Practice guideline update summary: Mild cognitive impairment

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Abstract

Objective
To update the 2001 American Academy of Neurology (AAN) guideline on mild cognitive impairment (MCI).

Methods
The guideline panel systematically reviewed MCI prevalence, prognosis, and treatment articles according to AAN evidence classification criteria, and based recommendations on evidence and modified Delphi consensus.

Results
MCI prevalence was 6.7% for ages 60–64, 8.4% for 65–69, 10.1% for 70–74, 14.8% for 75–79, and 25.2% for 80–84. Cumulative dementia incidence was 14.9% in individuals with MCI older than age 65 years followed for 2 years. No high-quality evidence exists to support pharmacologic treatments for MCI. In patients with MCI, exercise training (6 months) is likely to improve cognitive measures and cognitive training may improve cognitive measures.

Major recommendations
Clinicians should assess for MCI with validated tools in appropriate scenarios (Level B). Clinicians should evaluate patients with MCI for modifiable risk factors, assess for functional impairment, and assess for and treat behavioral/neuropsychiatric symptoms (Level B). Clinicians should monitor cognitive status of patients with MCI over time (Level B). Cognitively impairing medications should be discontinued where possible and behavioral symptoms treated (Level B). Clinicians may choose not to offer cholinesterase inhibitors (Level B); if offering, they must first discuss lack of evidence (Level A). Clinicians should recommend regular exercise (Level B). Clinicians may recommend cognitive training (Level C). Clinicians should discuss diagnosis, prognosis, long-term planning, and the lack of effective medicine options (Level B), and may discuss biomarker research with patients with MCI and families (Level C).
Mild cognitive impairment (MCI) is a condition in which individuals demonstrate cognitive impairment with minimal impairment of instrumental activities of daily living (IADL). Although MCI can be the first cognitive expression of Alzheimer disease (AD), it can also be secondary to other disease processes (i.e., other neurologic, neurodegenerative, systemic, or psychiatric disorders). The term amnestic MCI (aMCI) describes a syndrome in which memory dysfunction predominates; in nonamnestic MCI, impairment of other cognitive features (e.g., language, visuospatial, executive) is more prominent.

This practice guideline updates a 2001 American Academy of Neurology (AAN) practice parameter with recommendations concerning the diagnosis and treatment of MCI. The guideline focuses on presumed idiopathic or neurodegenerative MCI—particularly relating to AD—that may be secondary to other disease processes or Parkinson disease—MCI or vascular cognitive impairment, as these may have different epidemiologic and treatment spectra than AD. This article summarizes the guideline findings, conclusions, and recommendations. The full text of the guideline, including appendices e-1 through e-8, is available as supplemental data (links.lww.com/WNL/A125), as are tables e-1 through e-3 (links.lww.com/WNL/A34) and references e1–e50 (links.lww.com/WNL/A50).

The guideline addresses 4 questions:

1. What is the prevalence of MCI in the general population?
2. What is the prognosis for patients diagnosed with MCI for progression to a diagnosis of dementia, and how does this compare with an age-matched general population?
3. What pharmacologic treatments are effective for patients diagnosed with MCI?
4. What nonpharmacologic treatments are effective for patients diagnosed with MCI?

This guideline does not review the rapidly evolving field of biomarker research in MCI; the guideline panel determined that this should be the subject of a future guideline or systematic review. In addition, the potential psychological distress of a diagnosis of MCI (which has been discussed in the literature) was not one of the questions reviewed by the expert panel for this guideline.

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Glossary

AAN = American Academy of Neurology; AD = Alzheimer disease; aMCI = amnestic mild cognitive impairment; CI = confidence interval; CIND = cognitively impaired, no dementia; FDA = Food and Drug Administration; IADL = instrumental activities of daily living; MCI = mild cognitive impairment; RR = relative risk.

Description of the analytic process

This practice guideline principally follows the methodologies described in the 2004 edition of the AAN’s guideline development process manual. Conclusions and recommendations were developed in accordance with the process outlined in the 2011 guideline development process manual, as amended to include the updated scheme for classifying therapeutic articles. The complete guideline provides a description of the exact methodology followed, including the processes of convening the author panel, performing the literature search, and reviewing the evidence. In accordance with the 2011 guideline manual, recommendations were based not only on the evidence in the systematic review, but also on strong related evidence, established principles of care, and inferences. The level of obligation for each recommendation was based on the strength of these premises and the risk–benefit ratio of following the recommendation, with adjustments based on importance of outcomes, variation in patient preferences, feasibility/availability, and patient costs. Consensus was determined by a modified Delphi voting process in accordance with prespecified rules.

The panelists noted that various definitions of MCI and of related terms, such as cognitively impaired, no dementia (CIND), were used in the reviewed literature. Variation was based on different ascertainment methods, different neuropsychological measures, different measure thresholds, and requirements for different cognitive deficits. There was also variation in the use of aMCI and nonamnestic MCI in these studies. To address these discrepancies, the panelists reflected the specific definition used for a study where feasible in the evidence synthesis tables and guideline text, and provided specific comments on the potential effect of differing definitions.

Analysis of evidence

What is the prevalence of MCI in the general population?

Background

Various definitions of MCI have been used over time, reflecting an evolution of thought from primarily focusing on amnesia to including other cognitive deficits. Because memory deficits are the clinical hallmark of AD, some groups used criteria for MCI that required the presence of memory deficits in isolation (e.g., aMCI), and others included a broader definition that included either single-domain...
nonamnestic deficits or deficits in multiple cognitive domains, either with memory impairment (multidomain aMCI) or without (multidomain nonamnestic MCI). The definition of MCI is also affected by the psychometric properties of, and norms for, the tests used to identify thresholds between normal aging and MCI. Table e-1 (links.lww.com/WNL/A34) presents the characteristics of various definitions of MCI used in the literature evaluated here. Table e-2 shows the effect on frequency of MCI in the population when less or more stringent MCI criteria were applied.

**Analysis**

Twenty Class I studies and 14 Class II studies were identified. Eight of the Class I studies showed that a lower education level was significantly associated with a higher prevalence of MCI. Two of the Class I studies indicated that male sex was associated with the presence of MCI, but other studies found similar baseline prevalence in men and women.

A random-effects meta-analysis using Class I and II studies confirmed an increased prevalence with cohort age. The all-studies estimate for individuals aged 60–64 years was 6.7% (95% confidence interval [CI] 3.4%–12.7%, I² 11.0); for those aged 65–69, 8.4% (95% CI 5.2%–13.4%, I² 0); for ages 70–74, 10.1% (95% CI 7.5%–13.5%, I² 5.2); for ages 75–79, 14.8% (95% CI 10.1%–21.1%, I² 60.7); and for ages 80–84, 25.2% (95% CI 16.5%–36.5%, I² 0) (see table e-3, links.lww.com/WNL/A34).

**Conclusions**

MCI is common in older populations, and its prevalence increases with age (high confidence, multiple Class I and Class II studies, consistent meta-analysis) and lower educational level (high confidence, multiple Class I studies).

What is the prognosis for patients diagnosed with MCI for progression to a diagnosis of dementia, and how does this compare with an age-matched general population?

**Analysis**

Nine Class I studies evaluated prognosis for individuals with MCI, all showing an increased risk of progression to dementia when participants with MCI were compared with age-matched participants without MCI. A random-effects meta-analysis demonstrated that the cumulative incidence for the development of dementia in individuals with MCI/CIND older than age 65 followed for 2 years was 14.9% (95% CI 11.6%–18.1%, I² 0). In those with MCI/CIND vs age-matched participants at 2–5 years after, the relative risk (RR) of dementia (all types) was 3.3 (95% CI 2.5–4.5, I² 4.9); the RR of the diagnosis of AD was 3.0 (95% CI 2.1–4.8, I² 17.3).

Reversion to normal cognition in individuals with MCI

Four Class I studies showed reversion to normal cognition on follow-up in 14.4%, 33.3%, 19%, and 38% of participants with MCI. However, 2 studies documented increased rates of ultimate conversion to dementia in participants with MCI who reverted to normal cognition, suggesting that individuals who revert remain at a higher risk of progression back to MCI or dementia than individuals who have never received an MCI diagnosis (in these studies, 65% and 55% ultimately converted to dementia).

**Conclusions**

Persons with MCI are at higher risk of progressing to dementia than age-matched controls (high confidence, multiple concordant Class I studies, meta-analysis). Persons diagnosed with MCI may remain stable, return to neurologically intact, or progress to dementia (multiple Class I studies, 14.4%–55.6% reverting to normal).

What pharmacologic treatments are available for patients diagnosed with MCI, and are these treatments effective on cognitive measures of progression to dementia, excluding other symptomatic effects?

**Analysis**

One Class I study, 10 Class II studies described in 9 publications, and 3 Class III studies addressed pharmacologic treatment of MCI. Table 1 describes the available studies and conclusions for each pharmacologic intervention. Comprehensive descriptions of each study, including effect sizes and CIs, are available in the full-length guideline (links.lww.com/WNL/A125).

What nonpharmacologic treatments are effective for patients diagnosed with MCI?

**Analysis**

Two Class II studies were reviewed that used exercise as an intervention in individuals with MCI, and 1 Class II and 4 Class III studies investigated the use of various cognitive interventions. Table 2 describes the available studies and conclusions for each nonpharmacologic intervention; details are provided in the full-length guideline (links.lww.com/WNL/A125).

Putting the evidence into clinical context

Care for persons with cognitive impairment meeting various MCI criteria continues to evolve, with the area of biomarker research changing particularly rapidly. Even in the context of an evolving field, clinicians can provide high-quality care focusing on counseling, treatment, and comorbidity management. Where clinicians are not proficient in caring for the cognitive or behavioral/psychiatric needs of persons with MCI, referral to appropriate specialists is an important part of the treatment paradigm in line with the following recommendations.
Table 1  Evidence and conclusions for pharmacologic treatments for mild cognitive impairment (MCI)

<table>
<thead>
<tr>
<th>Agent</th>
<th>Classification of evidence</th>
<th>Conclusion</th>
</tr>
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<tbody>
<tr>
<td>Donepezil</td>
<td>3 Class II studies (Petersen 2005,e10 Doody 2009,e11 Salloway 2004,e12)</td>
<td>In patients with MCI, donepezil use over 3 years is possibly ineffective for reducing the chances of a progression to possible or probable Alzheimer dementia (low confidence in the evidence, single Class II study [Petersen 2005,e10]). In patients with MCI, it is unknown whether donepezil slows progression on various cognitive scales (very low confidence in the evidence based on 2 Class II studies with limited precision and small magnitude of effect) (Doody 2009,e11 Salloway 2004,e12). Study CIs could not exclude an important effect and the ADAS-Cog change was statistically significant but not clinically meaningful.</td>
</tr>
<tr>
<td>Galantamine</td>
<td>2 Class II studies (Winblad 2008,e14 both studies reported in 1 article)</td>
<td>In patients with MCI, galantamine use over 24 months is probably ineffective for reducing progression to dementia (moderate confidence in the evidence based on 2 Class II Studies).</td>
</tr>
<tr>
<td>Rivastigmine</td>
<td>1 Class II study (Feldman 2007,e14)</td>
<td>In patients with MCI, rivastigmine use up to 48 months is possibly ineffective for reducing the rate of progression to possible or probable Alzheimer dementia (low confidence in the evidence based on a single Class II study).</td>
</tr>
<tr>
<td>Flavonoid-containing drink</td>
<td>1 Class II study (Desideri 2012,e15)</td>
<td>In patients with MCI, there is insufficient evidence to support or refute the cognitive benefits of a drink with high-dose flavonoids (about 390 mg) on an integrated measure (cognitive z score) of overall cognitive function at 8 weeks (very low confidence in the evidence based on a single Class II study with CIs including unimportant effects; evidence of a dose response was also unclear).</td>
</tr>
<tr>
<td>Homocysteine-lowering B vitamins</td>
<td>1 Class II study (Smith 2010,e16)</td>
<td>In patients with MCI, there is insufficient evidence to support or refute the use of homocysteine-lowering therapies in patients with MCI (very low confidence in the evidence based on a single Class II study with decreased confidence in the evidence owing to use of a primary endpoint with unclear clinical significance).</td>
</tr>
<tr>
<td>Transdermal nicotine patch</td>
<td>1 Class I study (Newhouse 2012,e17)</td>
<td>Six months of transdermal nicotine (15 mg/d) use possibly improves cognitive test performance but not Clinical Global Impression of Change in patients with amMCI who do not smoke (low confidence in the evidence based on 1 Class I study with decreased confidence in the evidence owing to unclear clinical significance of the outcome of hit reaction time).</td>
</tr>
<tr>
<td>Piribedil</td>
<td>1 Class III study (Nagaraja 2001,e18)</td>
<td>Data are insufficient to support or refute an effect of piribedil on cognitive measures in MCI (very low confidence in the evidence based on 1 Class III study).</td>
</tr>
<tr>
<td>Rofecoxib*</td>
<td>1 Class II study (Thal 2005,e19)</td>
<td>Rofecoxib possibly increases the risk of progression to AD in patients with MCI (low confidence in the evidence based on 1 Class II study).</td>
</tr>
<tr>
<td>Tesamorelin injections</td>
<td>1 Class II study (Baker 2012,e20)</td>
<td>In patients with MCI, treatment with tesamorelin injections over 20 weeks is possibly effective to improve performance on various cognitive measures (low confidence in the evidence based on 1 Class II study).</td>
</tr>
<tr>
<td>V0191</td>
<td>1 Class III study (Dubois 2012,e21)</td>
<td>Data are insufficient to support or refute an effect of V0191 use on ADAS-Cog response rates in patients with MCI (very low confidence in the evidence based on 1 Class III study).</td>
</tr>
<tr>
<td>Vitamin E</td>
<td>1 Class II study (Petersen 2005,e22)</td>
<td>In patients with MCI, use of vitamin E 2,000 IU daily is possibly ineffective for reducing progression to AD (low confidence in the evidence based on a single Class II study).</td>
</tr>
<tr>
<td>Vitamin E + vitamin C</td>
<td>1 Class III study (Naeini 2014,e23)</td>
<td>In patients with MCI, combined use of oral vitamin E 300 mg and C 400 mg daily over 12 months is of uncertain efficacy (very low confidence in the evidence based on 1 Class III study).</td>
</tr>
</tbody>
</table>

Abbreviations: AD = Alzheimer disease; ADAS-Cog = Alzheimer’s Disease Assessment Scale–cognitive subscale; amMCI = amnestic mild cognitive impairment; CI = confidence interval.
References cited here can be found in the e-references (links.lww.com/WNL/A50) for the guideline summary article.
* Rofecoxib was removed from the market worldwide in September 2004. There are no data on whether other anti-inflammatory medications are effective or harmful in patients with MCI.
+ It is unclear from this study whether this effect is sustained beyond 20 years.

Practice recommendations
Section A: Recommendations for assessing for MCI

Recommendation A1
Rationale
Appropriate diagnosis of MCI is important because MCI becomes increasingly common as individuals age and is associated with an increased risk of progression to dementia, suggesting that this condition reflects a pathologic disease state rather than normal cognitive aging. Appropriate diagnosis of MCI is important in order to assess for reversible causes of cognitive impairment, to help patients and families understand the cause of their cognitive concerns, and to discuss the prognostic possibilities with the provider so they can plan accordingly, although sharing the diagnosis must be balanced with the potential harm of anxieties from diagnosing a patient with a condition that may not progress. Ascribing cognitive symptoms to normal aging without an assessment for MCI may
result in failure to assess for reversible causes of cognitive impairment or to provide patients and families with an accurate diagnosis that may affect life choices, or both. Although subjective cognitive complaints alone are insufficient to diagnose MCI, such complaints from either patients or their close contacts are core to most major MCI diagnostic criteria, as they may reflect a change in cognitive function.

Recommendation
For patients for whom the patient or a close contact voices concern about memory or impaired cognition, clinicians should assess for MCI and not assume the concerns are related to normal aging (Level B).

Recommendation A2
Rationale
In the United States, the Medicare Annual Wellness Visit requires an assessment to detect cognitive impairment. Subjective cognitive complaints alone can result in over-diagnosis or under-diagnosis of MCI and thus are insufficient to screen for MCI. Clinicians assessing for cognitive impairment should use a brief, validated cognitive assessment instrument in addition to eliciting patient and informant history regarding cognitive concerns.

Recommendation
When performing a Medicare Annual Wellness Visit, clinicians should not rely on historical report of subjective memory concerns alone when assessing for cognitive impairment (Level B).

Recommendation A3
Rationale
When screening or assessing for MCI, validated assessment tools should be used. Various instruments have acceptable diagnostic accuracy for detecting MCI, with no instrument being superior to another. Because brief cognitive assessment instruments are usually calibrated to maximize sensitivity rather than specificity, patients who test positive for MCI should then have further assessment (e.g., more in-depth cognitive testing, such as neuropsychological testing with interpretation based on appropriate normative data) to formally assess for this diagnosis. Diagnosis of MCI is based ultimately on a clinical evaluation determining cognitive function and functional status and not solely on a specific test score.

Recommendation
For patients for whom screening or assessing for MCI is appropriate, clinicians should use validated assessment tools to assess for cognitive impairment (Level B). For patients who test positive for MCI, clinicians should perform a more formal clinical assessment for diagnosis of MCI (Level B).

Recommendation A4
Rationale
In the presence of cognitive impairment, clinicians need to distinguish between a diagnosis of MCI and one of dementia, although the boundary is not always clear. Diagnosing dementia prematurely can lead to negative consequences for patients and families. Only a proportion of patients with MCI will proceed to dementia. In patients with cognitive

Table 2 Evidence and conclusions for nonpharmacologic treatments for mild cognitive impairment (MCI)

<table>
<thead>
<tr>
<th>Agent</th>
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</tr>
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<tr>
<td>Exercise</td>
<td>2 Class II studies (Nagamatsu 2012,e22 Suzuki 2013,e23)</td>
<td>In patients with MCI, treatment with exercise training for 6 months is likely to improve cognitive measures (moderate confidence in the evidence based on 2 Class II studies).</td>
</tr>
<tr>
<td>Cognitive interventions</td>
<td>1 Class II (Kinsella 2009,e24) and 4 Class III studies (Kinsella 2016,e25 Tsolaki 2011,e26 Nakatsuka 2015,e27 Lam 2015,e28)</td>
<td>There is insufficient evidence to support or refute the use of any individual cognitive intervention strategy (1 Class II study with results that are not statistically significant and with suspected imprecision, 4 Class III studies, each examining a different cognitive intervention strategy). When various cognitive interventions are considered as a group, for patients with MCI, cognitive interventions may improve select measures of cognitive function (low confidence in the evidence based on 1 Class II study with insufficient precision [Kinsella 2009,e24], 1 Class III study showing improvements in strategy knowledge, internal strategy use, and well-being but not external strategy or memory [Kinsella 2016,e25], 1 Class III study [Tsolaki 2011,e26] showing improvement on multiple cognitive measures, 1 Class III study [Nakatsuka 2015,e27] showing improvement on the MMSE but with some limitations, and 1 Class III study [Lam 2015,e28] showing no differences when all patients with MCI are considered, but with improvements in the integrated cognitive-physical training groups when considering the ADAS-Cog, fluency, and recall in patients with single-domain MCI and fluency in patients with multidomain MCI).</td>
</tr>
</tbody>
</table>

Abbreviations: ADAS-Cog = Alzheimer’s Disease Assessment Scale–cognitive subscale; MMSE = Mini-Mental State Examination.

References cited here can be found in the e-references (links.lww.com/WNL/A50) for the guideline summary article.
impairment, clinicians must carefully assess for evidence of functional impairment limiting independence in daily activities (e.g., by taking a careful history from the patient and a close contact), a requirement for all dementia diagnoses, to help distinguish between MCI and dementia. With a specific inquiry about functional impairment, clinicians may also identify dementia in patients when patients and family are less forthcoming about functional problems.

**Recommendation**

For patients with MCI, clinicians should assess for the presence of functional impairment related to cognition before giving a diagnosis of dementia (Level B).

**Recommendation A5**

**Rationale**

Diagnoses of MCI and dementia have important implications for patients and families. Appropriate diagnosis is important for informing evaluation for underlying causes, counseling on long-term prognosis, and recommending therapeutic strategies. Clinicians in many disciplines can have experience in caring for individuals with cognitive impairment, including family practice, geriatrics, internal medicine, neurology, psychiatry, and psychology. When clinicians without experience in cognitive impairment identify patients for whom there is a concern of MCI, they should refer these patients to a specialist with experience in cognition for further evaluation.

**Recommendation**

For patients suspected to have MCI, clinicians who lack the necessary experience should refer these patients to a specialist with experience in cognition (Level B).

**Recommendation A6**

**Rationale**

Although MCI is a high-risk state for progression to dementia, some patients with MCI remain stable and some improve. Some cases of MCI are associated with reversible causes of cognitive impairment, including medication side effects, sleep apnea, depression, and other medical conditions. Patients with MCI should undergo a medical evaluation for MCI risk factors that may be treatable.

**Recommendation**

For patients diagnosed with MCI, clinicians should perform a medical evaluation for MCI risk factors that are potentially modifiable (Level B).

**Recommendation A7**

**Rationale**

Because patients with MCI can improve, remain stable, or progress cognitively, identifying biomarkers that can stratify risk is expected to be particularly important for prognosis. The use of biomarkers in patients with MCI is a rapidly evolving field but to date, there are no biomarkers clearly shown to predict progression in patients with MCI.

**Recommendation A7a**

For patients and families asking about biomarkers in MCI, clinicians should counsel that there are no accepted biomarkers available at this time (Level B).

**Recommendation A7b**

For interested patients, clinicians may discuss the option of biomarker research or refer patients, or both, if feasible, to centers or organizations that can connect patients to this research (e.g., subspecialty centers, Trial Match, ClinicalTrials.gov) (Level C).

**Recommendation A8**

**Rationale**

Although MCI has no approved pharmacologic management, there are US Food and Drug Administration (FDA)–approved agents for treatment of Alzheimer dementia further emphasizing the importance of assessing for a change in cognitive status over time.

**Recommendation**

For patients diagnosed with MCI, clinicians should perform serial assessments over time to monitor for changes in cognitive status (Level B).

**Section B: Recommendations for management of MCI**

**Recommendation B1**

**Rationale**

Some patients with MCI improve or remain stable rather than progress. In addition, some cases of MCI are associated with reversible causes of cognitive impairment, including medication side effects, general medical conditions, sleep disturbance, and depression. Because these risk factors are treatable and have implications of their own, weaning patients from use of cognitively impairing medications where feasible and treating risk factors that may contribute to cognitive impairment should be the first steps in managing MCI, particularly because symptomatic treatment options are limited for impaired cognition.

**Recommendation**

For patients diagnosed with MCI, clinicians should wean patients from medications that can contribute to cognitive impairment (where feasible and medically appropriate) and treat modifiable risk factors that may be contributing (Level B).
Recommendation B2

Rationale
There are no FDA-approved medications for the treatment of MCI. Moreover, there are no high-quality, long-term studies identifying pharmacologic or dietary agents that either improve cognition or delay progression in patients with MCI.

Recommendation
For patients diagnosed with MCI, clinicians should counsel the patients and families that there are no pharmacologic or dietary agents currently shown to have symptomatic cognitive benefit in MCI and that no medications are FDA-approved for this purpose (Level B).

Recommendation B3

Rationale
Studies of cholinesterase inhibitors showed no benefit on cognitive outcomes or reduction in progression from MCI to dementia, although some studies could not exclude an important effect. In addition to lacking efficacy, side effects of cholinesterase inhibitors are common, including gastrointestinal symptoms and cardiac concerns.

Recommendation B3a
For patients diagnosed with MCI, clinicians may choose not to offer cholinesterase inhibitors (Level B).

Recommendation B3b
If clinicians choose to offer cholinesterase inhibitors, they must first discuss with patients the fact that this is an off-label prescription not currently backed by empirical evidence (Level A).

Recommendation B4

Rationale
Clinical trials provide an opportunity for interested patients to participate in identifying or testing new treatment options, which is of particular importance when no pharmacologic options are available.

Recommendation
For patients diagnosed with MCI who are interested in pharmacologic treatment, clinicians may inform these patients of centers or organizations that can connect patients to clinical trials (e.g., subspecialty centers, Trial Match, ClinicalTrials.gov) (Level C).

Recommendation B5

Rationale
Although long-term studies are unavailable, 6-month studies suggest a possible benefit of twice-weekly exercise for cognition in MCI. Exercise also has general health benefits and generally limited risk.

Recommendation
Clinicians should assess for behavioral and neuropsychiatric symptoms in MCI and treat with both pharmacologic and nonpharmacologic approaches when indicated (Level B).
Recommendation B8

Rationale

In patients with MCI, cognitive interventions may be beneficial in improving measures of cognitive function. It is good practice to offer nonmedication approaches to care.

Recommendation

In patients with MCI, clinicians may recommend cognitive interventions (Level C).

Suggestions for future research

The guideline panel recommends (1) the use of consistent diagnostic criteria for MCI and dementia in clinical trials, to improve the ability to apply and combine results; (2) the inclusion of patient cohorts with specific biomarker data in treatment studies targeted at specific pathologies (e.g., MCI due to AD); (3) the use of outcome measures that are direct measures of clinically meaningful patient outcomes (i.e., development of dementia, reduction of ability to undertake activities of daily living or IADL, patient or caregiver [or both] quality of life measures) or surrogate markers that have previously been shown to have a strong correlation with such measures; (4) standardized reporting of trial design in publications using CONSORT criteria; and (5) study of MCI thought to be secondary to AD and MCI related to other pathologies (e.g., vascular MCI, MCI related to Lewy body pathology); and (6) further study of early lifestyle and comorbidity modifications and the effects of such changes on the progression of MCI to different dementia subtypes.

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Conflict of interest

The American Academy of Neurology (AAN) is committed to producing independent, critical, and truthful clinical practice guidelines (CPGs). Significant efforts are made to minimize the potential for conflicts of interest to influence the recommendations of this CPG. To the extent possible, the AAN keeps separate those who have a financial stake in the success or failure of the products appraised in the CPGs and the developers of the guidelines. Conflict of interest forms were obtained from all authors and reviewed by an oversight committee prior to project initiation. The AAN limits the participation of authors with substantial conflicts of interest. The AAN forbids commercial participation in, or funding of, guideline projects. Drafts of the guideline have been reviewed by at least 3 AAN committees, a network of neurologists, Neurology peer reviewers, and representatives from related fields. The AAN Guideline Author Conflict of Interest Policy can be viewed at aann.com. For complete information on this process, access the 2004 AAN process manual.

Author contributions

Dr. Petersen: study concept and design, acquisition of data, analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content, study supervision. Dr. Lopez: study concept and design, analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content. Dr. Armstrong: analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content, study supervision. T.S.D. Getchius: study concept and design, study supervision. Dr. Ganguli: study concept and design, analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content. Dr. Gloss: analysis or interpretation of data, study supervision. Dr. Marson: study concept and design, analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content. Dr. Pringsheim: analysis and interpretation of data, study supervision. Dr. Day: analysis and interpretation of data, study supervision. Dr. Sager: study concept and design, analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content. Dr. Stevens: study concept and design, analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content. Dr. Rae-Grant: analysis or interpretation of data, drafting/revising the manuscript, critical revision of the manuscript for important intellectual content.
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