From mild cognitive impairment to subjective cognitive decline: conceptual and methodological evolution

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Abstract: Identification of subjects at the early stages of Alzheimer’s disease (AD) is fundamental for drug development and possible intervention or prevention of cognitive decline. The concept of mild cognitive impairment (MCI) evolved during the past two decades to define subjects at the transitional stage between normal aging and dementia. Evidence from cross-sectional and longitudinal studies has shown that MCI is associated with an increased risk of positive AD biomarkers and an increased annual conversion rate of 5%–17% to AD. The presence of AD biomarkers in subjects with MCI was associated with an even higher risk of progression to dementia. However, earlier clinical trials for pharmacotherapy in subjects with MCI were disappointing. To extend the spectrum of AD to an earlier stage before MCI, subjective cognitive decline (SCD) was introduced and was defined as self-reported cognitive decline before the deficits could be detected by cognitive tests. Subjects with SCD have an increased risk of underlying AD pathology. However, SCD can also develop secondary to other heterogeneous etiologies, including other neurodegenerative and psychiatric diseases, personality traits, physical conditions, and medication use. Several clinical and biomarker features were proposed to predict risk of conversion to AD in subjects with SCD. Further longitudinal studies are needed to support the validity of these high-risk features.

Keywords: mild cognitive impairment, subjective cognitive decline, preclinical Alzheimer’s disease, Alzheimer’s disease

Preclinical stages of Alzheimer’s disease as a potential therapeutic target

Alzheimer’s disease (AD) is the most important cause of dementia in the elderly population. Although much effort has been made in the development of therapies to stop the progression of AD, acetylcholinesterase inhibitors and the N-methyl-D-aspartate (NMDA) receptor antagonist memantine are the only two classes of medication that have modest positive effects on cognitive decline.1,2 Based on the evidence from neuroimaging, neuropathological and biochemical studies, it was established that the pathophysiological process of AD begins years or even decades before cognitive decline.3–5 One possible explanation for the failure of previous drug trials is that it may be too late to start treatment when there is evident cognitive impairment, and neuronal injury and synaptic dysfunction have advanced beyond the point of reversibility. Identification of subjects at an early stage is crucial for therapeutic intervention and possible prevention of cognitive decline. During the past several decades, two approaches have been used to identify subjects with early AD. One approach is to look for subtle cognitive changes before overt dementia, and the other approach is...
to look for surrogate biomarkers of Alzheimer’s pathology. On the clinical spectrum of AD, there is no definite cut-off point to discriminate between normal aging and dementia. By using more sensitive neuropsychological tools, subjects at the early end of the AD spectrum could be identified, but possibly at the expense of increased diagnostic uncertainty. The concept of preclinical AD evolved in response to the need to identify subjects with Alzheimer’s pathological process before the onset of significant cognitive decline. Preclinical AD was initially used to describe subjects with neuropathological evidence of AD without detectable cognitive changes. Following the development of biochemical and neuroimaging biomarkers, the National Institute on Aging and the Alzheimer’s Association (NIA-AA) recommended a staging schema for preclinical AD based on biomarker status. Subjects without cognitive decline are classified as stage I based on the presence of amyloidosis biomarkers, and as stage II if biomarkers for both amyloidosis and neuronal injury are present. Subjects with biomarkers for both amyloidosis and neuronal injury and subtle cognitive decline (well above the cut-off for mild cognitive impairment [MCI]) are classified as stage III. The application of biomarkers can help specify the earliest pathophysiological changes of AD and increase diagnostic certainty in subjects with subtle cognitive decline.

Mild cognitive impairment: the transition from normal aging to dementia

Since the early 19th century, researchers have attempted to identify the transitional state from normal aging to pathological cognitive decline. The evolution of diagnostic criteria for MCI is summarized in Table 1. The term mild cognitive impairment was first used to describe stage 3 of the global deterioration scale (GDS) for aging and dementia. At stage 3 of the GDS, subjects exhibit subtle deficits in cognition that affect complex occupational and social activities but do not yet meet the criteria for dementia. In 1999, Petersen et al redefined MCI as a syndrome of cognitive decline beyond that expected for an individual’s age and education level, but that does not notably interfere with activities of daily living. The original criteria focused on memory performance, which is often the earliest symptom of AD. However, decline in other cognitive domains can develop during or even before memory impairment. To broaden the concept of MCI, the Key Symposium was held in Sweden and published consensus criteria for MCI in 2004. To emphasize the heterogeneity of the clinical presentation and multiple underlying etiologies of MCI, the Key Symposium criteria expanded the definition of MCI beyond the memory domain and further classified MCI into three subtypes: amnestic, multiple domain, and single non-memory domain MCI. Early population-based studies showed that amnestic MCI affects $5\%$–$10\%$ of the elderly population. Adapting the expanded MCI criteria, the prevalence of MCI increased to $8\%$–$25\%$ of the elderly population aged $\geq 60$ years. In a systematic review published in 2013, the annual conversion rate of MCI to AD ranged from $7.5\%$ to $16.5\%$ per person-year for clinic-based studies, and from $5.4\%$ to $11.5\%$ for community samples. Consistent with previous findings that identified memory decline as the initial symptom of AD, amnestic MCI was associated with a higher risk of progression to AD in several longitudinal studies.

In 2011, the NIA-AA convened workgroups to revise diagnostic criteria for AD and its preclinical stages. The NIA-AA core clinical criteria for MCI due to AD were essentially adopted from the Key Symposium criteria, while incorporation of biochemical and neuroimaging biomarkers was recommended in research settings to stratify the level of certainty of underlying Alzheimer’s pathology. During the past 10 years, several biomarkers have been shown to predict the risk of progression to dementia. These risk factors include amyloid-specific biomarkers, neuronal injury biomarkers, and vascular comorbidities. The carrier status

### Table 1 Diagnostic criteria for mild cognitive impairment

<table>
<thead>
<tr>
<th>Core diagnostic criteria</th>
<th>GDS$^{1,2}$</th>
<th>Petersen et al$^{3}$</th>
<th>Key Symposium$^{11}$</th>
<th>NIA-AA 2011$^{15}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subjective cognitive complaint</td>
<td>+ Memory</td>
<td>+ Memory</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Objective cognitive impairment</td>
<td>Memory; &gt;1 SD below average</td>
<td>Memory; &gt;1.5 SD below average</td>
<td>$\geq 1$ cognitive domain; no recommended cut-off</td>
<td>$\geq 1$ cognitive domain; 1–1.5 SD below average</td>
</tr>
<tr>
<td>Preserved general cognitive performance</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Preserved functional independence</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Role of biomarkers</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>Incorporated in research criteria</td>
</tr>
</tbody>
</table>

Notes: $-$, criterion not required; $+$, required criterion.

Abbreviations: GDS, global deterioration scale; SD, standard deviation; NIA-AA, National Institute on Aging-Alzheimer’s Association.
of the apolipoprotein E (APOE) ε4 allele, which is the most important genetic risk factor for AD, has been associated with a more rapid conversion to AD in subjects with MCI. Abnormal CSF biomarkers, including lower β amyloid 1–42 (Aβ42), higher phosphorylated tau (P-tau), and higher total tau (t-tau) were observed in MCI patients who later converted to AD or other dementia compared with non-converters. Amyloid PET is another pathology-specific biomarker, and its positivity has been shown to predict later conversion to AD in subjects with MCI. Among biomarkers for neuronal injury, atrophy of medial temporal structures and hypometabolism in the temporo-parietal cortices were shown to predict a more rapid cognitive decline for subjects with MCI. Consistent with the risk stratification strategy proposed by the NIA-AA criteria, the combination of biomarkers for amyloid pathology and neuronal injury better predicted conversion to AD. In addition to biomarkers for Alzheimer’s pathology and neurodegeneration, the association between modifiable vascular risk factors and cognitive deterioration in MCI has gained considerable interest in recent years. Epidemiological studies showed an inconsistent association between metabolic syndromes and the incidence of MCI. Meanwhile, several studies showed a higher conversion rate from MCI to AD or other types of dementia in subjects with diabetes or other vascular risk factors. In addition, well-controlled diabetes, hypertension, and dyslipidemia were associated with a decreased risk of conversion to dementia. Whether vascular pathology is simply a comorbid condition with AD or is involved with the neurodegenerative process of AD is a burgeoning area of research.

**Subjective cognitive decline: moving toward a pre-MCI stage**

By defining MCI, recruited subjects could be objectively diagnosed at the earliest stage of cognitive impairment; however, randomized controlled trials for MCI patients have failed to find pharmacological treatments that are consistently effective at delaying cognitive decline. To stop the disease process, disease-modifying therapy may need to be initiated even before the onset of MCI. Prior to demonstrable cognitive impairment, many patients experience a subjective decline in memory or other cognitive domains. The subjective decline, even at the stage of normal cognitive performance, is associated with an increased risk of positive biomarkers for Alzheimer’s pathology and later conversion to dementia. Subjects with subjective cognitive decline may be a reasonable target for therapeutic trials. Variable terminology has been used in previous studies to describe this pre-MCI stage, including subjective cognitive impairment, subjective memory impairment, subjective memory impairment, and memory complaints. To generate comparability across studies, a consensus on the terminology and research criteria is crucial. In 2012, the Subjective Cognitive Decline Initiative (SCD-I) formed a working group to generate a common concept and terminology for SCD. The term “subjective cognitive decline” was suggested and was defined as a self-experienced persistent decline in cognitive capacity in comparison with the subject’s previously normal status, during when the subject had normal age-, gender-, and education-adjusted performance on standardized cognitive tests. While preclinical AD can account for the subjective decline in cognition, many different physical and mental conditions can influence self-experienced cognitive fitness in the elderly. Several community and clinic-based epidemiology studies have demonstrated the association between cognitive complaint and depression, anxiety, and personality traits. Other comorbidities that may contribute to cognitive complaints include physical health, sleep problems, and concurrent medication use.

As an unspecific syndrome with multiple possible underlying etiologies, SCD cannot be considered equal to a prodromal phase of AD. Subjects with both cognitive complaints and concurrent AD-associated pathological changes will be a better target for testing potential therapeutic agents, as they may be at a higher risk of further cognitive decline. The SCD-I working group used the term “SCD plus” to describe the following high-risk features: a subjective decline in memory, onset of SCD within the last 5 years, >60 years of age at SCD onset, concerns (worries) associated with SCD, feelings of worse performance than others in the same age group, confirmation of cognitive decline by an informant, presence of the APOE ε4 genotype, and biomarker evidence for AD. These proposed high-risk features were selected based on our current knowledge of AD, but they need further confirmation in longitudinal studies. A subjective complaint that is associated with concerns or supported by an informant may indicate a more significant decline from the baseline condition. In a community-based study that enrolled subjects aged ≥75 years, Jessen et al reported a twofold increase in the risk of developing AD or any dementia for SCD subjects with worries, compared with those without worries. Confirmation of cognitive decline by an informant was not included as core criteria for SCD proposed by SCD-I. However, some studies showed that cognitive decline reported by an informant or a mutual report from both the informant and the subject,
may better predict conversion to MCI or AD, compared with self-report. For subjects that later converted to MCI, Caselli et al observed that self-endorsed cognitive decline developed approximately 30 months earlier than that reported by the informants. The stronger predictive effect of an informant report may be partially explained by its association with a more advanced disease stage. Applying the SCD-plus criteria proposed by SCD-1, Fernandez-Blazquez et al showed a significantly higher risk of developing MCI in the following 13 months among subjects with SCD-plus (18.9%; adjusted HR=4.2), compared with SCD alone (5.6%).

**Incorporating preclinical AD into subjective cognitive decline: strength and bias**

Biomarkers, including CSF profiles of Aβ42 and tau, amyloid PET positivity, atrophy of medial temporal structures, and hypometabolism in the temporo-parietal cortices, have been shown to predict a more rapid progression to dementia in subjects with MCI. However, the associations between these biomarkers and long-term cognitive outcomes in SCD subjects were not strong. Published longitudinal studies accessing the predictive value of AD biomarkers for cognitive outcome in SCD subjects are summarized in Table 2. As shown in subjects with MCI, APOE ε4 allele carrier status is associated with an increased risk of conversion to MCI and AD in subjects with SCD. CSF level of Aβ42 and tau protein were the most studied AD biomarkers. While van Harten et al reported that low Aβ42 was the strongest predictor of conversion to MCI or AD, Hessen et al and Rolstad et al found that high t-tau rather than Aβ42 was correlated with future decline in memory and executive functions. Visser et al failed to find a significant association between abnormal Aβ42/t-tau ratios and conversion to dementia, but only one SCD subject developed dementia during the 2.5-year follow-up period for that study. Neuroimaging biomarkers, including visual rating of hippocampal atrophy and white matter hyperintensities, gray matter atrophy pattern, diffusion tensor imaging, and hypometabolism on FDG PET, were evaluated as predictors of cognitive decline in a few studies. Scheef et al reported a significant association between hypometabolism in the right precuneus and verbal episodic memory decline 35 months later. The same study group later reported that a gray matter atrophy pattern

**Table 2 Longitudinal studies for biomarkers and risk of cognitive decline in subjective cognitive decline subjects**

<table>
<thead>
<tr>
<th>References</th>
<th>Profile</th>
<th>Definition</th>
<th>Duration</th>
<th>Endpoint</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>van Harten et al 2013</td>
<td>n=127,</td>
<td>mean age 60,</td>
<td>SC and 1.5 SD</td>
<td>4 years</td>
<td>MCI or AD</td>
</tr>
<tr>
<td></td>
<td>F 64%, E4 33%</td>
<td></td>
<td></td>
<td></td>
<td>CR 10%; low Aβ42 (HR=16); high t-tau (HR=2.8); high p-tau (HR=2.6)</td>
</tr>
<tr>
<td>Visser et al 2009</td>
<td>n=60,</td>
<td>mean age 66,</td>
<td>SC and 1.5 SD</td>
<td>2.5 years</td>
<td>AD or non-Alzheimer's dementia; Aβ42/t-tau (ns)</td>
</tr>
<tr>
<td></td>
<td>F 48%, E4 53%</td>
<td></td>
<td></td>
<td></td>
<td>CR 0% to AD, 3% to non-Alzheimer's dementia; Aβ42/t-tau (ns)</td>
</tr>
<tr>
<td>Sierra-Rio et al 2016</td>
<td>n=55,</td>
<td>mean age 66,</td>
<td>SC and 1.5 SD</td>
<td>42 vs 34 months</td>
<td>MCI or AD</td>
</tr>
<tr>
<td></td>
<td>F 73%, E4 25%</td>
<td></td>
<td></td>
<td></td>
<td>CR 55% vs 18% (abnormal Aβ42/p-tau ratio; OR=27.1; pooled data from 55 SCD and 94 MCI subjects</td>
</tr>
<tr>
<td>Hessen et al 2015</td>
<td>n=122,</td>
<td>mean age 62.5,</td>
<td>SC and 1.3 SD</td>
<td>2 years</td>
<td>Decline of M or E for 0.5 SD</td>
</tr>
<tr>
<td></td>
<td>F 53%</td>
<td></td>
<td></td>
<td></td>
<td>Tau predicts M decline (P=0.046); Aβ42 (ns)</td>
</tr>
<tr>
<td>Rolstad et al 2013</td>
<td>n=82,</td>
<td>mean age 66,</td>
<td>SC and GDS stage 2</td>
<td>2 years</td>
<td>Decline of M, E, VS, V, or WM</td>
</tr>
<tr>
<td></td>
<td>F 54%</td>
<td></td>
<td></td>
<td></td>
<td>Tau predicts E decline (r²=0.07, P=0.03); Aβ42 (ns)</td>
</tr>
<tr>
<td>Scheef et al 2012</td>
<td>n=27,</td>
<td>mean age 67,</td>
<td>Worries (+), informant (+), and 1.5 SD</td>
<td>36 months</td>
<td>Decline of M or E</td>
</tr>
<tr>
<td></td>
<td>F 42%, E4 33%</td>
<td></td>
<td></td>
<td></td>
<td>PET hypometabolism at right precentral predicts M decline (P=0.029); MRI hippocampal gray matter (ns)</td>
</tr>
<tr>
<td>Peter et al 2014</td>
<td>n=26,</td>
<td>mean age 60,</td>
<td>Worries (+), informant (+), and 1.5 SD</td>
<td>34 months</td>
<td>Decline of M or E</td>
</tr>
<tr>
<td></td>
<td>F 75%, E4 29%</td>
<td></td>
<td></td>
<td></td>
<td>MRI AD gray matter pattern predicts M decline (P=0.12)</td>
</tr>
<tr>
<td>Selnes et al 2013</td>
<td>n=11,</td>
<td>mean age 61,</td>
<td>SC and GDS stage 2</td>
<td>2–3 years</td>
<td>MCI or AD; decline of MMSE</td>
</tr>
<tr>
<td></td>
<td>F 73%, E4 55%</td>
<td></td>
<td></td>
<td></td>
<td>CR 27% to MCI, 45% to AD; DTI and t-tau predict cognitive decline and medial temporal lobe atrophy (11 SCD and 43 MCI)</td>
</tr>
<tr>
<td>Hong et al 2015</td>
<td>n=129,</td>
<td>mean age 66,</td>
<td>SC and 1.0 SD</td>
<td>3.6 years</td>
<td>MCI or AD</td>
</tr>
<tr>
<td></td>
<td>F 65%, E4 29%</td>
<td></td>
<td></td>
<td></td>
<td>CR 22%; MRI visual rating of hippocampal atrophy and WMH (ns)</td>
</tr>
</tbody>
</table>

Notes: SD is used to define normal cognitive performance. +, criterion required to define study population.

Abbreviations: Aβ42, amyloid 1–42; AD, Alzheimer’s disease; CR, conversion rate; DTI, diffusion tensor imaging; E4, executive function; E4, apolipoprotein E ε4 allele carrier; F, female; GDS, Global Deterioration Scale; HR, hazard ratio associated with biomarkers; M, memory; MCI, mild cognitive impairment; MMSE, Mini-Mental State Examination; ns, nonsignificant; MRI, magnetic resonance imaging; OR, odds ratio associated with biomarkers; PET, positron emission tomography; p-tau, phosphorylated tau181; SC, subjective complaint; SCD, subjective cognitive decline; SD, standard deviation; t-tau, total tau; V, verbal function; VS, visuospatial function; WM, working memory; WMH, white matter hyperintensity.
similar to that observed in AD was associated with episodic memory decline.73 Hong et al tested visual rating scores for hippocampal atrophy and white matter hyperintensities to predict conversion to MCI and AD, but failed to show a significant association.79 In summary, current evidence suggests that low CSF Aβ42 and high t-tau or p-tau may help predict cognitive decline in SCD subjects, while the evidence for neuroimaging biomarkers is limited. The low conversion rate to dementia reported by previous studies reflects the early preclinical nature of SCD. Future studies should use longer follow-up periods or apply more sensitive measures for cognitive decline to establish the validity of biomarkers for cognitive prediction in SCD.

While incorporation of biomarkers for preclinical AD may help specify SCD subjects carrying an increased risk of progression to dementia, this approach is based on the hypothetical model of neurodegeneration following the amyloid cascade.46 In recent years, emerging evidence suggests that some subjects with subtle cognitive decline or MCI have biomarker profiles inconsistent with the hypothetical model for preclinical AD. These subjects, referred to as “suspected non-AD pathophysiology (SNAP),” present with neurodegeneration but are negative for biomarkers of amyloid pathology.78 Petersen et al analyzed MCI subjects from The Mayo Clinic Study of Aging cohort and found that 29% of MCI subjects exhibited neurodegeneration without amyloid deposition.79 In addition, these MCI subjects with neurodegeneration-only biomarker profiles had an increased risk of progression to AD, similar to subjects with both amyloid and neurodegeneration biomarkers. Alternatively, Landau et al extracted data from MCI and AD patients with negative florbetapir-PET from the Alzheimer’s Disease Neuroimaging Initiative cohort and found that these patients were less likely to be APOE ε4-positive, had less AD-specific hypometabolism, and had better longitudinal cognitive performance.80 These reports of conflicting biomarker profiles suggest a neurodegeneration pathway independent of amyloid pathology, or underlying non-AD pathology. More longitudinal studies will be needed to determine the long-term cognitive outcomes in amyloid-negative subjects. Studies using biomarkers as inclusion criteria may embed a selection bias that excludes SCD subjects with conflicting biomarker profiles.

Measurement of subjective changes
Various questionnaires have been used in previous studies to measure subjective changes in cognitive capacity. The SCD-I working group reviewed cognitive self-report measurements used in previous studies and found significant heterogeneity among studies.81 Measurements differ in response options, duration of reference timeframe, cognitive items assessed, and item specificity. Many measures were used only in a single study,82–84 while the Memory Complaint Questionnaire (MAC-Q)85 and the Everyday Cognition Scale (ECog)86 were the most commonly used assessments across different studies. Most of these questionnaires were developed recently and lack validation between different populations and cultures. In addition, very few studies have addressed the consistency and compatibility of different questionnaires.87 The SCD-I working group offered several recommendations for measurement selection.81 Measures should be validated for the target population, simple and easy to understand, inquire more about the cognitive issues that older adults encounter in their daily lives, sample cognitive domains beyond episodic memory, and have a specific and narrow reference period. Researchers should also use caution when comparing outcomes between studies using different measurements.

Conclusion and prospective
During the past two decades, the clinical spectrum of AD has been extended to include MCI and even further extended to include SCD. Extending the disease spectrum to the early stages reflects the need to identify and introduce disease-modifying therapy before an irreversible degenerative process occurs. While the concept of SCD was developed to broaden the AD spectrum, it also includes a broad range of underlying etiologies, including non-AD dementia, mood problems, and physical health conditions. The application of biomarkers to patients with SCD may help specify individuals with underlying Alzheimer’s pathology. Meanwhile, SCD can be applied to non-AD dementia and may provide an opportunity to study how different pathological processes interact to influence cognitive outcomes.

Disclosure
The authors report no conflicts of interest in this work.

References
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