Diagnosing Mild Cognitive Impairment (MCI) in clinical trials: a systematic review

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ABSTRACT

Objective: To describe how criteria for amnestic Mild Cognitive Impairment (aMCI) have been operationalised in randomised controlled clinical trials (RCTs).

Design: Systematic review.

Information sources: EMBASE, PubMed and PSYCHINFO were searched from their inception to February 2012. Electronic clinical trial registries were also searched (February 2012).

Study selection: RCTs were included where participant selection was made using Petersen et al-defined aMCI. There was no restriction on intervention type or the outcome tested.

Data extraction: For each trial, we extracted information on study design, demographics, exclusion criteria and the operationalisation strategy for the five aMCI diagnostic criteria including: (1) memory complaint, (2) normal general cognitive function, (3) memory impairment, (4) no functional impairment and (5) no dementia.

Results: 223 articles and 278 registered trials were reviewed, of which 22 met inclusion criteria. Various methods were applied for operationalising aMCI criteria resulting in variability in participant selection. Memory complaint and assessment of general cognitive function were the most consistently measured criteria. There was large heterogeneity in the neuropsychological methods used to determine memory impairment. It was not possible to assess the impact of these differences on case selection accuracy for dementia prediction. Further limitations include selective and unclear reporting of how each of the criteria was measured.

Conclusions: The results highlight the urgent need for a standardised approach to map aMCI. Lack of uniformity in clinical diagnosis, however, is not exclusively a problem for MCI but also for other clinical states such as dementia including Alzheimer’s disease, Lewy Body, frontotemporal or vascular dementia. Defining a uniform approach to MCI classification, or indeed for any classification concept within the field of dementia, should be a priority if further trials are to be undertaken in the older aged population based on these concepts.

ARTICLE SUMMARY

Article focus
- Accurate identification of individuals at risk of dementia or with predementia is important for clinical trial enrolment.
- Diagnosis of predementia is usually made using the amnestic form of Mild Cognitive Impairment (aMCI). While specific criteria for implementation exist, there is no operationalisation protocol.
- Research Question: How have criteria for aMCI been operationalised in randomised controlled clinical trials?

Key messages
- Various methods have been applied for operationalising aMCI criteria in randomised controlled clinical trials resulting in variability in participant selection.
- The results highlight the urgent need for a standardised approach to map aMCI.
- Lack of specific methods for clinical diagnosis is not a problem unique to the field of MCI. Across studies there continues to be inconsistency in the instruments and methodology used to diagnose Alzheimer’s disease and Vascular Dementia, including its prodromal stage, Vascular Cognitive Impairment no Dementia. Revision of diagnostic criteria should be a research priority.

Strengths and limitations of this study
- The review focuses on predementia defined using aMCI. However, not all clinical trials on predementia cognitive states have used this definition of MCI.
- We chose to focus on aMCI as this is one of the commonly applied definitions in clinical and research practice.

INTRODUCTION

As new preventative strategies for dementia are developed, methods to select persons accurately for clinical trial involvement will be needed. In this perspective, Mild Cognitive Impairment (MCI), an intermediate state between normal ageing and dementia, has
become a focus for trials to prevent or delay progression to Alzheimer’s disease. The expectation is that positive results are more likely to be achieved with earlier treatment initiation. While several different definitions exist for MCI, amnestic Mild Cognitive Impairment (aMCI) as defined by Petersen et al. is often used in clinical and research practice. However, despite being commonly applied, no standardised method for the operationalisation of each of the five component criteria (figure 1) necessary for an aMCI diagnosis exists, resulting in heterogeneity in diagnostic methods and case ascertainment across studies. Indeed, there are numerous possibilities for the measurement of the five criteria as highlighted in figure 1. The lack of an established diagnostic methodology for identifying cases for clinical trial enrolment is problematic as study-specific participant selection raises questions regarding the nature of the sample selected, while also making cross study comparison and generalisability of findings difficult.

We undertook a systematic review to explore the methods used to classify aMCI cases, defined using Petersen et al. criteria, in randomised controlled clinical trials (RCTs). The focus was on inclusion criteria and variation in the operationalisation of each of the five aMCI component criteria as outlined in figure 1.

**METHODS**

This review has been undertaken with adherence to the PRISMA statement. The review protocol is available on request.

**Search strategy**

EMBASE (including MEDLINE) and PSYCHInfo were searched using the following keywords and using Medical Subject Heading terms: (‘mild cognitive impairment’ OR MCI) AND (‘randomised controlled trial’ OR ‘randomised controlled trial’ OR RCT). Articles were searched from inception to 6 June 2011, with the search updated on 21 February 2012. Web-based searches, using the term ‘mild cognitive impairment’, were also undertaken in the ISRCTR trial registry (http://www.controlled-trials.com) and on www.clinicaltrials.gov (17 February 2012). Only studies that were published in English were included. Two investigators (BS and TM) independently searched publications using the following inclusion criteria: (1) the study was a RCT; (2) the trial had been completed (was not ongoing or terminated) and results published; (3) the authors report selecting participants using the definition of aMCI as reported in Petersen et al., and could include single or multidomain aMCI subtypes (amendments to criteria were allowed as long as they were stated and Petersen et al was referenced) and (4) the MCI group was analysed separately to the dementia or control groups. The protocol paper or the first publication reporting the primary outcome was selected in case of multiple publications using the same study sample. Titles and abstracts were searched first, followed by the full text of any identified articles. Reviews were also retained and the reference lists of these and each included paper were interrogated. Disagreements were resolved by consensus. Data quality was not assessed as all included studies were RCTs.

**Data extraction**

Data on the lead author, date of publication, study design (country, site, sampling framework, duration and intervention), demographics (age and gender distributions), trial exclusion criteria, dementia progression rates, outcomes tested and the methods used to operationalise each of the five component criteria for the diagnosis of aMCI were abstracted by two investigators (EP and TM) and checked by a third (MS).

**RESULTS**

A total of 223 articles were identified from the literature search. From the electronic search, 11 trials were identified from the ISRCTR trial registry and 267 from http://www.clinicaltrials.gov. Based on the title-abstract

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**Figure 1** Petersen criteria for amnestic Mild Cognitive Impairment.
search, 84 articles were identified for full text review. In total, 22 articles met inclusion criteria and were retained for this review. Figure 2 shows the selection process using the PRISMA (2009) Flow Diagram. As shown in Figure 2, articles were mainly excluded as the sample did not appear to be defined using the Petersen et al criteria or had inadequate details to support the use of Petersen et al criteria (e.g., only stated an objective cognitive deficit), or the article was a review. Online supplementary table S1A summarises the general characteristics, demographics and outcomes tested in each included article. Online supplementary table S1B summarises the operationalisation protocol used for identifying aMCI cases in each trial.

Trial exclusion criteria varied, but were mainly related to cerebrovascular and cardiovascular disease or health and psychiatric-related conditions that could be associated with cognitive decline. There were also differences in the population sampled (clinic vs community), site (single vs multicentre), duration (e.g., 90 days to 4 years) and sample demographics (e.g., age range: 50–90 years). Interventions included pharmacological agents and supplementation \(^6^{–17}\) (including: donepezil,
galantamine, rofecoxib, fluoxetine, lithium treatment, oestrogen treatment (E2), vitamin supplementation (E and B) and supplementation with ω-3 polyunsaturated fatty acids, arachidonic and docosahexaenoic acids), insulin therapy, physical activity (eg, aerobic exercise), cognitive training/rehabilitation programmes (eg, memory training, strategy learning) and combined therapies including cholinesterase inhibitor use combined with a cognitive training programme and physical activity combined with vitamin B supplementation. Outcomes varied extensively across studies and included assessment of cognitive function (in all studies either as a primary or secondary outcome, with no neuropsychological assessment applied consistently) in addition to non-cognitive measures (eg, vascular health such as blood pressure, quality of life, depression, cerebrospinal fluid (CSF) biomarkers of Alzheimer’s disease pathology and neuroimaging). Only five studies reported dementia progression rates, all of which varied: 16%/year, 5–6%/year, 24% over 1 year, 11.9% over a 24-week trial and 15% over 4 years. Most results were negative.

Operationalising MCI component criterion

Two studies did not report details of the operationalisation protocol for defining MCI.

Criterion 1: memory complaint

Five studies reported no details on how memory complaint was obtained. The memory complaint was obtained from the subject in four studies, whereas 11 studies utilised subject report and informant corroboration. One study gave unclear details on who had reported the complaint. In one study, this criterion was operationalised using a history of gradual onset and slow progressive decline in cognitive function, but how this was reported, for example, the subject or informant, was not stated. Three studies used specific scales rather than a single question to assess memory complaint. Smith et al used four items from the Cambridge Examination for Mental Disorders. Rapp et al used the Memory Functioning Questionnaire, which is a 64-item questionnaire assessing memory problems and the use of mnemonics. Van Uffelen et al used a positive response to a single item ‘Do you have memory complaints?’ or answering ‘sometimes’ at least twice on the cognition scale of Strawbridge.

Criterion 2: general cognitive function

This criterion was the one most consistently measured and was typically operationalised using the Mini Mental State Examination (MMSE), score either alone or in combination with other measures including: a structured interview with the patient and informant, the Dementia Rating Scale-II (DRS-II), the Mattis DRS (total score), the Telephone Interview for Cognitive Status (TICS), the Clinical Dementia Rating (CDR) score or the Alzheimer’s Disease Assessment Scale-Cognitive Subscale, in addition to the Clinician Interview-Based Impression of Change (CIBIC). One study used only the CDR score of 0.5.

The cut-off chosen for the MMSE varied from 23 to 26. Most studies used a cut-off value of ≥24, but ≥26, ≥23, ≤25 or a score adjusted for age/education, were also used. In one study, the protocol was modified during recruitment and the cut-off was adjusted from 24–30 to 24–28. One study used a 12-Item shortened MMSE with a cut-off score of ≥7. Three studies specified the use of the MMSE but did not report a cut-off score. Six studies did not specify operationalisation of this criterion.

Criterion 3: object memory decline

Five studies did not specify operationalisation of this criterion. Numerous different tests were used to assess cognition as shown in online supplementary table S2. In addition to inconsistency in test selection, there was no consistency in impairment severity (eg, 1SD, 1.5SDs or 2SDs below the mean). Further, it was not always stated whether cut-off scores for impairment were adjusted for age, education or premorbid ability. In one study, severity was adjusted from 1.5SDs below the mean (used in the first 6 months) to 1SD below the mean during the course of screening. Based on the nature of the objective deficit, three studies reported the inclusion of single or multidomain aMCI. One study reported the use of combined amnestic and non-amnestic (single and multidomain) cases.

In terms of non-memory performance, one study reported that this was tested and required to be unimpaired (defined using a cut-off >10th percentile). Another reported that performance was required to be relatively normal in non-memory domains. In one study, division of cases was unclear; the objective deficit in this study was defined as impairment on a total score comprising five domains (immediate and delayed memory, visuospatial/construction, language and attention) assessed using the Repeatable Battery for the Assessment of Neuropsychological Status.

Criterion 4: Activities of Daily Living (ADL)/Instrumental Activities of Daily Living (IADL)

Seven studies did not specify operationalisation of this criterion. In 11 studies, minimal or non-significant functional impairment was allowed. One study required that in MCI cases that had an MMSE score between 23 and 25, cognitive impairments did not significantly interfere with daily activities or social functioning, determined by a caregiver report. This restriction was not required in MCI cases with a MMSE score ≥26.

Functional impairment tended to be assessed by a self-report or informant report of the difficulty with ADLs or Basic ADLs. Specific scales were used for functional
assessment in some studies to include the Functional Autonomy Measurement System (SMAFQ), the Blessed Dementia Rating Scale (BDRS)-CERAD version, the Groningen Activity Restriction Scale and selected items from the Lawton and Katz scales or items from the Cambridge Behavioural Inventory (CBI). In only two studies did it appear that no evidence of any functional impairment was allowed; one, based on five items, related to ADLs from the CBI, and another specified no decline in ADLs without their measurement being specified.

Criterion 5: dementia diagnosis

Three studies did not specify operationalisation of this criterion. Fourteen studies used the Diagnostic and Statistical Manual (DSM-III-R/IV-TR/-IV), National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer’s Disease and Related Disorders Association (NINCDS-ADRDA) criteria or National Institute of Neurological Disorders and Stroke and Association Internationale pour la Recherche et l’Enseignement en Neurosciences (NINCDS-AIREN) criteria. Two studies used the CDR score, and one each used a self-report of a diagnosis, clinical judgement and the TICS combined with a MMSE score <24.

Additional measures

In some studies, additional measures, generally related to the assessment of global functioning (such as the CDR sum of boxes score) or dementia severity (eg, from none, mild, moderate and severe), were made in parallel to the mapping of the five aMCI criteria. For example, two studies administered the DRS, seven the CDR, one the BDRS, one the CIBIC, and one the Global Deterioration Scale (GDS). One study also had informants complete both the Informant Questionnaire on Cognitive Decline in the Elderly (IQCODE-Short form) and EuroQol (EQ-5D), a measure of health status.

DISCUSSION

This review highlights the lack of consistency in MCI case ascertainment in currently completed RCTs. How MCI was diagnosed was not always reported or clear, and varying operationalisation protocols make it impossible to determine similarity across the samples recruited in the different trials. A priority for clinical trial research is to agree on a uniform set of criteria to operationalise MCI. The recruitment protocols identified in this review could provide the basis for future work to determine best practice (eg, in terms of testing classification accuracy of the different methods used), in order to inform the development of a consistent recruitment methodology for MCI clinical trials.

The review highlights the continuing challenge of operationalising the current Petersen et al definition of aMCI. Without a standard operationalisation protocol for defining aMCI cases clinical trial recruitment will continue to be variable. Indeed, within the field of dementia, there is a lack of consistency in operationalisation protocols not only for aMCI, but its associated disorders (eg, Cognitive Impairment no Dementia), dementia and its subtypes (such as Alzheimer’s disease, Lewy Body dementia, frontotemporal dementia and vascular dementia), pre-MCI and other predementia states such as VCIND. For some dementias and their related conditions, it may however be difficult and unrealistic to have one set of operational criteria, precise assessment instruments or cut-off values. For example, a single set of criteria may not be possible for defining symptom fluctuations (eg, as seen in Lewy Body dementia), capturing variability in symptom profiles (eg, the different type of aphasic deficit presented in frontotemporal dementia) or reflecting differences in neuropsychological profiles (eg, for vascular dementia and VCIND, the type and location of vascular damage may result in variable symptom profiles). Different diagnostic criteria for MCI affect prevalence and progression. Similarly, for dementia, different criteria have been found to affect prevalence. Inconsistency in case classification for any health condition, whether it is within the field of dementia or any other disease category, can have an impact on research and trial recruitment and outcomes.

With regard to aMCI, consensus needs to be reached on five core issues relating to the measurement of each of the component criteria. First, whether memory complaint should be self-reported and/or informant reported and how it should be assessed (eg, single or multiple items). Second, how global cognitive function should be assessed with possible measures including the MMSE, CDR and GDS, and what the best cut-off score is (within and across cultures). Third, which neuropsychological test(s) should be used to assess memory, what should be the severity of cognitive impairment (1SD, 1.5SD) and whether covariate adjustment is needed. In addition, there is the question of whether both memory and non-memory domains should be tested. The possible tests identified in this review are outlined in online supplementary table S2. Fourth, how functional performance should be assessed (the type of questions), the nature of the task (eg, instrumental ADLs, basic ADLs), reporting (eg, patient, informant or clinician) and what is the maximum level of impairment (eg, none, mild, moderate or severe difficulty or significant difficulty in some areas but not in others). Fifth, how should dementia be defined for exclusion with examples used including: the DSM or NINCDS-ADRDA criteria, the CDR sum of boxes score ≥1 or via screening instruments (eg, the Telephone Screening Instrument). It should be noted that aMCI is not always operationalised as originally specified (eg, permissible significant functional impairment in some studies), and consensus needs to be reached on whether all five criteria are necessary. Further, whether modifications (if any) to criteria can be made and the
implications of making modifications, for example, in terms of dementia predictability and effect on generalisability, needs to be established.

Decision also needs to be reached on the best treatment target. The impairment captured in aMCI is not always progressive, with a proportion of cases reverting to normal or remaining stable at follow-up, particularly when mapped in population-based studies. Indeed, symptoms of MCI are not always a consequence of Alzheimer’s pathology, but rather can have multiple aetiologies such as depression or vascular disease, each with different outcomes (eg, dementia progression, improvement with treatment for the underlying health symptoms). Better methods are needed to determine the underlying cause of disease in this patient group to accurately identify those individuals whose MCI is associated with Alzheimer’s disease. One possibility could be defining aMCI as in the Alzheimer’s Disease Cooperative Study trial (based on a subjective memory complaint, MMSE score, impaired performance on the Logical Memory II Subscale, no functional impairment and a CDR score of 0.5) as strict implementation of this methodology has been found to result in a consistent rate of dementia progression (approximately 16% /year) across studies, including the multicentre Alzheimer’s Disease Neuroimaging Initiative. Further research is needed to test this method of operationalisation across cohorts (clinical and population based; across countries) and calculate prevalence and longitudinal course in order to determine the generalisability of these findings. Such results could have important implications in terms of identifying a standard protocol for all future aMCI clinical trials.

A recent task force on designing trials in early (prodromal) Alzheimer’s disease (AD) argues for the use of aMCI criteria in combination with biomarkers to improve case selection for clinical trials. Suggestions for possible biomarkers have included hippocampal or whole brain atrophy, CSF Aβ42 levels, PiB imaging, genetic screening (APOE e4 status) or behavioural deficits as each has been associated with dementia. Further, how dementia and AD are defined is currently undergoing revision, with the aim of improved stratification of patients. Where MCI now sits in the ever changing ‘lexicon’ of AD (ie, given there is currently no concrete border between preclinical and clinical disease) will have implications for who is targeted for clinical trial recruitment. For example, MCI, as defined by Petersen et al criteria, may no longer be considered at-risk, but as already AD and encompassed in the new-term ‘prodromal’; an early symptomatic stage where a patient shows evidence of memory impairment and positive ratings on pathophysiological and topographical markers of AD. Clinical trial research may therefore shift some focus to asymptomatic at-risk stages (eg, pre-MCI) where individuals are biomarker positive for AD but are otherwise healthy. However, like aMCI efforts are needed to standardise criteria and develop an operational protocol for any new stage of disease (eg, prodromal AD and pre-MCI) and to undertake validation across settings including oldest-old age groups and populations (vs clinical samples).

The review should be viewed in light of some limitations. First, we chose to focus on Petersen et al defined aMCI, as this is one of the commonly applied definitions in clinical and research practice. However, not all trials on preclinical cognitive states have used this definition of MCI, with some studies defining intermediate cognitive states using simply a MMSE score or using criteria that have made refinements to the original aMCI criteria. The main change has been in the acceptable level of functional impairment: from none to allowing minor problems, particularly in complex activities such as, for example, account keeping. Different definitions of MCI have different prevalence estimates and also vary in their risk of dementia progression (eg, more extensive patterns of cognitive changes have been associated with greater progression of MCI to dementia). Subtypes have also been defined depending on the neuropsychological profile including amnestic and non-amnestic single or multidomain MCI, and multidomain combined MCI that includes both memory and non-memory deficits. Which, if any, of the many different criteria and subtypes should be adopted in RCTs, or whether no distinction should be made between MCI and AD during recruitment requires further discussion.

CONCLUSION

Much work needs to be done on the characterisation of individuals at risk of dementia for clinical trial recruitment. Within this framework, attention is being focused on redefining the earliest stages of disease and generating new definitions of what constitutes ‘prodromal/pre-dementia’ and ‘at-risk’. Standardisation in the definition and development of an operational protocol will result in improvements in diagnosis and clinical trial methodology.

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