests	0 – 6 months (baseline)	3 – 3.5 years (first follow-up)	4.5 years (second follow-up)
MMSE	●	●	●
ADAS-cog	●		●
AQT		●	●
CDT		●	●
Cube-copying	● †	●	●
MemQoL*		●	●
RUD-Lite®**		●	

† Taken from the ADAS-cog. * Subjective memory impairment affecting quality of life.

** Resource utilization.

were administered at the Clinic of Neuropsychiatry at the Malmö University Hospital, Malmö, Sweden by several experienced nurses at the baseline visit and by the author at the 3- and 4.5-year follow-up visits.

The MMSE and the ADAS-cog were performed according to clinical routine for each test. The 85 point version of the ADAS-cog was used, which includes the valuable delayed word recall task. Low total score on the MMSE and high total score on the ADAS-cog reflect impaired cognitive functions. In AQT the participant is asked to name 40 figures as quickly as possible according to their colour, form and colour-form, respectively (Figure 14). The test administrator times each of the three tasks and notes any mistakes in naming. In the CDT the participant is asked to place the hours of the clock on a preset circle and the hands pointing at 10 minutes past 11 (Figure 8). To quantify the test performance we used the Ten-Point Clock Test rating scale, where points are given for placing the hours (1, 2, 4, 5, 7, 8, 10, 11) in the right sector (8 points) and the hands on ten minutes past eleven (2 points).¹⁶⁰ In cube copying the participant is asked to make a copy of a transparent cube below the original (Figure 9). The assessment of the drawings were based on a modification of a scoring system

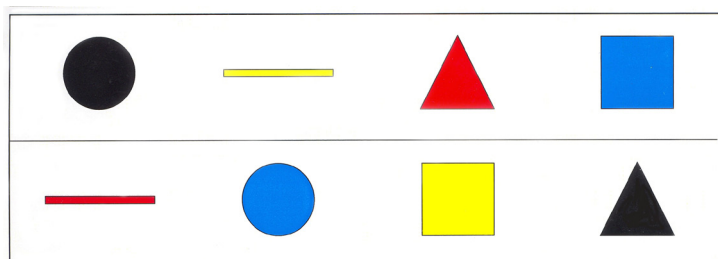


Figure 14. AQT, subtest colour-form. Short version used for giving instructions to the test subject

by Maeshima et al. where points are given for 8 correct vertexes and 12 correct parallel lines.¹⁶⁷ A vertex was defined as three lines connecting maximum 3 cm apart and the parallel limit was set to 10°. In the papers of this thesis the two separate measures have been combined into one numeric variable (20 points).

In quality of life assessment the Quality-of-Life Alzheimer's Disease instrument (QoL AD) was used, in which thirteen domains (physical health, energy, mood, living situation, memory, family, marriage, friends, self, ability to do chores, ability to do things for fun, money and life as a whole) are rated on a 4-point scale by the participant.³⁰⁴ Assessment levels are poor, not so good, good and excellent, and the first two levels were regarded as considerable decline. The memory item was used to assess the reduction of quality of life by developed subjective memory impairment (memQoL). By adding the disturbance factor of quality of life to subjective memory impairment, the use of the variable as a proxy marker for cognitive impairment is strengthened. Finally, the participants' current living situation was categorised according to a shortened version of Resource Utilization in Dementia (RUD-Lite[®]) specifying the type of housing.³⁰⁵ The 4 options available were regular apartment/house, service flats for senior citizens, old people's home, or nursing home.

CSF 4|4

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The same procedures were performed in the first and second CSF collection and analysis (Figure 15). Lumbar puncture was performed in the sitting position. The CSF samples were obtained in the L3/L4 or L4/L5 interspaces. After disposal of the first 1 ml of CSF, the following 10 ml were collected in plastic (polypropylene) tubes to avoid absorbance of amyloid β by the tube wall. All CSF samples were mixed gently to avoid possible gradient effects. No CSF sample contained more than 500 erythrocytes/ μ l. The CSF samples were centrifuged at 2.000 g at 4 °C for 10 min to eliminate cells and other insoluble material, and were then immediately frozen and stored at -80 °C pending biochemical analyses, without being thawed or refrozen. The CSF samples were analysed for A β 42, t-tau and p-tau (tau phosphorylated at threonine 181) with xMAP Luminex technology using the INNO-BIA AlzBio3 kit (Innogenetics, Ghent, Belgium) and the same batch of reagents.³⁰⁶ The results are presented in ng/l. In the longitudinal CSF study (paper III) random samples of stored baseline CSF were analysed together with the follow-up CSF in order to secure concordant assay values between the two analysis occasions. Serum samples were collected in conjunction to the first CSF collection and baseline CSF/serum albumin ratio was calculated for evaluation of the blood-brain barrier function.



Figure 15 A. Lumbar puncture for CSF collection. The L3-L4 interspace is palpated with the test subject in sitting position. *Photo by Medicinsk Informationsteknik*



Figure 15 B Lumbar puncture for CSF collection. After local anaesthetic infiltration in the skin, a needle is inserted in the L3-L4 interspace and into the spinal canal. *Photo by S. Palmqvist. With permission*



Figure 15C Lumbar puncture for CSF collection. Cerebrospinal fluid is tapped into specific test tubes before the needle is withdrawn. *Photo by S. Palmqvist With permission*

The lumbar punctures were performed at the Clinic of Neuropsychiatry at Malmö University Hospital, Malmö, Sweden by experienced physicians at baseline and the author at follow-up. The CSF samples were analyzed at the Unit of Neurochemistry at Sahlgrenska University Hospital, Göteborg, Sweden. The CSF/serum albumin ratio was determined at the Department of Clinical Chemistry at Lund University Hospital, Lund, Sweden. CSF cell counts were performed at the Department of Clinical Chemistry at Malmö University Hospital, Malmö, Sweden.

BLOOD TESTS 4|5

Baseline venous blood samples were collected from an arm vein by experienced nurses and separated into several test tubes due to the different requirements of each specific blood analyses. The samples were handled according to clinical practice at the Malmö University Hospital. The blood was analysed for blood cells, electrolytes, creatinine, liver-bile duct damage, blood lipids, vitamin B deficit, blood glucose, thyroid function, brain natriuretic peptide (BNP), prothrombin time (PT-INR), C-reactive protein (CRP), sexual hormones and APOE genotype. The blood samples were collected at the Clinic of Neuropsychiatry at Malmö University Hospital, Malmö, Sweden and the analyses were performed at the Department of Clinical Chemistry at the same hospital.

EEG 4|6

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EEG was recorded for 20 minutes from 19 electrodes with Nervus (Viasys Healthcare Inc, Madison WI) equipment, according to the 10–20 system and a sampling frequency of 256 Hz, high pass filter 0.16 Hz and low pass filter at 500 Hz (Figure 16). In order to certify that the analysis was performed on an EEG recorded with the patient fully awake, 10 sec epochs of artefact free EEG were selected in the eyes-closed situation within 5–20 sec after interaction with the patient either by verbal communication, or following eye closure on command. Post-interaction epochs were further used in order to optimize and standardize the influence of fluctuation in arousal, which predominantly causes variability in the theta activity. Several of these 10 s epochs were then analysed and one epoch was selected based on its alpha stability for further calculations.

Quantification of the EEG data reconstructed in CA mode, 2 sec epochs (Hamming filter), was performed by using commercially available software (Nervus Reader 3.4, Viasys Healthcare Inc Madison, WI). Peak frequency of

posterior dominant activity, log absolute power and relative power of delta (0.5-4 Hz), theta (4-8 Hz), alpha (8-13 Hz), beta (13-32 Hz) were calculated by FFT analysis for each quadrant of the skull: left anterior (F3,F7,T3, C3), right anterior (F4, F8, T4,C4), left posterior (P3,T5,O1), right posterior (P4,T6,O2). Average values for all four quadrants are presented as well as values for each quadrant (Figure 16).

The EEG recordings were performed at the Department of Neurophysiology at Malmö University Hospital, Malmö, Sweden. The subtraction, transformation and calculation of the EEG data were performed by an experienced physician and neurophysiologist, Professor Emeritus I. Rosén (co-author paper II).

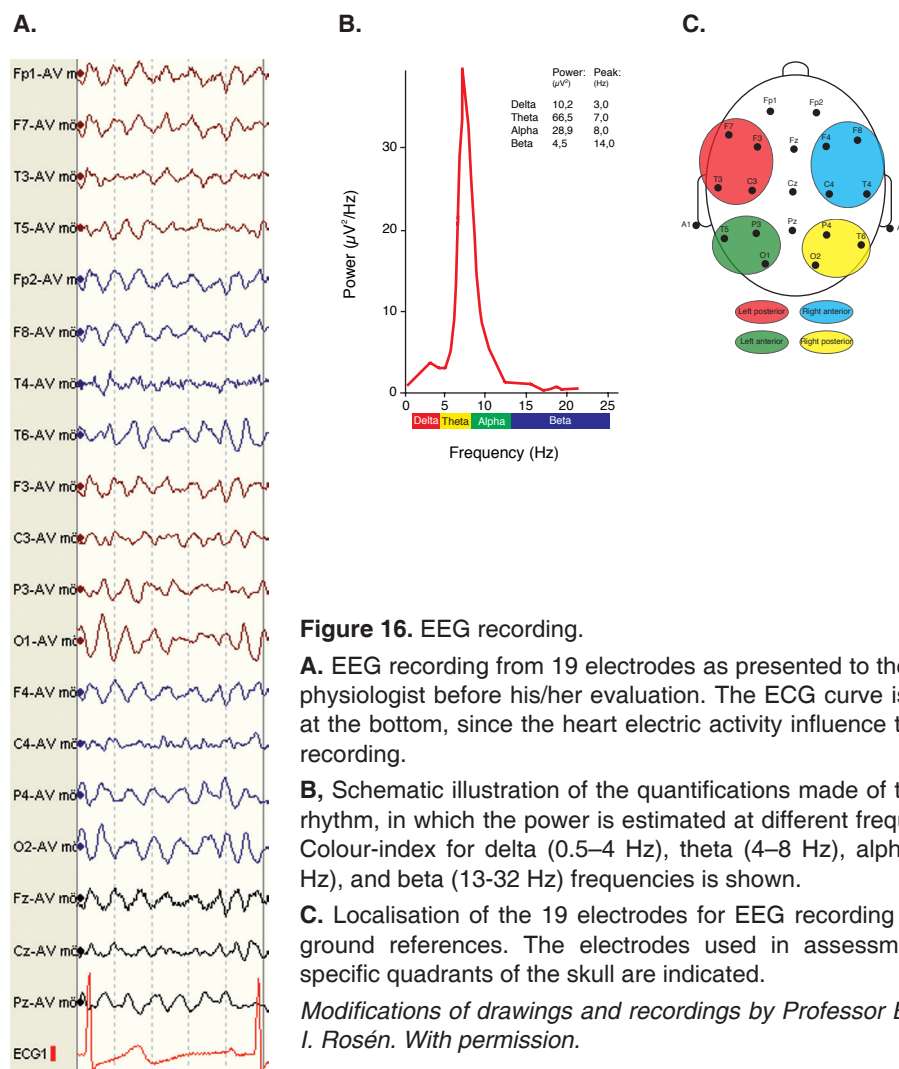


Figure 16. EEG recording.

A. EEG recording from 19 electrodes as presented to the neurophysiologist before his/her evaluation. The ECG curve is shown at the bottom, since the heart electric activity influence the EEG recording.

B. Schematic illustration of the quantifications made of the EEG rhythm, in which the power is estimated at different frequencies. Colour-index for delta (0.5-4 Hz), theta (4-8 Hz), alpha (8-13 Hz), and beta (13-32 Hz) frequencies is shown.

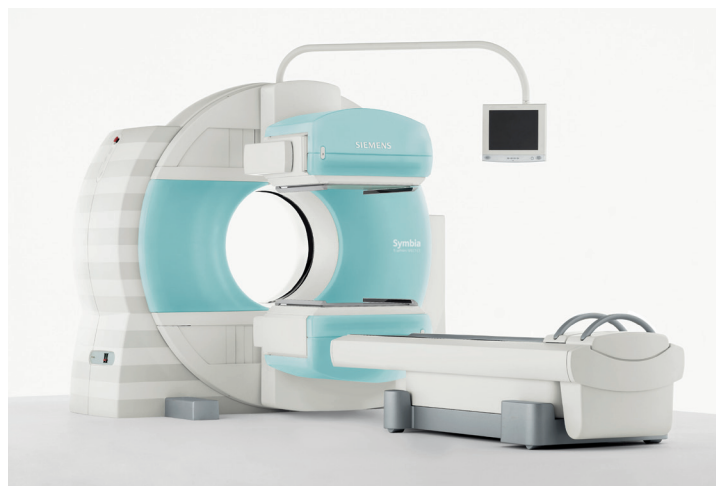
C. Localisation of the 19 electrodes for EEG recording and the ground references. The electrodes used in assessments for specific quadrants of the skull are indicated.

Modifications of drawings and recordings by Professor Emeritus I. Rosén. With permission.

SPECT-CT 4|7

^{99m}Tc -exametazime and SPECT-CT (Siemens Symbia[®] T2, Siemens Medical Solutions) were used to measure the rCBF (Figure 17). The participants were given an intravenous injection of 900 MBq ^{99m}Tc -exametazime (Ceretek[®], GE Health Care) in a well-lit setting apart from ambient noise, resting in a supine position, awake with eyes open. Image acquisition began 30 minutes later with a total acquisition time of about 40 min. The gamma camera was equipped with low energy high-resolution collimators, rotating in a non-circular orbit of 360 degrees and recording 128 projections per detector head. The recorded images were reconstructed into a $128 \times 128 \times 128$ voxel matrix using 3D-OSEM (Flash 3D, Siemens Medical Solution) with DEW scatter correction and attenuation correction based on the sequentially performed low dose CT measurements. The spatial resolution of the images was about 10 mm FWHM (full-width half maximum) at the centre of rotation.

Statistical parametric mapping (SPM5) (The Wellcome Trust Centre for Neuroimaging, London, UK) together with MATLAB 7.1 (The Mathworks) were used for voxel-based analyses. The SPECT images were normalized into a standard stereotactic space.^{307, 308} The images were then smoothed by means of an isotropic Gaussian filter (16-mm FWHM). In the statistical analysis voxels were normalized to the global level of CBF. Analysis of covariance between



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Figure 17. The SPECT-CT scan. The circular component of the machine in the back is the CT scan. The upper and lower cubical components in front of the circular component are the dual head gamma camera. The test subject lies on the bed in the front during scanning.
With permission from Siemens Medical Solution, Siemens AG

the SPECT measurements versus CSF biomarker levels and results on cognitive tests was performed by entering a regressor of respective covariate. Voxels were considered significant at a threshold of $p < 0.001$, uncorrected. Clusters were considered significant at a threshold of $p < 0.05$, corrected for multiple comparisons.

The ^{99m}Tc -HMPAO SPECT-CT scans were performed at the Department of Neurophysiology at Lund University Hospital, Lund, Sweden. Image analysis with SPM was performed at the Division of Psychiatry, Department of Clinical Neuroscience, Karolinska Institutet, Stockholm, Sweden by PhD A. Forsberg (co-author paper IV).

STATISTICS 4|8

Statistical analysis was performed using the SPSS statistics, which in 2009 was renamed PASW statistics (SPSS Inc., Chicago, IL, USA). The versions 12.0.1 (paper I), 14.0.1 (paper II), 15.0.1 (paper III) and 17.0.1 (paper IV) for Windows were used.

Non-parametric tests and median values were used when the variables were not normally distributed. Spearman rank correlation coefficient was used to test the degree of linear correlation between continuous variables (cognitive assessments, CSF biomarkers level, EEG rhythm and age). Dichotomized data was analyzed with Mann-Whitney U test (gender, memQoL, longitudinal CSF biomarkers change, outliers and dropouts). Kruskal-Wallis test was used to test nominal data with more than two labels (APOE genotype). Prediction of cognitive decline (paper I) was estimated and described with the sensitivity and specificity numbers when using cut-off-levels that optimized the Youden index³⁰⁹ as well as with the odds ratio using a logistic regression model. Level of significance was set to $p < 0.05$, where not stated otherwise. 71

ETHICS 4|9

Each performed investigation in this study was approved by the regional ethics committee of Lund University, Lund, Sweden. (LU 383-98, LU 184-01, LU 450/2006) The participants gave their written consent to participate at each assessment occasion. None of the participants had obligations towards or were dependant upon any of the clinics or co-workers involved.

On each test occasion the participants received financial reimbursement according to the investigations involved. Transportation expenses to and from

the test clinics were always compensated. At baseline the participants received a sum of 1000 SKR. The same amount was received at the 4.5-year follow-up. At the 3-year follow-up, which only included cognitive testing and the non-invasive EEG recording, the participants received gift certificates for a value of 250 SKR. The CSF samples were stored in a registered biobank (reg. nr. 68). The SPECT investigation involved exposure to radiation and a specific approval by the Isotope Committee at Lund University Hospital was acquired (IK 623).

MAIN RESULTS 5

The full version of each paper is placed in the appendix of the thesis. The main features of each paper are summarized below.

PAPER I 5|1

HYPOTHESIS

AD-associated changes in the CSF biomarkers A β 42, t-tau and p-tau correlate with and precede cognitive decline in a group of cognitively healthy elderly individuals.

SETTING

CSF analysis and cognitive assessment were performed at baseline with an additional cognitive follow-up three years later. Fifty-seven cognitively healthy elderly individuals underwent repeated assessments and CSF was successfully collected in 54 of these individuals. The CSF was analyzed for A β 42, t-tau and p-tau 181. The cognitive assessment included the MMSE, AQT, subjective cognitive impairment and current living situation.

RESULTS

No participant developed dementia or MCI during the 3 years. However subjective memory impairment affecting quality of life (memQoL) was reported in 25 % of the participants at follow-up. Participants with pathologic memQoL were slower on follow-up AQT colour-form compared to those without subjective memory impairment (Mann Whitney-U test, $p < 0.01$) and moreover had lower baseline CSF A β 42 levels (Mann Whitney-U test, $p < 0.05$). CSF A β 42 predicted pathologic memQoL with an area under the curve of 0.722 using a ROC-curve (Mann Whitney-U test, $p < 0.05$). Pathologic memQoL was best predicted by the combination of CSF A β 42 and CSF p-tau with a sensitivity of 71.4 % and specificity of 75.7 % if using cut-off levels CSF A β 42 ≤ 670 ng/l and CSF P-tau ≥ 55 ng/l (Youden index 0.471). Moreover, the same cut-off levels had a maximum odds ratio of 9.5 ($p = 0.004$) for future pathologic memQoL. Low baseline CSF A β 42 levels further correlated with a higher prevalence of living in service flats for senior citizens as well as lower total MMSE scores at follow-up. However, the MMSE score range was narrow with a 1 point difference between the 25th and 75th percentile and included the maximum score. CSF t-tau and p-tau levels individually did not correlate with any cognitive measurements.

COMMENTS

In this group of cognitively healthy elderly individuals there were participants with CSF biomarker levels that in cognitively impaired individuals would be regarded as pathologic. Additionally, the CSF biomarkers were not normally distributed which would be expected of biological markers in strictly healthy individuals. The predictive ability for cognitive impairment of low CSF A β 42 levels separately as well as in combination with high CSF P-tau levels are in alignment with findings in other current and recently published studies on healthy elderly.¹³⁸⁻¹⁴¹ These changes in CSF biomarkers have also been associated with AD dementia and conversion to AD dementia in MCI individuals.^{171, 178, 180} In this paper memQoL was used as a proxy marker for cognitive impairment since no participant developed MCI or dementia and the association with cognitive speed results supported this usage. In light of the subjective nature and heterogeneity of memQoL it is impressive that CSF biomarker levels were able to indicate future cognitive decline in a sample such as this. In conclusion, CSF biomarker levels in this paper might indicate neurodegeneration and hence AD-related biological signs could possibly be detectable already in healthy elderly individuals.

CONCLUSION

AD-associated changes in CSF biomarkers could precede cognitive decline in a group of cognitively healthy elderly individuals.

PAPER II 5|2

HYPOTHESIS

74 AD-associated changes in the CSF biomarkers A β 42, t-tau and p-tau are related to a concurrent change in the brain electric activity in a group of cognitively healthy elderly individuals.

SETTING

EEG measurement and CSF collection were performed on 33 elderly individuals with repeated normal cognitive test scores over 4.5 years, including total MMSE scores of 27 or higher. The CSF collection was performed at the 4.5-year follow-up and the EEG at most one year earlier. Cognitive assessments were performed in conjunction with both investigations and consisted of the MMSE, the ADAS-cog (only at the 4.5-year follow-up), cube copying, clock drawing test and AQT. The CSF was analyzed for A β 42, t-tau and p-tau 181. The EEG recordings underwent quantification with calculation of mean peak frequency and

relative power of the delta (0.5–4 Hz), theta (4–8 Hz), alpha (8–13 Hz) and beta (13–32 Hz) frequencies. The frequency power was calculated for each quadrant of the skull as well as average value for the entire skull.

RESULTS

Increased CSF p-tau and t-tau levels correlated with increased relative theta activity ($r_s = 0.556$; $p < 0.001$) whereas no correlation was seen for CSF A β 42. However, combined as a ratio, CSF p-tau/A β 42 and t-tau/A β 42 correlated stronger with relative theta activity ($r_s = 0.622$; $p < 0.001$). Consistently, p-tau had stronger correlations compared to t-tau. The correlations remained for each quadrant of the skull, with the strongest in the right posterior ($r_s = 0.643$; $p < 10^{-4}$) quadrant followed by the left posterior ($r_s = 0.591$; $p < 0.001$), the right anterior ($r_s = 0.575$; $p < 0.001$) and the left anterior quadrant ($r_s = 0.510$; $p < 0.01$). Both increased relative theta activity and high p-tau/A β 42 or t-tau/A β 42 ratios correlated with slower results on AQT ($r_s > 0.461$; $p < 0.01$). Also, with cognitive speed the strongest correlation for relative theta activity was seen in the posterior quadrants ($r_s > 0.503$; $p < 0.01$). No correlation was observed with the other frequency bands or cognitive tests. In the theta frequency statistical outliers were observed that in a clinical setting would be regarded as pathological EEG measurements. None of these participants differed in cognitive performance and the correlations remained even if they were excluded.

Main results

COMMENTS

In this paper cerebral function visualized by EEG rhythm and cognitive speed correlated with neuropathologic processes represented by CSF biomarkers. The correlations observed furthermore included changes previously associated with AD (i.e. increased CSF tau, theta activity and cognitive speed). The findings are in alignment with the only previous study of CSF and EEG in healthy elderly, which reported a correlation between CSF t-tau and slowing of EEG rhythm.²⁴⁸ Furthermore, several previous studies have suggested theta activity to change early in AD development^{224, 227, 230, 235, 237, 238} and especially in the posterior regions.^{227, 249} The posterior localisation of the correlations is also in alignment with the classical AD neuropathology distribution. The primary location for the first tau pathology in AD is the medial temporal lobe (MTL)⁴⁴ whereas A β 42-associated pathology is first seen in dorso-frontal regions.^{129, 171} This could partly explain the difference in the correlation patterns of CSF p-tau and t-tau versus that of CSF A β 42, since relative theta power has been associated with MTL atrophy. Moreover, neuronal cell degeneration is often suggested as the origin of

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increased relative theta activity^{230, 243, 249, 250} and the stronger correlation of p-tau in the current paper would then suggest degeneration of neurons with a pathologic hyperphosphorylation state. In conclusion CSF biomarkers and EEG theta activity might indicate early abnormal degenerative changes in the brain already in cognitively healthy elderly.

CONCLUSION

AD-associated changes in CSF biomarkers could be associated with a specific slowing of the EEG activity in a group of cognitively healthy elderly individuals.

PAPER III 5|3

HYPOTHESIS

Individuals with a progressive deterioration towards an AD pattern in the CSF biomarkers A β 42, t-tau and p-tau levels will perform cognitively worse than those with stable levels.

SETTING

In this longitudinal study CSF collection and cognitive assessment was performed at baseline and four years later. At baseline, 54 participants had a successful CSF collection and 37 of these had an additional second successful CSF collection at follow-up and are therefore included in the paper. Baseline CSF collection was performed within 6 months after the cognitive assessment. The CSF was analyzed for A β 42, t-tau and p-tau 181. Longitudinal assessments included mean difference in CSF biomarker levels as well as dichotomization in regard to the magnitude of a possible change in CSF biomarker levels. Dichotomization was done for 10 %, 15 % and 20 % change in an AD-associated pathologic direction for each of the CSF biomarkers. The cognitive assessments included the MMSE and the ADAS-cog with the addition of AQT and clock drawing at follow-up.

RESULTS

CSF biomarkers and cognitive performance did not correlate at baseline, whereas low CSF A β 42 levels correlated with lower delayed word recall score ($r_s = -0.437$, $p < 0.01$) and AQT score ($r_s > -0.385$, $p < 0.05$) at follow-up. Baseline CSF A β 42 levels could not predict cognitive performance at follow-up, which could suggest occurrence of individual changes in CSF A β 42 levels during the follow-up period. The mean longitudinal change in A β 42 levels were however not associ-

ated with cognitive performance. Instead, the participants who decreased more than 15 % in CSF A β 42 levels performed worse on ADAS-cog delayed word recall at follow-up ($z = -2.18$, $p < 0.05$) and those who decreased more than 20 % performed worse on AQT at follow-up ($z = -2.35$, $p < 0.05$). CSF p-tau and t-tau levels did not correlate with cognitive performance at follow-up, however those participants that increased more than 20 % in CSF p-tau levels performed worse on AQT at follow-up ($z = -2.13$, $p < 0.05$). No additional effect was seen if combinations of CSF A β 42, p-tau and t-tau were used. The 17 dropouts had slightly higher baseline ADAS-cog delayed word recall score (i.e. worse) than the remaining participants but otherwise did not differ. Of possible confounders age showed small associations with some cognitive test performances and presence of APOE- ϵ 4 allele was associated with higher follow-up CSF T-tau levels and a greater longitudinal decrease in CSF A β 42 levels.

COMMENTS

In this paper CSF A β 42 levels correlated with cognitive performance and individuals with a relative extensive longitudinal decrease in CSF A β 42 levels had poorer cognitive results compared to those with stable levels. This is in alignment with the findings in the three other papers of this thesis, namely that CSF A β 42 has been primarily associated with cognitive performance whereas CSF p-tau has primarily been associated with other cerebral functions such as electric activity and blood flow. However, in contrast to some of the other studies, no additional effect was observed if the AD-associated combination of CSF A β 42 and p-tau was used, even though longitudinal increase in CSF p-tau levels correlated by itself with cognitive speed. In this paper it was delayed episodic memory and cognitive speed, cognitive functions known to be affected early in AD, that correlated with CSF biomarkers. Thus, a possible concurrent decrease in CSF A β 42 levels and cognitive performance could have taken place during the follow-up period. Such a transition in markers for AD has even been proposed to occur preclinically since longitudinal stability of CSF biomarker levels has not correlated to the extent of cognitive decline in manifest disease.^{195, 196, 204} In summary, this paper suggests that CSF biomarkers previously associated with AD appear to correlate with decline in cognitive functions known to be affected early in AD development in a group of cognitively healthy elderly individuals. Hence, the results could implicate that CSF biomarkers might be able to detect the very early neurodegenerative processes of AD.

CONCLUSION

A progressive deterioration in CSF A β 42 levels could be related to decline in cognitive performance in a group of cognitively healthy elderly individuals.

PAPER IV 5|4

HYPOTHESIS

AD-associated changes in the CSF biomarkers A β 42, t-tau and p-tau are related to a concurrent change in rCBF in a group of cognitively healthy elderly individuals.

SETTING

In this cross-sectional study, 32 cognitively healthy elderly individuals underwent CSF collection, SPECT imaging and cognitive assessment. The CSF was analyzed for A β 42, t-tau and p-tau 181. The SPECT images were analysed with SPM to investigate covariance between rCBF versus CSF biomarkers and cognitive performance. The cognitive assessments included the MMSE, the ADAS-cog and AQT.

RESULTS

High CSF p-tau and t-tau levels correlated with decreased rCBF in the right superior posterior medial frontal lobe ($p < 0.001$) and increased rCBF in the left fronto-parieto-temporal border zone area ($p < 0.05$). The covariance between rCBF versus CSF p-tau had a higher degree of significance at cluster level compared to that versus CSF t-tau. CSF A β 42 was not associated with rCBF. Neither CSF p-tau and t-tau levels nor rCBF in the current right frontal and left posterior locations were associated with cognitive performance.

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COMMENTS

In this paper CSF p-tau and t-tau levels were associated with rCBF. This association between CSF tau and rCBF has previously been both reported and denied.^{272, 278, 285} The negative covariance between CSF p-tau and t-tau levels versus rCBF was found in the right medial frontal lobe at the location of the supplementary motor cortex (SMC). Previous rCBF studies on AD and MCI individuals have reported a similar frontal rCBF decrease, however most often as secondary findings concurrent with decreases in more posterior areas of the brain.^{274, 278, 280, 282, 288} In addition, the SMC is highly involved in executive functions^{310, 311}, which is one of the earliest cognitive functions to be affected in AD development.^{89, 95}

Meanwhile, the positive covariance in the paper was located in the left fronto-parieto-temporal area. Increased rCBF has often been reported in AD and MCI individuals^{274, 277, 312, 313} and this increase has been hypothesized to be a possible initial compensatory mechanism in situations of neuronal stress, for example incipient AD.^{312, 313} Hence, in the current paper, biological markers related to AD (i.e. CSF biomarkers and rCBF) appear to already correlate in cognitively healthy elderly individuals. These findings could hereby suggest that these biomarkers might indicate preclinical neurodegenerative changes in the brain.

CONCLUSION

AD-associated changes in CSF p-tau and t-tau levels could be related to specific right medial frontal and left fronto-parieto-temporal changes in rCBF in a group of cognitively healthy elderly individuals.

COMMENTS 6

In the previous chapter, comments were made for each specific paper. However, there are more general aspects and circumstances, which influence the entire study. In the “Thesis at a glance” section a structured summary of the thesis is presented. Here below certain issues will be discussed more closely. First, it should be recognized that the explorative nature of this study limits one from drawing extensive conclusions. The findings in this study are only representative for this sample and can not be generalized to explain definite patterns in population-based samples. Instead the conclusions in this study are valuable in a wider perspective because they can be used to generate hypotheses for future research in other study samples.

Comments

THE OBJECTIVES 6|1

1st OBJECTIVE

The first objective of this thesis was to study the distribution of markers. The variance in cognitive performance was, for obvious reasons, limited due to the cognition-based selection process. However, the biological markers in this study did not show the same narrow distribution. Instead the values of the biological markers ranged over a relatively large interval. Furthermore, it could be noticed that the CSF biomarkers did not have a normal distribution, which could be expected from a biological marker in a control sample without extensive disease pathology. In fact the distribution of CSF biomarker levels in this group of cognitively healthy individuals resembles the distribution in groups with AD individuals.³¹⁴ The characteristic of this specific distribution is the occurrence of a prolonged tail of individuals towards the direction of levels associated with AD (Figure 18).

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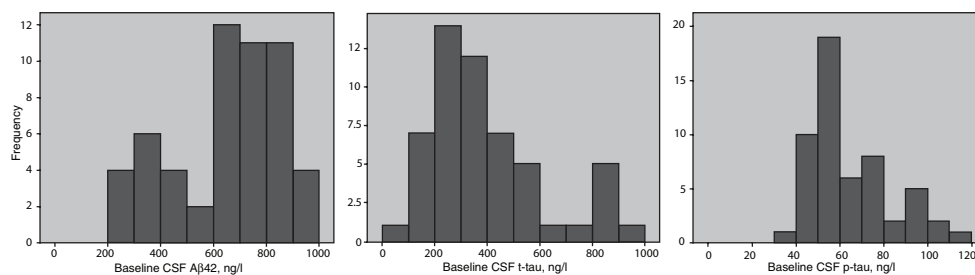


Figure 18. Distribution of CSF Aβ42, t-tau and p-tau levels in the study sample of cognitively healthy elderly individuals. A prolonged tale of the distribution can be seen in the directions that are associated with AD dementia.

Finally, within the current study sample statistical outliers (> 1.5 SD of the median value) were observed in the papers for some biomarkers, further suggesting a relatively wide range of values. If the individuals in the sample were assessed for each separate biomarker with AD dementia clinical reference standards from the participating neurophysiology and neurochemistry departments, several clinically pathologic biomarker levels were observed. Four clinically pathologic EEG recordings were qualitatively assessed by the neurophysiologist. Ten participants had pathologic rCBF when qualitatively assessed by the SPECT-supervising neurophysiologist and around the same number had participants CSF biomarker levels outside the cut-offs used at the memory-clinic.

2nd OBJECTIVE

The second objective was to study the relationship between biological markers and cognitive performance. The study suggests that in the sample there is a longitudinal relationship between these variables, which gives rise to a cross-sectional relationship during the follow-up. As has been previously stated in the thesis, the crude cognitive tests used created difficulties in detecting true differences. Despite this there were cognitive functions that correlated with the biological markers.

The first interesting aspect is that it was cognitive functions associated with early AD development, i.e. delayed episodic memory and cognitive speed, which correlated with the biomarkers. Secondly, it is not surprising that the MMSE scores did not provide any relevant correlations with the biological markers in this study. Thirdly, in the first paper, impairments in a subjective measurement of cognitive function at the 3-year follow-up were related to CSF biomarker levels. Later at the 4.5-year follow-up, it was instead impairments in objective measurements of cognitive functions that correlated with CSF biomarker levels. Hence, this shift could strengthen the belief that a true association between CSF biomarker levels and cognitive decline exists in this group of cognitively healthy elderly. Finally, it should be held in mind that the variance in cognitive performance could have other underlying causes than neurodegenerative disease and in the current study these possible confounding factors could not be excluded or completely adjusted for.

3rd OBJECTIVE

The third objective was to study the relationship between the different biological markers. The study suggests that in this sample the different biomarkers have an association to one another and as for the cognitive variables it is changes in the biomarkers previously related to AD dementia that correlated with one another.

The first comment that could be made concerns the difference in correlation patterns of the CSF biomarkers seen in this study. The CSF tau levels appear to be more closely related to the other biological markers whereas the CSF A β 42 levels appear more closely related to the cognitive performance. Despite the limitations in the ability to draw conclusions the findings are in alignment with findings in other studies. It is primarily CSF A β 42 levels that have correlated to cognitive performance in samples of healthy elderly.^{139, 141} In AD dementia and MCI it is instead CSF tau levels that have correlated to the cognitive performance.⁴³ The few studies that have investigated the relationship between rCBF or EEG activity to CSF biomarkers have reported correlations only to CSF tau.^{248, 272, 285} A difference in the correlation pattern between the CSF biomarkers is further supported by the findings of Fagan et al. where whole-brain atrophy correlated to CSF A β 42 levels in the healthy controls but to CSF tau levels in individuals with very mild dementia.¹⁹⁹ Hence, the difference in correlation patterns of the CSF biomarkers between the papers of this study does not make the relationships less likely to reflect true associations.

The second comment of this third objective concerns the difference in spatial location of the correlations. CSF tau levels correlated primarily with right frontal and left posterior rCBF but right posterior EEG activity. This difference in locations could be explained by that the changes in the biological markers with spatial assessments are possibly caused by remote effects that originate from the regions of the first neuropathologic changes.^{152, 276, 289} In addition, the localisations of each correlation were never in opposition to what had been previously reported for that specific biomarker.

The third comment regards the fact that several previous studies have not been able to observe the current correlations between biological markers and cognitive function within manifest AD dementia samples. There can be several reasons for this but one of them can depend on the kind of marker that is studied. A state marker that reflects disease or no disease will, at one point in the disease development, change to reach a plateau-level where it stabilizes. Hence, from this moment the state marker will not correlate to a rate marker that reflects the progression of the disease and therefore continuously change.

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THE STUDY SAMPLE 6|2

At the baseline of this study the aim was to recruit a cognitively optimized control group, i.e. “supernormal” individuals. The result of this is that conversion to MCI or dementia would be less likely compared to that in a population-based

or hospital-based control sample. As calculated in the discussion of paper I, the theoretical likelihood of someone developing dementia in this study is low. The fate of the dropouts is further unknown and conversion to MCI or dementia in that group has not been possible to assess. Nevertheless, at baseline the dropouts had significantly worse cognitive test scores on the MMSE (1 point) and the ADAS-cog (1 point) compared to the remaining participants at the 4.5-year follow-up. In summary, it could be reasonable that conversion to MCI and AD dementia was not available as outcome measurements in this study.

However, three participants at the 4.5-year follow-up were on the borderline to fulfil the MCI criteria due to a deviant score on a single cognitive test. These individuals were re-assessed after an additional one year but did not deteriorate further and instead some improved their deviant score. In light of this, these few individuals were not treated as a separate group in the longitudinal analyses but they were excluded from the cross-sectional follow-up analyses as specified in the papers concerned. On a general note it should be mentioned that the sample could never be as cognitively optimized at follow-up as it was aimed to be at baseline.

THE AIM AND THE FUTURE 6|3

The aim of this study was to investigate biological and cognitive markers for AD in a group of cognitively healthy elderly individuals. All papers of this study could suggest that biological markers and foremost CSF biomarkers might indicate neuropathology in this sample. In most of the cases the findings of this study can be linked with a possible connection to AD and primarily its early stages. Thus, the study could suggest that biological markers and foremost CSF biomarkers might indicate early neurodegenerative changes of the brain in this sample.

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The key word in the previous sentence is “*could*”, even though this potential ability of the biological markers is more probable now than it was before this study was performed. Hence, the study confirms that biological markers are an interesting area of future preclinical AD research. It reinforces the importance of planning and performing new preclinical studies, which should be larger and better powered than the current one. It also suggests that it could be relevant to continue the cognitive follow-up of the participants in the current study. Finally, it supports the need of thorough control selections in the future, if the sample is to be treated as “free from disease”. To conclude, the pathological processes prevailing in AD might bridge the clinically created arbitrary division of normal and non-normal aging of the brain.

CONCLUSIONS 7

Specific hypotheses were in advance proposed for each paper as described in the “Aims, objectives and hypotheses” of the thesis. Based on these hypotheses, the study concludes that:

- In a group of cognitively healthy elderly individuals there existed individuals with deteriorated levels in AD-associated biological markers.
- These deteriorations in biological marker levels could in part be associated with worse cognitive performance.
- Biological markers previously associated with AD might indicate early neurodegenerative changes in the brain also in cognitively unimpaired individuals.

ACKNOWLEDGEMENTS 8

There are several people that have contributed to the making of this thesis. To these people, I would like to express my deepest gratitude.

A critical, curious and ambitious scientist is not created on its own. As supervisor, **Elisabet Londos** gives her postgraduate students the unique possibility to develop these important skills. Without her inspiration and faith this thesis would not have been possible and I would not be doing research. It has been an honour and an enormous privilege to work with you.

I have had the opportunity to learn and delve into the research field of dementia disorder and to do this as an occupation. I have also have had the opportunity to take part in a fantastic longitudinal research project. All of this was possible thanks to my co-supervisor and moreover my executive director **Lennart Minthon**. You make things possible.

As co-supervisor, **Oskar Hansson** has been the role model of a critical, curious and ambitious scientist. You have added an additional dimension to my research.

I started this journey together with my friend **Maria Andersson**. As postgraduate students we have enjoyed many rewarding and amusing moments together. For me, dementia research will always be associated with you. I would also like to thank the entire group of current and previous postgraduate students at the clinic, **Cecilia, Carina, Eva, Fredrik, Gustav, Henrietta, Kajsa, Viktoria, Peder, Sebastian** and **Åsa**, and the senior scientist, **Katarina**. It has been fun and I look forward to more collaboration in the future!

I would like to give my warmest regards to everyone involved in the execution of the study and those who contributed greatly to the data collection. **Eva Falk-Langebros** and **Tarja Tikkanen**, I enjoy working with you. The clinical trial group at the Department of Neuropsychiatry, **Annacarin, Cissi, Marie and Pia**, you make collaboration fun and easy. **Petra Henriksen**, come back soon.

To all co-writers I am grateful and honoured that you wanted to work with me, **Kaj Blennow, Anton Forsberg, Douglas Hägerström, Ingmar Rosén, Erik Ryding**, and **Henrik Zetterberg**.

I would like to thank **Roberta Boson** for linguistic support and **Nadja Andrade Paes Maya** for the art work in this thesis.

This thesis would not have been possible if it wasn't for the flexibility of my employers at my clinical workplaces, **Department of Neuropsychiatry** at UMAS in Malmö, **Kalmar County Hospital** and **Emmaboda Health Centre**.

Jag vill tacka min **familj** och alla mina **vänner** för att ni finns där alla de stunder då arbetet inte lockar och jag behöver ert sällskap. Forska vore inte lika roligt om jag inte visste att ni fanns som belöning efter en hård dags arbete.

Finally, I would like to emphasize the incredible effort and contribution all of the **research subjects** have made. They have not been simple investigations we have put you through. I hope that you have gained as much from our meetings as I have.

This thesis was made possible by unconditional grants from Region Skåne, The County Council of Kalmar, and Lund University.

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SUPPLEMENTS 10

Supplement 1

CSF articles analysis matrix

Supplement 2

Evidence for diagnostic biomarkers

Supplement 1: CSF articles analysis matrix

Article	Journal	Impact factor	Study design	Number of controls	Volunteers	Randomized selection	Subjective memory impairment	Other patient group	Specification	Cognitive assessment	Physical examination	Psychiatric examination	Blood sampling
Rota et al. 2006 ³¹⁵	Neurol Sci	0.8	P - C	25				●	Neurologic complaints	●			
Bibl et al. 2006 ³¹⁶	Brain	7.3	P - C	23			●	●	Neurologic complaints	●	●	●	
Biroccio et al. 2006 ³¹⁷	Proteomics	6.1	P - C	27				●	Neurologic complaints		●		
Castaño et al. 2006 ³¹⁸	Neurol Res	1.6	R - C	43						●			
Mielke et al. 2006 ³¹⁹	Alzheimer Dis Assoc Disord	2.0	R - C	86		●					●	●	●
Hansson et al. 2006 ¹⁷⁸	Lancet Neurol	12.2	P - L	39	●					●	●	●	
Galimberti et al. 2006 ³²⁰	Neurology	4.9	P - L	40				●	Neurologic complaints		●		
Strekalova et al. 2006 ³²¹	Neurobiol Aging	5.3	P - C	46				●	Exclude subarachnoidal hemorrhage				
Blasko et al. 2006 ³²²	Dement Geriatr Cogn Disord	2.6	P - C	27				●	Surgical intervention	●			
de Leon et al. 2006 ²⁰⁵	Neurobiol Aging	5.3	P - L	9			●	●	Subjective memory impairment	●	●	●	●
Kapaki et al. 2005 ³²³	Int J Geriatr Psychiatry	2.2	P - C	50				●	Surgical intervention	●	●	●	●
Schoonenboom et al. 2005 ³²⁴	Ann Neurol	7.6	P - C	26			●	●	Subjective memory impairment	●			
Maruyama et al. 2005 ³²⁵	Ann Neurol	7.6	P - L	27			●	●	Subjective memory impairment	●			
Zhang et al. 2005 ³²⁶	J Alzheimers Dis	5.1*	P - C	31						●	●	●	●
Georganopoulou et al. 2005 ³²⁷	Proc Natl Acad Sci U S A	10.2	P - L	15						●			
Ahmed et al. 2005 ³²⁸	J Neurochem	4.6	P - C	18				●	Neurologic complaints		●		
Walter et al. 2004 ³²⁹	Neurobiol Aging	5.3	P - C	30				●	Surgical intervention				
de Leon et al. 2004 ¹³⁴	J Intern Med	4.0	P - L	2						●			
Schoonenboom et al. 2004 ³³⁰	Neurology	4.9	P - L	21			●	●	Subjective memory impairment	●			
Lewczuk et al. 2004 ³³¹	J Mol Neurosci	2.6	P - C	9				●	Neurological / psychiatric complaints	●	●	●	●
Ryberg et al. 2004 ³³²	Neurochem Int	3.0	P - C	19						●	●	●	
Clark et al. 2003 ³³³	Arch Neurol	4.9	P - L	69			●	●	Subjective memory impairment	●	●	●	●
Puchades et al. 2003 ³³⁴	Brain Res Mol Brain Res	1.6	P - C	7			●	●	Psychiatric disorder	●	●	●	●
Ganzer et al. 2003 ³³⁵	J Neural Transm	2.5	P - C	68			●	●	Dementia disorder	●	●	●	●
Sunderland et al. 2003 ¹⁷⁵	JAMA	23.3	P - C**	72	●					●	●		●
Sáez-Valero et al. 2003 ³³⁶	J Neurosci Res	3.2	P - C	106						●	●	●	●
Bagli et al. 2003 ³³⁷	Eur Arch Psychiatry Clin Neurosci	2.3	P - C	25			●	●	Dementia disorder	●	●	●	●
Schönknecht et al. 2003 ³³⁸	Neurosci Lett	1.9	P - C	16				●	Surgical intervention	●		●	
Kapaki et al. 2003 ³³⁹	Eur J Neurol	2.2	P - L	46				●	Surgical intervention	●	●		●
Andreasen et al. 2003 ³⁴⁰	Acta Neurol Scand Suppl	0.3	P - L	32			●	●	Subjective memory impairment	●	●	●	
Buerger et al. 2003 ³⁴¹	Am J Psychiatry	8.3	P - C	21						●	●	●	
Total					2	1	10	21		25	20	16	12
Mean impact factor (without impact > 12)		4.9 (4.0)			17.8	2.0	4.4	3.9		5.1	5.2	4.0	4.8

P = prospective, R = retrospective, C = cross-sectional, L = longitudinal. * as of 2009-06-23. ** manuscript including a separate meta-analysis part

Supplement 2: Evidence for diagnostic biomarkers

	NINCDS-ADRDA ³ 2007	SBU ¹²⁴ 2006
CT	n/a	● ● ●
MR	● ● ●	● ● ●
PET	● ● ●	● ●
SPECT	● ●	● ●
CSF	● ● ●	● ● ●
qEEG	n/a	●
Gene mutation testing	● ● ●	n/a
APOE-ε4 allele	n/a	●

- Limited diagnostic value
- ● Moderate diagnostic value
- ● ● High diagnostic value
- n/a not applicable