Evidence-based Clinical Practice Guideline for Deprescribing Cholinesterase Inhibitors and Memantine in People with Dementia

Recommendations

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Disclaimer:
This document is a general guide, to be followed subject to the clinician’s judgement and the person’s preference in each individual case. The guideline is designed to provide information to assist decision making and is based on the best evidence available at the time of developing this publication.

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Plain English Summary

Dementia describes a syndrome that is characterised by a progressive loss in cognition, function and behaviour [1]. Worldwide, the number of people living with dementia is increasing every year [2]. There are currently two classes of medications available to treat the symptoms of dementia: cholinesterase inhibitors (ChEIs: donepezil, rivastigmine and galantamine) and the N-methyl-D-aspartate (NMDA) receptor antagonist, memantine [3]. These medications can have important benefits to people with dementia and their carers.

All medications come with the potential for benefits as well as risks, and these risks and benefits can change over time, such as during long-term use. Therefore, appropriate use of ChEIs and memantine involves both prescribing these medications to individuals who are likely to benefit, and deprescribing (withdrawing) them for individuals where the risks outweigh the benefits. However, deprescribing itself also has the potential for both benefit and harm to the individual. Thus, the purpose of this guideline is to assist healthcare professionals (particularly prescribers) to determine when it might be suitable to trial withdrawal of these medications for an individual. These recommendations only apply to individuals already taking one of the described medications (donepezil, rivastigmine, galantamine and/or memantine).

The main points of this guideline are as follows:

- A proportion of people who have used these medications for over 12 months or outside an approved indication may be able to stop the medication with minimal clinically relevant negative consequences. Discontinuation of ChEIs and/or memantine may lead to a worsening of cognitive function in certain populations of users. The limited data on person-important outcomes, such as quality of life and function, suggest that these outcomes may not be altered by discontinuation. However, there is considerable uncertainty in the benefits and harms of both prescribing and deprescribing in the individual.

- It is important to consider the values, preferences and experiences of the person with dementia and/or their carer/family when determining if trial deprescribing is appropriate. Carers have expressed fears associated with medication discontinuation, and individuals may feel that deprescribing is ‘giving up’ or a signal that they are no longer worth treating. Good communication between clinicians and people with dementia and/or carers/family on the benefits and harms of continuing versus discontinuing, in the context of their values and preferences, is necessary when discussing a potential trial of deprescribing.

- ChEIs and memantine have been found to be cost-effective in treating approved indications in some populations and settings, based on the data from short-term studies. The cost implications of deprescribing may include reduced medication costs, reduced costs of
treating adverse drug effects, and an uncertain benefit or cost if there is a change in function that increases or decreases health service utilisation. Further research is required in this area.

- There are numerous clinical considerations when deprescribing ChEIs and/or memantine, including how to assess for ongoing benefit, how to conduct withdrawal and monitoring (plus actions to follow monitoring), and implementation of non-pharmacological management strategies.
Executive Summary

We followed the process of developing class-specific deprescribing guidelines [4], which are based on a comprehensive checklist for successful guideline development (Guideline 2.0) [5] and the AGREE-II criteria [6]. We also incorporated the requirements for the Australian National Health and Medical Research Council (NHMRC) external guideline approval [7]. This process involves a systematic review and uses the GRADE process to assess the quality of the evidence and convert the evidence into recommendations (see Methods). Developing the recommendations involved considering the quality of the evidence, the risks and benefits of deprescribing, the risks and benefits of continuation, consumer values and preferences, and economic considerations (see individual sections, plus Appendix 2 in main guideline document).

The recommendations below are classed as one of three possible types of recommendations: Evidence-based Recommendations (EBR), Consensus-based Recommendations (CBR) or Practice Points (PP). In this guideline, we employ CBR, which are recommendations based on a systematic review where there is limited or low-quality evidence, as well as PP, which are recommendations outside the scope of the systematic review based on expert opinion and non-systematically reviewed evidence.

Each recommendation contains a rating of the quality of the evidence and strength of the recommendations. The recommendations below are rated as based on low- or very low-quality evidence. The major limitations to the quality of the evidence were a high risk of bias and a lack of generalisability (for details, see Appendix 2 in main guideline document). We have rated the strength of the recommendations as ‘Strong’. A strong recommendation is provided when all or most individuals would be best served with that course of action, and the outcomes align with their values and preferences. A weak recommendation reflects that consideration of the individual’s values and preferences and treatment goals is required before proceeding with the recommended course of action (such as the individual’s preference on competing interests). There is considerable heterogeneity in the population of people with dementia in terms of both their condition and their values and preferences. The rating of strong is primarily based on the evidence presented (despite its low quality) and a reasonable judgement of the limited potential for harm in a carefully monitored trial of discontinuation.

This document is a general guide. Implementation of recommendations should only be conducted by qualified/trained personnel in consultation with appropriate parties (such as the prescriber, family, nurses and care staff). The people involved in these parties for consultation will vary by setting and should be considered in the local context considering scopes of practice of health care professionals.
Recommendations

NB: this is not a treatment guideline—the recommendations below should not be applied to assist in the decision to initiate medication. They should not be used to dissuade against prescribing these medications or as reason to prescribe them.

The recommendations below apply to adults who have already been prescribed and have been regularly taking a ChEI and/or memantine for a sufficient amount of time at the maximum tolerated dose. These recommendations are to ‘trial deprescribing’.

**Trial deprescribing** refers to slowly reducing the medication dose (tapering) prior to complete cessation, with monitoring throughout the process. If the person has a noticeable decline after dose reduction/cessation (after exclusion of other causes), then the medication should be restarted at the previous minimum effective dose.

These recommendations should be considered in the context of the individual. People with dementia vary in their condition (such as progress, age of onset, symptom profile and aetiology), overall health state (such as comorbidities, polypharmacy, frailty and life expectancy), values, preferences and treatment goals. It is also important to consider their previous response to the medication. Improvement, stabilisation and even reduced rate of decline in cognition can all be considered benefits of treatment, and this can have an important impact on the person with dementia and their family. However, it is very difficult to quantify the ongoing benefit of long-term use in the individual. Trial withdrawal may help identify individuals who are still benefiting from the medication. Decisions surrounding deprescribing should be conducted as shared decision making with the person with dementia and/or their family/carer, ensuring that they are informed of the likely potential benefits and harms of both continuing and discontinuing these medications. Other healthcare professionals may need to be consulted to determine the appropriateness to trial withdrawal, or to ensure monitoring is conducted throughout the process. Application of these recommendations may need to be adapted depending on the context in which they are used, i.e. depending on the health care organisation, and professionals involved.

We present these recommendations for clinicians to consider within the context of each individual:

**PP:** Deprescribing of cholinesterase inhibitors and/or memantine should be a trial discontinuation, with periodic monitoring (such as every four weeks) and re-initiation of the medication if the individual evidences clear worsening of condition after withdrawal.

**PP:** The dose of the cholinesterase inhibitors and/or memantine should be tapered prior to discontinuation by halving the dose every four weeks to the lowest available dose, followed by discontinuation.

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CBR: For individuals taking a cholinesterase inhibitor (donepezil, rivastigmine or galantamine) for Alzheimer’s disease, dementia of Parkinson’s disease, Lewy body dementia or vascular dementia for greater than 12 months, we recommend trial discontinuation if:

- cognition and/or function has significantly worsened over the past six months (or less, as per the individual)
- no benefit (improvement, stabilisation or decreased rate of decline) was seen at any time during treatment
- the individual has severe/end-stage dementia.

(Level of evidence: Low; Strength of recommendation: Strong)

CBR: For individuals taking a cholinesterase inhibitor (donepezil, rivastigmine or galantamine) for an indication other than Alzheimer’s disease, dementia of Parkinson’s disease, Lewy body dementia or vascular dementia, we recommend trial discontinuation. (Level of evidence: Low; Strength of recommendation: Strong).

CBR: For individuals taking memantine for Alzheimer’s disease, dementia of Parkinson’s disease or Lewy body dementia for greater than 12 months, we recommend trial discontinuation if:

- cognition and/or function has significantly worsened over the past six months (or less, as per the individual)
- no benefit (improvement, stabilisation or decreased rate of decline) was seen at any time during treatment
- the individual has severe/end-stage dementia.

(Level of evidence: Very Low; Strength of recommendation: Strong)

CBR: For individuals taking memantine for indications other than Alzheimer’s disease, dementia of Parkinson’s disease or Lewy body dementia, we recommend trial discontinuation. (Level of evidence: Very Low; Strength of recommendation: Strong).

PP: Other situations in which trial deprescribing of cholinesterase inhibitors and/or memantine can be considered include: a decision by a person with dementia and/or their family/carer to discontinue the medication, a person with dementia’s refusal or inability to take the medication, non-adherence that cannot be resolved, drug–drug or drug–disease interactions that make treatment risky, severe agitation/psychomotor restlessness, and non-dementia terminal illness.
Areas of Major Debate

There has been significant debate between our Guideline Development Team (GDT) members about the need to tailor the recommendations to the individual. Some stakeholder GDT members could recall previous individuals who had been treated and for whom the recommendation would not be appropriate. For example, the lines between underlying causes of dementia (such as Alzheimer’s disease (AD) versus non-AD dementia) are not always clear. There was tension between wishing to add qualifiers and keeping the recommendations clear and straightforward for end-users. To resolve this debate, we have included a preamble to the recommendations to ensure that users of the guideline are focused on the individual and aware of the significant variability among people with dementia. This debate, in turn, led to a discussion about the rating of the strength of the recommendations of ‘Strong’ versus ‘Weak’. According to the GRADE process [8,9]:

[a] strong recommendation [should be made] when ... all or almost all informed people would make the recommended choice for or against an intervention.

[a] weak recommendation [should be made] when ... most informed people would choose the recommended course of action, but a substantial number would not.

Much debate can be held over what constitutes a ‘substantial number’, as we agree that some people with dementia and/or their family/carer may not wish to discontinue the ChEI and/or memantine. However, we feel that the majority of people in the situations outlined by the recommendations who are informed would agree to a trial discontinuation. This is based on assumptions with value based on adopting a ‘less is more’ approach (as outlined in Error! Reference source not found. Error! Reference source not found. and Error! Reference source not found.). The recommendations may also be complicated by the life-limiting nature of dementia and lack of alternative treatments, with significant hope being placed on these medications by people with dementia and their family. It is also important to remember that the strength of the recommendation is based not only on the systematic review evidence, but also on the review of benefits and harms, consumer values and preferences, and economic considerations.

The GDT also encountered tension between wishing to trial discontinuation to determine if the medication is still having a benefit, versus not wishing to ‘rock the boat’ for people with dementia who are otherwise stable. In the absence of external measures to determine if the medication is having a benefit, some clinicians view trial discontinuation as an appropriate measure to determine the need to continue the medication, while others prefer evidence of harm to trigger discontinuation, with avoidance of potential harm from deprescribing more highly valued.
References


