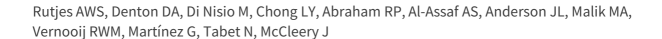


Cochrane Database of Systematic Reviews

Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life (Review)



Rutjes AWS, Denton DA, Di Nisio M, Chong LY, Abraham RP, Al-Assaf AS, Anderson JL, Malik MA, Vernooij RWM, Martínez G, Tabet N, McCleery J.

Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life. *Cochrane Database of Systematic Reviews* 2018, Issue 12. Art. No.: CD011906.

DOI: 10.1002/14651858.CD011906.pub2.

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[Intervention Review]

Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

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Editorial group: Cochrane Dementia and Cognitive Improvement Group. **Publication status and date:** New, published in Issue 12, 2018.

Citation: Rutjes AWS, Denton DA, Di Nisio M, Chong LY, Abraham RP, Al-Assaf AS, Anderson JL, Malik MA, Vernooij RWM, Martínez G, Tabet N, McCleery J. Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life. *Cochrane Database of Systematic Reviews* 2018, Issue 12. Art. No.: CD011906. DOI: 10.1002/14651858.CD011906.pub2.

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ABSTRACT

Background

Vitamins and minerals play multiple functions within the central nervous system which may help to maintain brain health and optimal cognitive functioning. Supplementation of the diet with various vitamins and minerals has been suggested as a means of maintaining cognitive function, or even of preventing dementia, in later life.

Objectives

To evaluate the effects of vitamin and mineral supplementation on cognitive function in cognitively healthy people aged 40 years or more.

Search methods

We searched ALOIS, the Cochrane Dementia and Cognitive Improvement Group's (CDCIG) specialised register, as well as MEDLINE, Embase, PsycINFO, CINAHL, ClinicalTrials.gov and the WHO Portal/ICTRP from inception to 26th January 2018.

Selection criteria

We included randomised controlled trials that evaluated the cognitive effects on people aged 40 years or more of any vitamin or mineral supplements taken by mouth for at least three months.

Data collection and analysis

Study selection, data extraction, and quality assessments were done in duplicate. Vitamins were considered broadly in the categories of B vitamins, antioxidant vitamins, and combinations of both. Minerals were considered separately, where possible. If interventions and outcomes were considered sufficiently similar, then data were pooled. In order to separate short-term cognitive effects from possible longer-term effects on the trajectory of cognitive decline, data were pooled for various treatment durations from 3 months to 12 months and up to 10 years or more.

Main results

In total, we included 28 studies with more than 83,000 participants. There were some general limitations of the evidence. Most participants were enrolled in studies which were not designed primarily to assess cognition. These studies often had no baseline cognitive assessment and used only brief cognitive assessments at follow-up. Very few studies assessed the incidence of dementia. Most study reports did not mention adverse events or made only very general statements about them. Only 10 studies had a mean follow-up > 5 years. Only two studies had participants whose mean age was < 60 years at baseline. The risk of bias in the included studies was generally low, other than a risk of attrition bias for longer-term outcomes. We considered the certainty of the evidence behind almost all results to be moderate or low.

We included 14 studies with 27,882 participants which compared folic acid, vitamin B12, vitamin B6, or a combination of these to placebo. The majority of participants were aged over 60 years and had a history of cardio- or cerebrovascular disease. We found that giving B vitamin supplements to cognitively healthy adults, mainly in their 60s and 70s, probably has little or no effect on global cognitive function at any time point up to 5 years (SMD values from -0.03 to 0.06) and may also have no effect at 5-10 years (SMD -0.01). There were very sparse data on adverse effects or on incidence of cognitive impairment or dementia.

We included 8 studies with 47,840 participants in which the active intervention was one or more of the antioxidant vitamins: ß-carotene, vitamin C or vitamin E. Results were mixed. For overall cognitive function, there was low-certainty evidence of benefit associated with ß-carotene after a mean of 18 years of treatment (MD 0.18 TICS points, 95% CI 0.01 to 0.35) and of vitamin C after 5 years to 10 years (MD 0.46 TICS points, 95% CI 0.14 to 0.78), but not at earlier time points. From two studies which reported on dementia incidence, there was low-certainty evidence of no effect of an antioxidant vitamin combination or of vitamin E, either alone or combined with selenium. One of the included studies had been designed to look for effects on the incidence of prostate cancer; it found a statistically significant increase in prostate cancer diagnoses among men taking vitamin E.

One trial with 4143 participants compared vitamin D3 (400 IU/day) and calcium supplements to placebo. We found low- to moderate-certainty evidence of no effect of vitamin D3 and calcium supplements at any time-point up to 10 years on overall cognitive function (MD after a mean of 7.8 years -0.1 MMSE points, 95% CI -0.81 to 0.61) or the incidence of dementia (HR 0.94, 95% CI 0.72 to 1.24). A pilot study with 60 participants used a higher dose of vitamin D3 (4000 IU on alternate days) and found preliminary evidence that this dose probably has no effect on cognitive function over six months.

We included data from one trial of zinc and copper supplementation with 1072 participants. There was moderate-certainty evidence of little or no effect on overall cognitive function (MD 0.6 MMSE points, 95% CI -0.19 to 1.39) or on the incidence of cognitive impairment after 5 years to 10 years. A second smaller trial provided no usable data, but reported no cognitive effects of six months of supplementation with zinc gluconate.

From one study with 3711 participants, there was low-certainty evidence of no effect of approximately five years of selenium supplementation on the incidence of dementia (HR 0.83, 95% CI 0.61 to 1.13).

Finally, we included three trials of complex supplements (combinations of B vitamins, antioxidant vitamins, and minerals) with 6306 participants. From the one trial which assessed overall cognitive function, there was low-certainty evidence of little or no effect on the TICS (MD after a mean of 8.5 years 0.12, 95% CI -0.14 to 0.38).

Authors' conclusions

We did not find evidence that any vitamin or mineral supplementation strategy for cognitively healthy adults in mid or late life has a meaningful effect on cognitive decline or dementia, although the evidence does not permit definitive conclusions. There were very few data on supplementation starting in midlife (< 60 years); studies designed to assess cognitive outcomes tended to be too short to assess maintenance of cognitive function; longer studies often had other primary outcomes and used cognitive measures which may have lacked sensitivity. The only positive signals of effect came from studies of long-term supplementation with antioxidant vitamins. These may be the most promising for further research.

PLAIN LANGUAGE SUMMARY

Vitamin and mineral supplementation for preventing cognitive deterioration in cognitively healthy people in mid and late life

Review question

This review aimed to find out whether people aged 40 years or older could maintain their mental abilities or reduce their risk of dementia by taking vitamin or mineral supplements.

Background

There is great interest in anything people might do to keep themselves mentally healthy as they age and, especially, to reduce their risk of developing dementia. Various vitamin and mineral supplements have been suggested as ways to do this. Vitamins and minerals have many functions in the body and the ways they might affect brain health are not well understood. Therefore, we were interested in the effects of any vitamins or minerals, either alone or in combination.

What we did

We searched up to January 2018 for all trials which had studied the effect on cognition (thinking and remembering) of any vitamin or mineral supplements in people aged 40 years or more who had no cognitive problems at the beginning of the study. The studies had to include a comparison group which had not taken the supplement being studied and, to be sure the comparison was fair, it should have been decided at random whether people got the supplement or not.

What we found

We grouped the 28 included trials according to the kind of supplement they used and how it might work.

Most of the trials were not originally designed to study cognition or dementia and used only simple measures of cognition. Very few studies investigated whether participants developed dementia. Long-term studies are probably needed to find effects on the risk of dementia or cognitive decline but only 10 studies had an average length of follow-up longer than five years. The studies were generally well-conducted although the longer trials had difficulty following up all of the participants and this could have biased some of the longer-term results.

There were 14 trials of B vitamins (folic acid, vitamin B6, vitamin B12) with nearly 28,000 participants, mainly in their 60s and 70s. Most of these trials were quite short (less than two years). We found no evidence that B vitamins had any effect on cognition.

There were 8 trials of antioxidant vitamins (beta-carotene/vitamin A, vitamin C, vitamin E) with approximately 47,000 participants. These trials tended to be longer than the B vitamin trials so may have had more chance of detecting effects on dementia and cognitive decline. The results were mixed. We found low-certainty evidence of better overall cognitive function after an average of 18 years taking beta-carotene and after five years to 10 years taking vitamin C, but no effects after shorter periods of treatment. There were also small benefits of beta-carotene, vitamin C, and antioxidant combinations on memory at some time points but not others. There was no evidence of any benefits from vitamin E alone. Two studies examined the risk of developing dementia. One found no effect of a combination of antioxidant vitamins and the other found no effect of vitamin E, either alone or combined with the mineral selenium. Most of the studies did not report any information about harmful effects. One included trial was designed to look for an effect on the risk of prostate cancer; it found a higher risk among the men taking vitamin E.

There was a small trial of vitamin D supplements which found they probably had no effect on cognition over six months. There were longer trials of vitamin D with calcium (one trial), zinc and copper (one trial), and complex multivitamins (three trials). All lasted between five and 10 years, but none of them found any evidence of beneficial effects on cognition. One trial found no effect of selenium taken for approximately five years on the risk of developing dementia.

Conclusions

We found no good evidence to suggest that middle-aged or older people can preserve cognitive function or prevent dementia by taking vitamin or mineral supplements. There were a few positive results associated with long-term use of antioxidant vitamins, particularly beta-carotene and vitamin C, although the effects were small. Further research into the effects of these vitamins may be worthwhile.

SUMMARY OF FINDINGS FOR THE MAIN COMPARISON [Explanation]

B Vitamins compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life

Patient or population: cognitively healthy people in mid and late life

Setting: community

Intervention: B Vitamins (folic acid, vitamins B6 and vitamin B12, alone or in combination)

Comparison: placebo

Outcomes	()		Relative effect (95% CI)	№ of participants (studies)	Certainty of the evi- Comments dence (GRADE)
	Risk with placebo	Risk with B Vitamins			(GRADE)
Overall cognitive functioning 3 to 12 months of supplementation (final measurement data) assessed with: MMSE or TICSm		SMD 0 (0.08 lower to 0.08 higher)	-	2556 (3 RCTs)	⊕⊕⊕⊝ MODERATE ¹
Overall cognitive functioning 3 to 12 months of supplementation (change from baseline data) assessed with: MMSE or TICSm		SMD 0.03 lower (0.26 lower to 0.19 higher)	-	317 (2 RCTs)	⊕⊕⊖⊝ LOW ¹²
Overall cognitive func- tioning > 1 to 2 years of supplementation (fi- nal measurement data) assessed with: MMSE, TICS, or TICSm		SMD 0.15 higher (0.17 lower to 0.47 higher)	-	6956 (5 RCTs)	⊕⊕⊕⊝ MODERATE ¹

Overall cognitive func- tioning > 1 to 2 years of supplementa- tion (change from base- line data) assessed with MMSE	•	SMD 0.06 higher (0.2 lower to 0.31 higher)		241 (1 RCT)	⊕⊕⊖⊖ LOW ¹²
Overall cognitive functioning > 2 to 5 years of supplementation (final measurement data) assessed with MMSE, TICS, F-TICSm or a composite score	•	SMD 0.01 lower (0.11 lower to 0.09 higher)		3985 (4 RCTs)	⊕⊕⊕⊖ MODERATE ¹
Overall cognitive func- tioning - > 5-10 years of supplementation (fi- nal measurement data) assessed with MMSE, TICS or TICSm		SMD 0.01 lower (0.09 lower to 0.07 higher)		10249 (3 RCTs)	⊕⊕⊖⊖ LOW ¹³
Incidence of MCI or de- mentia > 2 to 5 years of supplementation assessed with: MMSE < 24 on 2 or more occa-			RR 0.89 (0.67 to 1.18)	1803 (1 RCT)	⊕⊕⊕⊝ MODERATE ¹
sions follow-up: median 2.8 years	103 per 1.000	91 per 1.000 (69 to 121)			

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

TICS: Telephone Interview for Cognitive Status

TICS m: modified Telephone Interview for Cognitive Status

F-TICSm: modified Telephone Interveiw for Cognitive Status (French version)

¹ Downgraded for indirectness due to restricted inclusion criteria for participants in included studies

 $^{^{2}}$ Downgraded for imprecision due to small sample size and wide confidence intervals

³ Downgraded for study limitations due to missing data from 26% of participants in SEARCH 2010 MMSE:Mini Mental State Examination

BACKGROUND

Description of the condition

Cognitive health, mild cognitive impairment, and dementia

There is no consensus definition of cognitive health or successful cognitive ageing, but it has been described as "not just the absence of cognitive impairment, but the development and preservation of the multidimensional cognitive structure that allows the older adult to maintain social connectedness, an ongoing sense of purpose, and the abilities to function independently, to permit functional recovery from illness and injury, and to cope with residual cognitive deficits" (Depp 2012; Hendrie 2006). Successful cognitive ageing is distinct from mild cognitive impairment (MCI) and dementia.

Dementia is a syndrome of cognitive and functional decline that is usually progressive. It affects multiple cognitive domains, among which memory is usually prominent, but orientation, executive function, language, perception, and social cognition are also often affected. In most cases, the onset of dementia is gradual. It is usually preceded by a stage of mild cognitive impairment when cognitive deficits beyond those of normal ageing are detectable, but function is not significantly impaired. The term dementia is used once the cognitive deficits begin to have a significant effect on daily activities. As the syndrome progresses, those affected become increasingly dependent on others for all activities of daily living.

Types of MCI and dementia

Subtypes of dementia are distinguished by the underlying pathology. The four most common subtypes are Alzheimer's disease dementia (AD) (accounting for an estimated 60% to 70% of all dementia cases); vascular dementia (VaD); dementia with Lewy Bodies (DLB); and frontotemporal dementia (FTD). Accurate diagnosis of the subtypes may be difficult. Mixed pathology is common, with more than 80% of cases having some features of Alzheimer's disease (Jellinger 2006; WHO 2012). However, the proportion of dementia attributable to Alzheimer's disease reduces with age (Savva 2009).

There are numerous different operational definitions of MCI, with different scope (e.g. memory or non-memory impairments) (Matthews 2007), prevalence (Stephan 2007), and risk of progression to dementia (Matthews 2008). Currently, there is no single, universally accepted definition of MCI (Stephan 2013), but adaptations of the criteria suggested by Petersen 1999 are commonly used in clinical trials. Categorisation of MCI may also be based on the suspected underlying cause of the cognitive deficits, e.g. MCI due to Alzheimer's disease and MCI due to vascular disease (termed 'vascular cognitive impairment, no dementia': VCIND).

It is increasingly recognised that some dementia syndromes, including AD, have a very long preclinical phase with pathological changes beginning years, or even decades, before the onset of symptoms (Sperling 2011). Attempts have been made to develop criteria to capture preclinical states as early as possible. For example, the label 'pre-MCl' has been applied to people with impaired executive function and language, elevated apathy scores, and reduced volumes of left brain areas of the hippocampus, an area involved in consolidation of information from short-term memory to long-term memory but also spatial memory, enabling orientation. (Duara 2011). It is likely that biomarker tests, tests that measure some biological state, will be needed to identify people in the earliest stages of dementing illnesses.

Prevalence of MCI and dementia

In the UK Medical Research Council's population-based Cognitive Function and Ageing Study (CFAS), which mapped 18 different definitions of MCI, the range of prevalence estimates was found to be variable (0.1% to 42.0%), and conversion rates to dementia generally low (Stephan 2007). Prevalence rates and rates of conversion to dementia have been reported to be higher in specialist settings than in population-based studies (adjusted conversion rate from MCI to dementia 9.6% versus 4.9%) (Mitchell 2009). The risk of dementia increases with age. According to a World Health Organization (WHO) report, only 2% to 10% of cases start before the age of 65 years (WHO 2012). The same report estimated that there were 35.6 million people with dementia in the world in 2010, and that this figure would double every 20 years to reach 65.7 million in 2030 (WHO 2012). However, there is a degree of uncertainty about the expected increase in prevalence of dementia. Recent research by Matthews 2013 in the UK and by Christensen 2013 in Denmark indicates that age-specific prevalence of dementia may be falling in developed countries, suggesting the possibility that dementia risk may be modifiable. Nevertheless, because of population ageing, the overall prevalence continues to rise.

Risk factors

Generally, risk factors for dementia can be divided into modifiable and non-modifiable categories. Non-modifiable risk factors include age, genetic factors, and gender (females are at higher risk). At a population level, age is the most significant risk factor. In people older than 65 years, the risk of AD (the commonest cause of dementia) doubles every five years (Launer 1999; McCullagh 2001; Van den Berg 2012; Van der Flier 2005). A pooled analysis of four prospective studies in Europe found that the incidence of AD among people aged 90 years and over was 63.5/1000 personyears (Launer 1999). Genetics plays a major role in early onset AD, but a lesser role in the much commoner late onset disease. Many potentially modifiable risk factors are common to AD and vascular dementia. These include diabetes mellitus, midlife obe-

sity, midlife hypertension, smoking, and physical inactivity (Prince 2014; WHO 2012). The fact that risk factors are active in midlife is consistent with what is known of the long preclinical phase of dementia and implies that there is a long period in which preventive efforts could be effective.

At present, there is no cure for any subtype of dementia, but identifying and targeting modifiable risk factors might offer opportunities to modify its onset and course. Studies to test this hypothesis have investigated cognitive stimulation, exercise, diet, and other nutritional interventions, and the management of vascular risk factors, such as hypertension, diabetes, obesity, smoking, and physical inactivity (Lindsay 2002; Lourida 2013; Norton 2014; Wilson 2002).

Description of the intervention

This review focusses on the effect of vitamins or minerals (or both) on cognitive functioning. Vitamins are organic compounds that are essential for the normal physiological process in the body and play important roles in growth and development (Kennedy 2011). MInerals are inorganic elements and as nutrients are also needed by the body to grow and develop normally (Centers for Disease Control and Prevention 2014). Most of these essential nutrients are available naturally in the diet. Dietary supplements are any consumed products that aim to provide nutrients additional to those obtained in the regular diet.

How the intervention might work

Vitamins and minerals have numerous important roles in human physiology. Recognised biological roles of all those considered for this review are summarised briefly in Appendix 1.

The most commonly cited mechanisms by which vitamins and minerals may influence cognitive function are (1) via effects on specific plasma amino-acids called homocysteine for B vitamins (folic acid, vitamin B6, and vitamin B12) and (2) via antioxidant effects, reducing the concentration of free radicals that may damage human cells, for vitamins C, E, and β-carotene, possibly vitamin D, and some minerals such as selenium. These hypotheses about possible mechanisms are based largely on preclinical studies and on observations of associations between elevated homocysteine levels or elevated markers of oxidative stress and cognitive decline or dementia (Clarke 2007a, Bennett 2009). Detailed understanding of mechanisms is, however, lacking.

Why it is important to do this review

The prevalence and high societal costs of dementia are such that even small reductions in its age-specific incidence would have a large impact on healthcare costs and on the overall burden it causes to patients, their families, and communities. In order to decide where preventive efforts should be concentrated, robust assessments are needed of the effects of interventions and of the 'dose' and duration of intervention necessary to achieve an effect.

For individuals, fear of cognitive decline and dementia may be powerful motivators to seek preventive interventions. Vitamin and mineral supplementation and cognitive activities (e.g. computerised 'brain training' games), in particular, are subject to promotion by those with commercial interests. It is important for people to know whether time, effort, and money they might invest to prevent cognitive decline is likely to be well spent. Information about adverse effects is also important. Although nutritional and behavioural interventions are often perceived to be 'low risk', they are not necessarily without the potential to cause harm. For example, trials have found high doses of vitamin E to be associated with higher rates of side effects than placebo (Bjelakovic 2012; Brigelius-Flohe 2007; Miller 2005).

OBJECTIVES

To evaluate the effects of vitamin and mineral supplementation on cognitive function in cognitively healthy people aged 40 years or more.

METHODS

Criteria for considering studies for this review

Types of studies

We included randomised or quasi-randomised controlled trials, published or unpublished, reported in any language. We included trials with a duration of intervention of 12 weeks or more. Trials in cognitively healthy people with a duration as short as 12 weeks typically investigate cognitive enhancement rather than maintenance of cognitive function. We included these trials in order to give a full picture of the data, although it is recognised that the relationship between short-term cognitive enhancement and maintenance of cognitive function over longer periods of time, or incidence of dementia, is unclear.

Types of participants

We included trials in which the participants were cognitively healthy people from midlife onwards, i.e. participants aged 40 years and over without a dementia diagnosis or cognitive impairment at baseline ('midlife' was defined for the purposes of disease classification, in DSM-IV 1994, as between 40 years and 65 years). The cognitive status of participants was determined by the

trialists' own definitions of 'cognitively healthy'. These definitions were recorded. We contacted trialists if further clarification was needed to determine health status. If trials included a cognitively mixed population, we required at least 80% of participants to be free of cognitive impairment (including subjective cognitive impairment) at baseline.

Types of interventions

We included studies comparing the effect of orally-administered vitamin or mineral supplements (or vitamin and mineral supplements) with control interventions which were not expected to have an effect on cognition (placebo or no intervention/usual care). Eligible vitamins and minerals are listed in Appendix 1. These vitamins and minerals could be given singly or in combination and at any dose, for a minimum of 12 weeks. Trials of vitamins or minerals (or vitamins and minerals) given in combination with other nutrients (e.g. fatty acids, amino acids) or with drugs were excluded unless the design allowed the effects of vitamins or minerals to be isolated. For example, a trial comparing vitamin X to amino acid Y was ineligible, but a trial comparing vitamin X and amino acid Y to amino acid Y only was eligible.

Types of outcome measures

Primary outcomes

Mean overall cognitive functioning measured with any internationally accepted and validated measure: for example, Alzheimer's Disease Assessment Scale - cognitive subscale (ADAS-cog); Mini Mental State Examination (MMSE); Repeatable Battery for the Assessment of Neuropsychological Status (RBANS); Cambridge Cognition Examination (CAMCOG).

Secondary outcomes

Any internationally accepted and validated measures of:

- specific cognitive functioning subdomain: episodic memory:
- specific cognitive functioning subdomain: executive functioning;
- specific cognitive functioning subdomain: speed of processing;
 - incidence of MCI or all-cause dementia;
 - quality of life, either generic or disease-specific;
- clinical global impression (a scale measuring a clinician's judgement on overall symptom severity);
 - functional performance;
- number of participants experiencing one or more serious adverse events (SAE);
 - mortality.

If a trial provided data on more than one cognitive scale for a specific outcome, we applied a hierarchy of cognition-related outcomes and used data on the cognitive scale that was highest on this hierarchy. For example, if a trial reported results on both the Mini Mental State Examination and the Clinical Dementia Rating scale (CDR), we used the outcome data from MMSE in our quantitative analyses. The rank of a scale in the hierarchy was determined by the frequency of its use in a large set of 79 trials, evaluating vitamin and mineral supplementation, dietary interventions, and physical exercise interventions.

Search methods for identification of studies

Electronic searches

We searched ALOIS (www.medicine.ox.ac.uk/alois), the Cochrane Dementia and Cognitive Improvement Group's (CD-CIG) specialised register.

ALOIS is maintained by the Information Specialist for the CD-CIG, and contains studies that fall within the areas of dementia prevention, dementia treatment and management, and cognitive enhancement in healthy elderly populations. The studies are identified through:

- 1. Monthly searches of a number of major healthcare databases: MEDLINE, Embase, CINAHL, PsycINFO and LILACS;
- 2. Monthly searches of a number of trial registers: ISRCTN; UMIN (Japan's Trial Register); the WHO portal (which covers Clinical Trials.gov; ISRCTN; the Chinese Clinical Trials Register; the German Clinical Trials Register; the Iranian Registry of Clinical Trials and the Netherlands National Trials Register, plus others);
- 3. Quarterly search of *The Cochrane Library*'s Central Register of Controlled Trials (CENTRAL);
- 4. Six-monthly searches of a number of grey literature sources: ISI Web of Knowledge Conference Proceedings; Index to Theses; Australasian Digital Theses.

To view a list of all sources searched for ALOIS see About ALOIS on the ALOIS website (www.medicine.ox.ac.uk/alois).

Details of the search strategies run in healthcare bibliographic databases, used for the retrieval of reports of dementia, cognitive improvement, and cognitive enhancement trials, can be viewed in the 'methods used in reviews' section within the editorial information about the Cochrane Dementia and Cognitive Improvement Group.

We ran additional searches in MEDLINE, Embase, PsycINFO, CINAHL, ClinicalTrials.gov and the WHO Portal/ICTRP to ensure that the searches for each suite of reviews was as comprehensive and as up-to-date as possible to identify published, unpublished, and ongoing trials. The search strategies that we created and used can be seen in Appendix 2.

Searching other resources

We screened reference lists of all included trials. In addition, we screened reference lists of recent systematic reviews, health technology assessment reports, and subject-specific guidelines identified through www.guideline.gov. The search was restricted to those guidelines meeting National Guideline Clearinghouse's (NGC) 2013 inclusion criteria published in this year or later.

We contacted experts in the field and companies marketing included interventions, in order to provide additional randomised trial reports that were not identified by the database search.

Data collection and analysis

Selection of studies

We used crowdsourcing to screen the search results. Details of this have been described here: http://www.medicine.ox.ac.uk/ alois/content/modifiable-risk-factors. In brief, teams of volunteers performed a 'first assessment' on the search results. The volunteers were recruited through the author team's institutions. They screened the results using an online tool developed for Cochrane Embase project but tailored for this programme of work. The crowd decided, based on a reading of title and abstract, whether the citation was describing a randomised or quasi-randomised trial, irrespective of the citations topic. It is estimated that this removes 75% to 90% of results retrieved. The remaining titles and abstracts were then screened by the author team, followed by full text screening. Two clinicians in the author team screened independently (AA, DD, RA). In the event of disagreement that could not be resolved by discussion, a methodologist (LYC) and additional clinician (NT or JA) arbitrated on eligibility and reasons for exclusion. All consensus rulings and reasons for exclusion were subsequently verified by methodologists (AR and MdN or LYC). Citations and full text papers from the update searches conducted in 2017 and 2018 were screened independently by two clinicians (JM and GM), who reached consensus through discussion if necessary.

Data extraction and management

Two review authors, working independently, extracted trial information using a structured data extraction form. Discrepancies were resolved by discussion, or by the involvement of a third reviewer. Where possible, we extracted (as a minimum) the following information related to characteristics of participants, intervention, and study design:

Participant characteristics:

- gender;
- baseline age (range, median, mean);
- baseline cognitive function;
- cognitive diagnostic status.

Intervention characteristics:

- nature of the intervention; generic and trade name of intervention;
 - description of the control condition;
 - duration of treatment;
 - dosage and frequency;
 - any concomitant treatments;
 - treatment adherence.

Methodological characteristics:

- trial design (individual or cluster randomisation; parallel group, factorial, or crossover design);
 - number of participants;
 - outcome measures used;
 - duration of follow-up as measured from randomisation;
 - duration of follow-up as measured from end of treatment;
 - source of financial support;
 - publication status.

If eligible cognitive outcome data were available at multiple time points, we grouped outcomes into short-term (12 weeks/three months to one year), medium-term (> one year to two years) and longer-term results (> two to five years, > five to 10 years and >10 years). For the secondary outcome all-cause dementia, only outcome data after one or more years of follow-up were considered. Within these time periods, the longest available data reported by the study were extracted (for example, if study reported data at six months, 9 months and one year, only the one-year data were extracted and analysed for the short-term time point.

For dichotomous outcomes (such as incident dementia or mortality), we extracted from each trial the number of participants with each outcome

For continuous outcomes, we extracted the number of participants in whom the outcome was measured, and the mean and standard deviation of the change from baseline or of the mean value at each relevant time point. When necessary, means and measures of dispersion were approximated from figures in the reports.

For crossover trials, we extracted data on the first treatment period only.

Whenever possible, we extracted intention-to-treat data i.e. analysing all patients according to the group randomisation. If this was not available, then we extracted and reported data from available case analyses. If these data were also not available, we considered data from 'per protocol' analyses. We contacted the trialists if we could not obtain the necessary data from the trial report.

Assessment of risk of bias in included studies

Two authors independently assessed the risk of bias in each of the included trials using Cochrane's 'Risk of bias' tool (Higgins 2011). We resolved disagreements by consensus. We assessed the risk of bias potentially introduced by suboptimal design or conduct in the domains of sequence generation, concealment of allocation,

blinding of participants and caregivers, blinding of outcome assessors, selective outcome reporting, and incomplete outcome data, including the type of statistical analyses used (true intention-to-treat versus other). The general definitions that were used are reported in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011); the review-specific definitions described in Appendix 3 are in part derived from a previously published systematic review (Rutjes 2012).

Measures of treatment effect

The measure of treatment effect for continuous outcomes was a mean difference (MD) if all studies measuring that outcome used the same scale and a standardised mean difference (SMD), defined as the between-group difference in mean values divided by the pooled standard deviation (SD), if an outcome was measured with a variety of scales. The treatment effect for dichotomous outcomes was expressed as a relative risk (RR).

Unit of analysis issues

We included no cluster-randomised trials and identified no other unit of analysis issues.

Dealing with missing data

Missing data in the individual trials may put the study estimates of effects at a high risk of bias, and may lower the overall certainty of the evidence according to GRADE (Higgins 2011). We dealt with missing data in our 'Risk of bias' assessments and planned to evaluate attrition bias in stratified analyses of the primary outcomes (Appendix 3). We thus analysed the available information and did not contact authors with a request to provide missing information. We did not impute missing data ourselves.

Assessment of heterogeneity

We considered differences in trial populations, interventions (including duration of treatment), and outcomes when assessing clinical heterogeneity and deciding which data to pool. We used visual inspection of forest plots and the tau² and I² statistics when considering whether to downgrade our confidence in results for inconsistency between trials, see also Data synthesis.

Assessment of reporting biases

Due to the low number of studies included in meta-analyses, we did not perform any funnel plot analyses to explore possible reporting biases.

Data synthesis

We pooled interventions which have been postulated to share a main mechanism of action in preventing development of dementia. Specifically, we pooled data from trials of:

- the antioxidant vitamins C, E, and ß-carotene, and
- B vitamins (thiamine (B1), pyridoxine (B6), folic acid (B9) and cobalamin (B12)) on the grounds that these are cofactors for key metabolic processes and are involved in reducing high homocysteine levels.

Whenever possible, we used standard inverse-variance randomeffects meta-analysis to combine outcome data across the trials (DerSimonian 1986) (see Data extraction and management for definitions of time points). When indicated, we converted scales to a consistent direction of effect across all trials, so that negative estimates of effect referred to those favouring active supplementation, while positive estimates referred to those favouring control. We visually inspected forest plots for the presence of heterogeneity and calculated the variance estimate tau2 as a measure of betweentrial heterogeneity (DerSimonian 1986). We prespecified a tau² of 0.04 to represent low heterogeneity, 0.09 to represent moderate heterogeneity, and 0.16 to represent high heterogeneity between trials (Spiegelhalter 2004). The I² statistic and the corresponding Chi² test were depicted in addition (Higgins 2003) for the benefit of readers more familiar with this statistic. I² describes the percentage of variation across trials attributable to heterogeneity rather than chance, with values of 25%, 50%, and 75% typically being interpreted as low, moderate, and high between-trial heterogeneity. Tau2 was preferred over I2 in the interpretation of betweentrial heterogeneity, as the interpretation of I2 can be affected by the precision of trials included in the meta-analysis (Rücker 2008). All P values were two-sided. Statistical analysis was done in Review Manager 5 (Review Manager 2014); and in STATA, release 13 (StataCorp, College Station, Texas).

Subgroup analysis and investigation of heterogeneity

Where we identified substantial inconsistency between trials in a meta-analysis, we inspected forest plots to look for obvious sources of inconsistency and reviewed the trials for clinical heterogeneity. Because of the small numbers of trials from which data could be pooled, we conducted no subgroup analyses.

Sensitivity analysis

We did not conduct any sensitivity analyses.

GRADE and summary of findings table

We constructed 'Summary of Findings' (SoF) tables for comparisons of B vitamins, antioxidant vitamins, Vitamin D + calcium, Zinc + Copper and complex multivitamins with placebo. For each comparison, we included in the SoF tables results related to overall cognitive function and the incidence of dementia at each time

point. If a study presented results for the same participants at different time points, and these could not be pooled with results from other studies, then we presented in the SoF table only the results from the longest period of follow-up.

We used GRADE to describe the certainty of the overall body of evidence for each outcome in the 'Summary of findings' table (Higgins 2011; Guyatt 2008).

Certainty refers to the degree of confidence which can be placed in the estimates of treatment benefits and harms. There are four possible ratings: 'high', 'moderate', 'low' and 'very low'. The GRADE approach rates evidence from RCTs which do not have serious limitations as 'high-certainty'. However, several factors can lead to the downgrading of the evidence to 'moderate', 'low' or 'very low'. The degree of downgrading is determined by the seriousness of these factors: study limitations (risk of bias); inconsistency; indirectness of evidence; imprecision; and publication bias (Higgins 2011; Guyatt 2008). Rating evidence 'high-certainty' implies that we are confident in our estimate of the effect, and further research is very unlikely to change this. A rating of 'very low'-certainty indicates that we are very uncertain about the obtained summary estimate of the effect.

RESULTS

Description of studies

See Characteristics of included studies and Characteristics of excluded studies.

Results of the search

We conducted searches in December 2014, July 2015, March 2016, August 2016, March 2017, and January 2018. In total, we retrieved 7,257 records from the six searches. After de-duplication, 5211 records remained. A Crowd and the CDCIG information specialist assessed these at title and abstract level. In total, 725 results remained after this assessment. The review team then screened these records. Of these, we assessed 74 full-text articles for eligibility and included 28 studies. One study was placed in the section 'Awaiting classification'; although it was marked as 'completed' in the trial registry, we were unable to find any results or to get any response from the authors. This process is depicted in Figure 1.

0 additional 7257 records identified through records identified database through other searching sources 5211 records after duplicates removed 5211 records 4486 records screened excluded 725 records screened by review team 45 full-text articles excluded: 6: Wrong outcomes 18: Wrong population 16: Wrong intervention 3: Wrong study design 1: Study terminated before enrollment 1: No further information could be found 74 full-text articles assessed for 1 article <u>awaiting</u> eligibility assessment 28 studies included in qualitative synthesis 27 studies included in quantitative synthesis (meta-analysis)

Figure I. Study flow diagram.

Included studies

We included 28 studies with a total of more than 83,000 participants in the review.

Studies of B vitamins

We included 14 studies with 27,882 participants in which the active intervention was folic acid, vitamin B12, vitamin B6, or a combination of these (Andreeva 2011; Dangour 2015; Durga 2007; Eussen 2006; Ford 2010; Hankey 2013; Kang 2008; Lewerin 2005; McMahon 2006; SEARCH 2010; Stott 2005; Toole 2004; Van der Zwaluw 2014; Walker 2012). Seven of the 14 studies had been designed with cognition as a primary outcome.

- Study design. Nine studies compared the B vitamin or B vitamin combination to placebo in a two-arm, parallel group design. In one two-arm, parallel group study (Toole 2004), the intervention was a multivitamin containing the United States' Food and Drug Administration (FDA) recommended daily allowances (RDA) of vitamins supplemented with high doses of folic acid, vitamin B6, and vitamin B12 while the comparator arm was a multivitamin containing the RDAs only. One study (Eussen 2006) had a three-arm, parallel group design with active intervention groups receiving B12 and (folic acid + B12); we combined these to make a single active intervention group. Four studies had factorial designs with placebo controls. SEARCH 2010 had a 2 x 2 factorial design including (folic acid + B12) and simvastatin. Andreeva 2011 had a 2 x 2 factorial design including (folic acid + B6 + B12) and omega fatty acids. Stott 2005 had a 2 x 2 x 2 factorial design including (folic acid + B12), B2, and B6 and reported results for all eight groups separately; for comparability with other studies, we included the (folic acid + B12) and the (folic acid + B12 + B6 groups), combining these to make a single active intervention group. Kang 2008 had a 2 x 2 x 2 x 2 factorial design. It reported on a cognitive substudy of the Women's Antioxidant and Folic Acid Cardiovascular Study (WAFACS), which was established by adding a fourth arm -(folic acid + B6 + B12) - to the Women's Antioxidant Cardiovascular Study (WACS). The active interventions in WAFACS were therefore (folic acid + B6 + B12), vitamin E, vitamin C, and \(\mathscr{G}\)-carotene. Kang 2008 reported on the effects of the B vitamins.
- Interventions. The active interventions varied in their constituents and dose. In one trial, the active intervention of interest was folic acid alone; in one trial, it was B12 alone. In five trials, the active intervention was (folic acid + B12) and in six trials (folic acid + B12 + B6), although in one of these trials the B12 dose was minimal. Daily doses of folic acid ranged from 0.4 mg to 2.5 mg, of B6 from 3 mg to 50 mg and of B12 from 0.02

mg to 1 mg. All studies had a placebo control, although for most of the factorial studies the comparison was 'B vitamin' versus 'no B vitamin' where the 'no B vitamin' group were taking a variety of other interventions. In one trial, all participants were also given 600 IU of vitamin D3 daily.

- *Population*. The study populations varied.
- \circ Six studies recruited participants with a history of cardio- or cerebrovascular disease (Andreeva 2011, Hankey 2013; MRC/BHF 2002, SEARCH 2010, Stott 2005, Toole 2004) and one recruited men with hypertension (Ford 2010). Three studies included only participants with elevated serum homocysteine levels (> 12 or 13 μ mol/L) at baseline (Durga 2007, McMahon 2006, Van der Zwaluw 2014). Two studies included only participants with mild to moderate B12 deficiency (Dangour 2015; Eussen 2006). One study included only diabetic participants (Anand 2011). In one study, participants were selected on the basis of elevated psychological distress (Walker 2012).
- o In all studies, there was evidence that most participants were cognitively healthy at baseline, but inclusion of small numbers of participants with cognitive impairment was possible. The inclusion decision was marginal for two studies. Stott 2005 reported median MMSE at baseline of 28 with an IQR of 26 to 29; hence 25% of participants in this UK study had an MMSE < 26. In Toole 2004, all participants had had a 'non-disabling' ischaemic stroke and baseline MMSE was 26.9 (SD 3.4) so it is likely that some participants had a degree of cognitive impairment at baseline. Only participants with 'severe cognitive impairment' were excluded. We decided to include both of these studies, but to consider the population as a possible source of heterogeneity.
- The age of participants also varied. In six studies, the mean age at the onset of supplementation was between 60 years and 70 years; in eight studies the mean age was between 70 years and 80 years.

Studies of antioxidant vitamins

We included eight studies with 47,840 participants in which the active intervention was one or more of the antioxidant vitamins: vitamin E, vitamin C, \(\beta\)-carotene (Anand 2011; Grodstein 2007; Kang 2006; Kang 2009; Kesse-Guyot 2011; Kryscio 2017; MRC/BHF 2002; Yaffe 2004). Only one of these studies had been designed primarily to assess cognition.

• *Study design*. One study (Anand 2011) compared vitamin E to no vitamin E in a randomised, open-label design. The remaining studies were placebo-controlled. One (Kang 2006) was a 2 x 2 factorial comparison of vitamin E and low dose aspirin. One (Kryscio 2017) was a 2 x 2 factorial comparison of

vitamin E and selenium supplements (this trial was also included in a comparison of selenium with placebo, below). Grodstein 2007 reported data on \(\mathcal{B}\)-carotene. It started as a two-arm parallel group study (ß-carotene and placebo, the Physicians' Health Study I (PHS-I)) which lasted for 13 years. After a gap of up to 18 months in some cases, approximately half the participants in PHS-I and some new recruits entered the Physicians' Health Study II (PHS-II) and a cognitive assessment was added. The original randomisation of participants in PHS-I to ß-carotene or placebo was retained in PHS-II while vitamin C, vitamin E, and a complex multivitamin were added in a factorial design. The paper therefore reported data after both long-term (mean 18 years, range 15 to 20 years) and shorter-term (mean one year, range three months to three years) treatment with ß-carotene. Kang 2009 studied vitamin E, vitamin C, and ß-carotene in a 2 x 2 x 2 factorial design. In the remaining three studies, the active intervention was a combination of antioxidant vitamins. MRC/BHF 2002 had a 2 x 2 factorial design comparing (vitamin E + vitamin C + \(\mathcal{B}\)-carotene) and simvastatin. Yaffe 2004 was a four-arm parallel group study; in this comparison, we included the (vitamin E + vitamin C + \(\mathbb{G}\)-carotene) and placebo arms (Yaffe 2004 had two additional arms: one treated with zinc and cupric oxide, which we included in another comparison, and one treated with the antioxidant combination plus zinc and cupric oxide, which we excluded). Finally, Kesse-Guyot 2011 compared a combination of vitamin E, vitamin C, ß-carotene, zinc, and selenium to placebo. In this case, it was not possible to separate the antioxidant vitamins from the minerals. This study also differed in the timing of the cognitive assessment which took place six years after the end of the eight-year supplementation period.

- Interventions. There were data on vitamin E from four trials, on vitamin C from one trial, and on \(\beta\)-carotene from two trials. Most of these data were from factorial trials in which the comparison group was receiving a variety of interventions. In two trials, the active intervention was the antioxidant combination (vitamin E + vitamin C + \(\beta\)-carotene), in one trial, it was the same combination with zinc and selenium and in one trial there were both vitamin E and (vitamin E + selenium) arms. All vitamin E was given as alpha-tocopherol. Doses of vitamin C ranged from 120 mg to 500 mg daily, of vitamin E from 600 IU on alternate days to 660 IU daily (except in Kesse-Guyot 2011 in which the vitamin E dose in the combined antioxidant was only 45 IU daily) and of \(\beta\)-carotene from 6 mg to 50 mg daily.
 - Population. The study populations varied.
- Two studies included women only (Kang 2006, Kang 2009) and two studies included men only (Grodstein 2007;
 Kryscio 2017). In Anand 2011, all participants had type 2 diabetes and were taking oral hypoglycaemics. In Kang 2009, all participants had cardiovascular disease or three or more coronary risk factors. In MRC/BHF 2002, participants were recruited on

the basis of a high risk of coronary heart disease death over the next five years. In Yaffe 2004, all participants had age-related macular degeneration.

- o Reflecting the fact that these studies were overwhelmingly cognitive 'add-ons' to studies with other purposes, only one study (Anand 2011) conducted any cognitive assessment at baseline. In this study, which was conducted in an economically-deprived population in India, a score of 24 or more on the Tamil version of the MMSE was considered normal cognition. Although we could not be sure that the other studies did not include some participants with cognitive impairment, we considered it likely in all cases that the study populations were very largely free of dementia or significant cognitive impairment and that the data were informative for our review question.
- o The age of participants included in this comparison was generally lower than in the B vitamin comparison. In three studies (Anand 2011, the long-term component of Grodstein 2007 and Kesse-Guyot 2011), the mean age at randomisation was between 50 and 60 years. In four studies, the mean or median age at randomisation was between 60 and 70 years. For the shorter-term component of Grodstein 2007, the mean age at randomisation was 71 years. MRC/BHF 2002 did not report a mean age of participants; the range at baseline was 40 years to 80 years.

Studies of vitamin D

We included one trial with 4143 participants which compared (vitamin D3 + calcium carbonate) with placebo (Rossom 2012, the Women's Health Initiative Memory Study, or WHIMS) and a small pilot study with 60 participants comparing vitamin D3 and placebo (Aspell 2017). Cognition was a primary outcome in Aspell 2017. Although Rossom 2012 was an ancillary study to a larger study with different primary aims, participants did have a detailed assessment to exclude dementia or cognitive impairment prior to enrolment in WHIMS.

- *Study design*. Both studies were two-arm parallel group studies. The mean duration of treatment and follow-up in Rossom 2012 was 7.8 years; in Aspell 2017, it was six months.
- Intervention. The doses differed markedly in these two studies. In Rossom 2012, the experimental intervention was 400 IU vitamin D3 and 1000 mg calcium carbonate daily. Use of personal supplemental vitamin D up to 600 IU (later 1000 IU) and of calcium up to 1000 mg was also permitted. All participants were also enrolled in WHI Hormone Therapy trials and may have been taking estrogen with or without progestin under those protocols. In Aspell 2017, the experimental intervention was 4000 IU of vitamin D3 on alternate days.
- *Population*. The participants in Rossom 2012 were 4143 women with a mean age of 71 years (range 65 to 80 years). The participants in Aspell 2017 were 60 community-dwelling men and women with a mean age of 68.5 years.

Studies of zinc and copper

We included two trials comparing zinc or zinc and copper supplements with placebo (Maylor 2006, Yaffe 2004), but only Yaffe 2004 with 1072 participants reported usable data. Maylor 2006 was designed to assess cognitive function, but Yaffe 2004 was part of a trial whose primary outcome was the effect of supplementation on age-related macular degeneration.

- Study design and interventions. Maylor 2006 was a threearm, parallel-group trial with 232 participants which compared two doses of zinc gluconate (15 mg and 30 mg a day) with placebo over six months. This study was eligible for inclusion, but did not report data in sufficient detail for extraction and inclusion in analyses. Yaffe 2004 was a four-arm, parallel group trial in which participants were treated for a median of 6.9 years. For this comparison, we used data from the active intervention arm in which participants were treated with 80 mg of zinc as zinc oxide and 2 mg of copper as cupric oxide.
- *Population*. The participants in Maylor 2006 were 387 people aged 55 to 87 years. The cognitive inclusion criterion was MMSE > 23, but in fact mean MMSE among participants was > 28 with low variability. The participants in Yaffe 2004 were 2166 people with age-related macular degeneration. Their median age at baseline was 69 years (range 55 to 80 years). There was no baseline cognitive assessment.

Studies of selenium

Kryscio 2017 (PREADVISE) was a 2 x 2 factorial study of vitamin E and selenium supplements. For this comparison (selenium versus placebo), we considered only the 3711 participants in the selenium only and placebo only groups. The participants who received vitamin E and (vitamin E + selenium) were considered in the antioxidant vitamins versus placebo comparison. This was an ancillary trial to SELECT, the primary aim of which was the prevention of prostate cancer.

Studies of supplementation with B vitamin, antioxidant vitamin and mineral combinations ('complex multivitamins')

We included three studies with 6306 participants in which the interventions were supplements which included both B vitamins and antioxidant vitamins, with or without additional minerals (Cockle 2000, Grodstein 2013, Wolters 2005). For ease of reference, we called these 'complex multivitamins'. Only the two smaller trials (Cockle 2000, Wolters 2005, combined n=359) were designed to assess cognition.

• *Study design*. Cockle 2000 and Wolters 2005 were small, two-arm, placebo-controlled, parallel group studies lasting 24 weeks and six months respectively. Grodstein 2013 reported the multivitamin data from a cognitive substudy (n = 5947) of the large Physicians' Health Study II (PHS-II), which is described under 'antioxidant vitamins' above. It had a 2 x 2 x 2 x 2 factorial

design. The mean time from randomisation to first cognitive assessment was 2.5 years (range 0.18 to 5.3 years) and to final cognitive assessment 8.5 years (range 0.3 to 14.2 years).

• Interventions.

- o In Cockle 2000, the daily dose of the experimental intervention contained: vitamin A palmitate 3334 IU, thiamine mononitrate (B1) 14 mg, riboflavin (B2) 16 mg, pyridoxine (B6) 22 mg, vitamin B12 0.03 mg, vitamin C 600 mg, dl-alphatocopherol acetate (vitamin E) 100 mg, folic acid 4 mg, d-biotin 2 mg and nicotinamide 180 mg. The authors stated that "[t]he dosing was calculated to provide the equivalent of one US RDA (National Research Council, 1980) of vitamin A and ten US RDAs of the remaining vitamins." The study was planned to last for 48 weeks, but due to very high dropout after week 24, no data were reported after 24 weeks.
- In Grodstein 2013, the experimental intervention was Centrum Silver ® (Pfizer). Its composition is shown in Appendix
- o In Wolters 2005, the experimental intervention was Nobilin Q10 ® (Medicom Pharma) providing daily doses of 150 mg vitamin C (calcium ascorbate), 50 mg magnesium (magnesium carbonate), 36 mg vitamin E (D- alfa-tocopherol acetate), 34 mg niacin (nicotinamide), 16 mg pantothenic acid, 9 mg β-carotene, 3.4 mg pyridoxine (pyridoxine hydrochloride), 3.2 mg riboflavin, 2.4 mg thiamine (thiamine mononitrate), 400 μg folic acid (pteroyl glutamic acid), 200 μg biotin, 60 μg selenium (selenium-enriched yeast), and 9 μg cobalamin (cyanocobalamin).

• Population.

- Cockle 2000 randomised 139 participants, Grodstein 2013 randomised 5947, and Wolters 2005 reported results for 220 (not clear how many were randomised).
- Wolters 2005 included women only and Grodstein 2013 included men only.
- o The participants in Cockle 2000 had a mean MMSE at baseline of 28.8 ± 1.5 years. There was no baseline cognitive assessment in Grodstein 2013. In Wolters 2005, baseline cognitive assessment supported the authors' assertion that subjects were cognitively healthy.
- The age of participants at baseline was median 63 years in Wolters 2005, median 70 years in Cockle 2000 and mean 72 years in Grodstein 2013.

Excluded studies

Forty-five studies were excluded (See Characteristics of excluded studies). Studies were excluded if they did not meet the inclusion criteria relating to: population (ACTRN12607000321448, Alavi 2014, Anonymous 2003, Benton 1995, Corless 1987, De Jager 2012, Flicker 2011, Hvas 2004, Imaoka 2016, Krikorian 2010, Kwok 2017, Li 2015, Macpherson 2012, Murray-Kolb

2011, NCT00903695, NCT02185222, Petersen 2005, Smith 1999, Van Uffelen 2007), outcomes (Abbasi 2013, Ford 2008, Miller 2016, NCT02467153), intervention (Arwert 2003, Bryan 2002, Chan 2010, Chew 2015; Dhesi 2004, Harris 2015, NCT01095211, NCT01708005, Ng 2017, Ong 2016, Pase 2015, Pathansali 2006, Pettersen 2017, Schietzel 2016, Summers 2010, Wouters-Wesseling 2005), or study design (Cheng 2016, Quick 2013, Rietsema 2014). One study was withdrawn prior to enrolment (Liu 2015). Maniam 2004 was a potentially eligible RCT described in an abstract published in 2004, but we were un-

able to identify any related full report. Some of the studies were excluded in this review, but were included in the Cochrane review on vitamin and mineral supplementation in people with mild cognitive impairment (McCleery 2018).

Risk of bias in included studies

Our judgments about the risks of bias in the included studies are detailed in the Characteristics of included studies tables and shown graphically in Figure 2.

Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	• Other bias
Anand 2011	•	?	•	•	?	•	•
Andreeva 2011	•	•	•	?	•	?	•
Aspell 2017	?	?	•	•	•	•	•
Cockle 2000	?	?	•	?	•	•	•
Dangour 2015	•	•	•	•	•	•	•
Durga 2007	?	•	•	•	•	?	•
Eussen 2006	?	?	•	•	•	•	•
Ford 2010	•	•	•	•	•	•	•
Grodstein 2007	•	?	?	?	•	•	•
Grodstein 2013	•	•	?	?	•	•	•
Hankey 2013	•	•	•	•	•	•	•
Kang 2006	?	?	•	?	?	?	•
Kang 2008	?	?	•	•	•	•	•
Kang 2009	?	?	•	•	?	•	•
Kesse-Guyot 2011	•	•	•	•	•	?	
Kryscio 2017	•	•	•	•	?	•	•
Lewerin 2005	?	?	•	?	?	?	•
Maylor 2006	?	?	?	?			?
McMahon 2006	•	?	•	?	•	•	•
MRC/BHF 2002	•	•	•	•	•	•	•
Rossom 2012	•	?	•	•	•	•	•
SEARCH 2010	•	•	?	•	•	•	•
Stott 2005	•	•	•	•	•	•	•
Toole 2004	•	•	•	•	?	•	•
Van der Zwaluw 2014	•	•	•	•	•	•	•
Walker 2012	•	•	•	•	•	•	•
Wolters 2005	?	?	•	?	?	•	?
Yaffe 2004	•	•	•	•		?	•

Allocation

Overall, selection bias was not a major concern. We judged all studies to be at low or unclear risk of bias in this domain. In a number of papers, our judgment was 'unclear' because there was a lack of detail given about the allocation methods used, but in some cases these were large, multicentre studies in which the methods of random sequence generation and allocation concealment were likely to have been adequate.

Blinding

One study (Anand 2011) was explicitly an open-label study and we therefore considered it to be at high risk of performance and detection bias. All other studies were described by their authors as being double-blind, although again there was, in some cases, a lack of reporting of the methods used to ensure blinding, especially of outcome assessment. In most cases, the experimental intervention and placebo were described as being identical. We judged all studies other than Anand 2011 to be at either low or unclear risk of bias in these domains.

Incomplete outcome data

Incomplete outcome data posed a significant risk of bias to the longer-term results. For some studies (Ford 2010, Grodstein 2013, Kang 2008), we judged the risk to be low or unclear for shorter-term outcomes, but high for longer-term outcomes when there had been greater loss of participants to follow-up.

Selective reporting

The included studies had a variety of primary outcomes, In a number of large studies, cognitive outcomes were not specified in protocols, but were added at a later stage. However, in these cases, simple cognitive measures were generally used and were reported. We judged five studies to be at high risk of selective reporting bias. Cockle 2000 did not report all the relevant outcomes described in the Methods section of the paper. Maylor 2006 did not report results in sufficient detail for inclusion in analyses. Stott 2005 mentioned functional outcomes in the trial registry entry which were not reported in the paper. Toole 2004 reported MMSE scores at 12 months but without the number of participants in each group, so we could not include the data in meta-analysis, and did not report the MMSE data at 24 months. Walker 2012 did not report all outcomes in sufficient detail for inclusion.

Other potential sources of bias

We identified no high risks of bias due to other factors.

Effects of interventions

See: Summary of findings for the main comparison B vitamins compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life; Summary of findings 2 Antioxidants compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life; Summary of findings 3 Vitamin D3 compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life; Summary of findings 4 Zinc + copper compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life; Summary of findings 5 Complex multivitamin compared to control for maintaining cognitive function in cognitively healthy people in mid and late life

The minimum clinically important difference on cognitive tests for healthy participants in prevention trials is unknown and is likely to vary with the age of participants and the duration of follow-up. We are therefore unable to judge the likely clinical importance of differences detected on any cognitive test. When describing the results, we refer to a difference between groups when P < 0.05

B vitamins

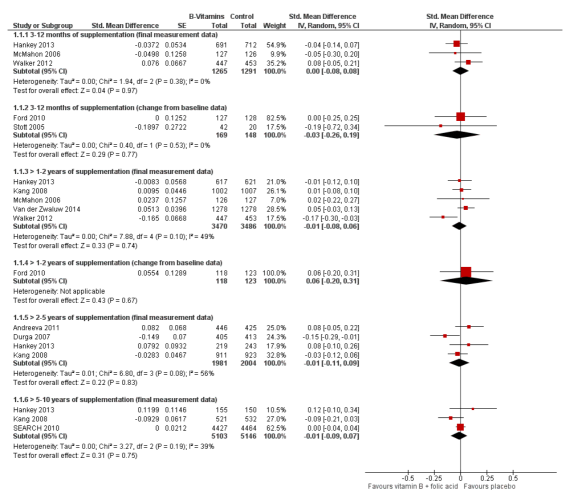
We compared B vitamins to placebo at four time points: 3-12 months of treatment, > 1-2 years, > 2-5 years and > 5-10 years. Some studies did not assess cognitive function at baseline and reported final measurement data. Some studies reported only change from baseline data. Because we were pooling outcomes measured with a variety of instruments, we used standardised mean differences and therefore pooled final measurement and change from baseline data separately.

Negative SMDs favoured B vitamins; positive SMDs favoured placebo.

Overall cognitive function

See Figure 3; Analysis 1.1

Figure 3. Forest plot of comparison: I B vitamins versus control, outcome: I.I Overall cognitive functioning.



3-12 months of supplementation:

Three studies providing final measurement data showed that there is probably no effect of B vitamins on overall cognitive function (SMD 0.00, 95% CI -0.08 to 0.08, n = 2556; moderate-certainty evidence). A fourth study (Toole 2004) also provided final measurement MMSE data on 3097 participants, but did not report the number of participants in each intervention group; it reported no significant difference between groups (P = 0.19). Two smaller studies providing change from baseline data confirmed that there may be no effect of B vitamins (SMD -0.03, 95% CI -0.26 to 0.19, n = 317; low-certainty evidence).

> 1-2 years of supplementation:

Five studies providing final measurement data showed that there is probably no effect of B vitamins (SMD -0.01, 95% CI -0.08 to 0.06, n = 6956; moderate-certainty evidence). One study reporting only change from baseline data similarly showed that there may be no effect of B vitamins (SMD 0.06, 95% CI -0.20 to 0.31, n = 241; low-certainty evidence).

> 2-5 years of supplementation:

Four studies provided final measurement data and suggested that there is probably no effect of B vitamins (SMD -0.01, 95% CI - 0.11 to 0.09, n = 3985; moderate-certainty evidence).

> 5-10 years of supplementation:

Three studies contributed data to this comparison; low-certainty evidence indicated that there may be no effect of B vitamins (SMD -0.01, 95% CI -0.09 to 0.07, n = 10249).

0.46, n = 536). A single study providing change from baseline data also favoured placebo, but the certainty of this evidence was very low (SMD 0.39, 95% CI 0.07 to 0.71, n = 171).

Episodic memory

See Analysis 1.2

3-12 months of supplementation:

Four studies providing final measurement data showed that B vitamins probably have no effect on episodic memory (SMD 0.02, -0.08 to 0.12, n = 1442; moderate-certainty evidence). Two studies providing change from baseline data confirmed that there may be no effect (SMD 0.04, 95% CI -0.15 to 0.23, n = 430; low-certainty evidence).

> 1-2 years of supplementation:

Four studies providing final measurement data suggested there may be no effect of treatment (SMD 0.05, 95% CI -0.03 to 0.13, n = 5626; low-certainty evidence). The one study providing only change from baseline data on this outcome confirmed an absence of effect (SMD 0.06, 95% CI -0.19 to 0.32, n = 241; low-certainty evidence).

> 2-5 years of supplementation:

Three studies providing final measurement data suggested there may be no effect of treatment (SMD -0.05, 95% CI -0.15 to 0.05, n = 3523; low-certainty evidence).

> 5-10 years of supplementation:

From one study, there was low-certainty evidence that there may be little or no effect of treatment (SMD -0.10, 95% CI -0.22 to 0.02, n = 1053).

Executive function

See Analysis 1.3

3-12 months of supplementation:

From three studies providing final measurement data, the effect estimate slightly favoured placebo, but there was uncertainty about the size and direction of the effect (SMD 0.14, 95% CI -0.18 to

> 1-2 years of supplementation:

Very low-certainty evidence from three studies providing final measurement data suggested no effect of treatment (SMD -0.01, 95% CI -0.15 to 0.13, n = 2976). There was inconsistency in this analysis in which one study (McMahon 2006) tended to favour placebo.

> 2-5 years of supplementation:

Two studies providing final measurement data showed that there may be no effect of treatment (SMD -0.06, 95% CI -0.13 to 0.02, n = 2648; low-certainty evidence).

> 5-10 years of supplementation:

From one study, there was low-certainty evidence that there may be no effect of treatment (SMD 0.02, 95% -0.10 to 0.15, n = 1049).

Speed of processing

See Analysis 1.4

3-12 months of supplementation:

Two studies provided final measurement data on this outcome and showed that there may be no effect of treatment (SMD -0.04, 95% CI -0.28 to 0.19, n = 289; low-certainty evidence). Pooled data from three studies providing change from baseline data gave a similar result, but with very low-certainty evidence (SMD 0.12, 95% CI -0.34 to 0.58, n = 487).

> 1-2 years of supplementation:

Two studies provided final measurement data and showed that there is probably no effect of treatment (SMD 0.03, 95% CI - 0.09 to 0.15, n = 1030; moderate-certainty evidence). A further study providing change from baseline data confirmed this (SMD -0.04, -0.29 to 0.21, n = 241; low-certainty evidence).

> 2-5 years of supplementation:

One study provided final measurement data and showed no effect of treatment (SMD -0.08, 95% CI -0.22 to 0.06, n = 818; high-certainty evidence).

Quality of life

> 1-2 years of supplementation:

Ford 2010 reported that participants in the B vitamin and placebo groups did not differ significantly on any of the four mental summary scales of the SF-36 after 24 months.

No other included studies reported on quality of life.

Clinical global impression

No included studies reported on clinical global impression.

Functional performance

No included studies reported on functional performance.

Incidence of MCI or dementia

See Analysis 1.5.

Hankey 2013 (VITATOPS study) reported on the incidence of cognitive impairment (defined as MMSE score < 24 during follow-up on two or more occasions) after a median of 2.8 years of supplementation. Of those who had at least two MMSE measurements during follow-up, 50/908 (5.51%) in the B vitamin group and 49/895 (5.47%) in the placebo group met this criterion for cognitive impairment (RR 1.01, 95% CI 0.69 to 1.48, n = 1803; moderate-certainty evidence).

Ford 2010 treated participants for two years, but also followed up the participants who were still alive approximately eight years after the baseline study assessment, assessing 73 of the 299 randomised participants with the Telephone Interview for Cognitive Status scale (TICS) and ascertaining dementia diagnoses from a population register for all 172 survivors. Defining cognitive impairment as TICS \leq 27, they found an OR 0.72 (95% CI 0.25 to 2.09) favouring the B vitamin group. For either a dementia diagnosis or TICS \leq 27, the OR was 0.76 (95% CI 0.37 to 1.56) also favouring B vitamins. Due to serious imprecision and risk of attrition bias, this was very low-certainty evidence.

SEARCH 2010 investigators recorded diagnoses of dementia at final follow-up (after a mean of 6.7 person-years) "based on reports from a participant/relative or doctor, there was no further confirmation or adjudication" (communication from author). The paper stated that "similar numbers of participants in each treatment

group were reported to have developed dementia during follow-up (vitamins, 43 (0.7%), vs placebo, 38 (0.6%))".

Serious adverse events

Most papers made only general statements about an absence of serious adverse events in both treatment groups.

All-cause mortality

See Analysis 1.6

3-12 months of supplementation:

In one study (Dangour 2015), one participant died in the placebo group and no participants in the B vitamin group.

> 1-2 years of supplementation:

In Toole 2004, in which all participants had a history of recent ischaemic stroke, 99/1821 participants (5.4%) died in the B vitamin group compared to 117/1847 participants (6.3%) in the comparator group over 2 years of follow-up (RR 0.9, 95% CI 0.7 to 1.1, n = 3668).

> 2-5 years of supplementation:

In one study (Andreeva 2011), 24/585 participants (4.1%) died in the placebo group and 44/586 participants (7.5%) in the B vitamin group (RR 1.8 favouring placebo, 95% CI 1.17 to 3.08, n = 1171; very low-certainty evidence).

> 5-10 years of supplementation:

In SEARCH 2010, in which all participants had a history of myocardial infarction, 983/6033 participants (16.3%) died in the B vitamin group compared to 951/6031 participants (15.8%) in the placebo group over a mean of 6.7 years of follow-up (RR 1.03, 95% CI 0.95 to 1.12, n = 12064; moderate-certainty evidence). Mortality data was not clearly reported in any other study.

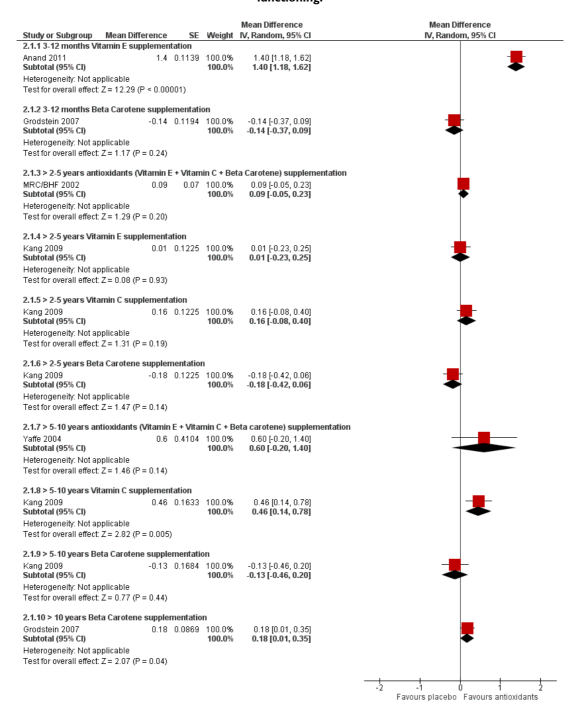
Antioxidant vitamins

We compared antioxidants to placebo at four time points: 3-12 months of treatment, > 2-5 years, > 5-10 years and > 10 years. (We had no data for the > 1-2 year time point).

Overall cognitive function

See Analysis 2.1; Figure 4

Figure 4. Forest plot of comparison: 2 Antioxidants versus control, outcome: 2.1 Overall cognitive functioning.



Positive values favoured antioxidants; negative values favoured placebo.

3-12 months of supplementation:

One study of vitamin E in middle-aged adults with type 2 diabetes (Anand 2011) found a higher MMSE in the intervention group, but this was the only open-label study and the certainty of this evidence was very low due to risk of bias, imprecision and indirectness (MD 1.4, 95% CI 1.18 to 1.62, n = 74).

One study of beta-carotene (Grodstein 2007) included an older population (mean age 73 years) who were treated for a mean of 12 months, but a range of two months to three years. Among these participants, there was probably no effect of beta-carotene on overall cognitive function measured with the TICS (MD -0.14, 95% CI -0.37 to 0.09, n = 1904; moderate-certainty evidence).

> 2-5 years of supplementation:

One study using a combination of vitamin C, E, and beta-carotene (MRC/BHF 2002) found that there was probably no effect of the antioxidant treatment on the TICS after a mean five years of follow-up (MD 0.09, 95% CI -0.05 to 0.23, n = 20536; moderate-certainty evidence).

Kang 2009 was a factorial study which reported TICS results separately for vitamin C, E, and beta-carotene after a mean of 3.5 years of treatment. It reported the mean difference (with 95% CI) between each vitamin group and placebo, but reported only the total number of participants (2824) and not the number in each treatment group. This study showed that there may be no effect of vitamin C (MD 0.16, 95% CI -0.08 to 0.40, n not reported; low-certainty evidence), vitamin E (MD 0.01, 95% CI -0.23 to 0.25, n not reported; low-certainty evidence) or beta-carotene (MD -0.18, 95% CI -0.42 to 0.06; n not reported; low-certainty evidence).

> 5-10 years of supplementation:

One study using a combination of vitamins C and E and betacarotene for a median of 6.9 years (Yaffe 2004) found that there may be no effect on the modified MMSE (3MS) (MD 0.6, 95% CI -0.2 to 1.40, n = 1100; moderate-certainty evidence).

Kang 2009 also reported outcome data from a final assessment after a mean of 8.9 years. However, again it reported only the total number of participants (n = 1586) at this time point and not the number in each group. A second study (Kang 2006) reported TICS data after approximately 10 years of treatment with vitamin E. We were unable to pool data vitamin E data from these two studies because of the unknown number of relevant participants in Kang 2009. In Kang 2006, the first cognitive assessment was conducted 5.6 years (range 4.4 to 6.8) after randomisation and the last 4.0 (range 2.6 to 5.7) years later. The mean difference in TICS scores between groups at final cognitive assessment was 0.04 points (95% CI -0.12 to 0.21, n = 5226). The mean difference in cognitive decline on the TICS between the vitamin E and placebo groups from first to last cognitive assessment was 0.02 points (95% CI -0.02 to 0.06; P = 0.31). In the vitamin E arm from Kang 2009, there was also no evidence of an effect on final TICS score (MD -0.16, 95% CI -0.49 to 0.16; n not reported, P = 0.33). We considered the overall certainty of this evidence on vitamin E to be low. In the vitamin C arm from Kang 2009, there may have been a small beneficial effect of vitamin C on final TICS score (MD 0.46, 95% CI 0.14 to 0.78, n not reported, P = 0.006; low-certainty evidence). In the beta-carotene arm from Kang 2009, there was no evidence of an effect of treatment (MD -0.13, 95% CI -0.46 to 0.19, n not reported, P = 0.42; low-certainty evidence).

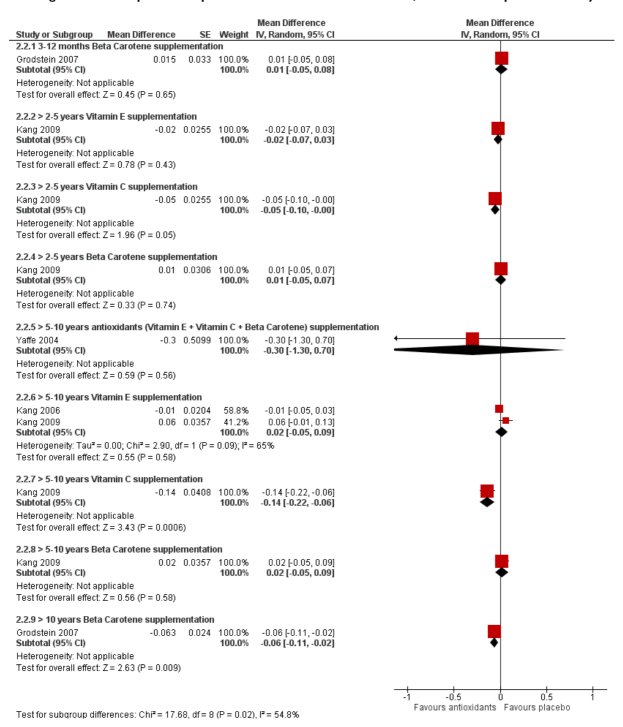
> 10 years of supplementation:

Grodstein 2007 included 4052 participants who had been treated with beta-carotene for a mean of 18 years (range 15 to 20 years), starting at a mean age of 56 years. There was moderate-certainty evidence of a very small beneficial effect of beta-carotene treatment on the TICS (MD 0.18, 95% CI 0.01 to 0.35, n = 4052).

Episodic memory

See Analysis 2.2; Figure 5

Figure 5. Forest plot of comparison: 2 Antioxidants versus control, outcome: 2.2 Episodic memory.



3-12 months of supplementation:

Grodstein 2007 found that there was probably no effect of beta-carotene (MD 0.01, 95% CI -0.05 to 0.08, n = 1904; moderate-certainty evidence) on a composite verbal memory measure after a mean of 12 months (range two months to three years).

> 2-5 years of supplementation:

From Kang 2009, there was probably a very small beneficial effect of vitamin C on the TICS 10-word recall test (MD -0.05, 95% CI -0.10 to 0; n = 2824; low-certainty evidence), but probably no effect of vitamin E (MD -0.02, 95% CI -0.07 to 0.03, n = 2824; low-certainty evidence) or beta-carotene (MD 0.01, 95% CI -0.05 to 0.07, n = 2824; low-certainty evidence).

> 5-10 years of supplementation:

Yaffe 2004, using combined vitamin C and E and beta-carotene, found no evidence of an effect on the delayed recall logical memory test (MD -0.30, 95% CI -1.30 to 0.70, n = 1100; low-certainty evidence).

A pooled analysis of two studies using vitamin E provided very low-certainty evidence of no effect (MD 0.02, 95% CI -0.05 to 0.09, n = 6812, 2 studies). There was low-certainty evidence from a single study (Kang 2009) that there may be a small beneficial effect of vitamin C (MD -0.14, 95% CI -0.22 to -0.06, n = 1583) and no effect of beta-carotene (MD 0.02, 95% CI -0.05 to 0.09, n = 1583).

Kesse-Guyot 2011 treated participants for eight years with a combination of vitamin C, vitamin E, beta-carotene, zinc, and selenium and assessed cognition six years after the end of treatment. On a delayed cued-recall test (the RI-48), there was a small benefit for the vitamin group (MD 0.61, 95% CI 0.02-1.20, n = 4447, moderate-certainty evidence).

> 10 years of supplementation:

Grodstein 2007 found evidence of a small beneficial effect of beta-carotene on a composite verbal memory measure after treatment for a mean of 18 years (range 15 to 20 years), starting at a mean age of 56 years (MD -0.06, 95% CI -0.11 to -0.02; n = 4052; moderate-certainty evidence).

Executive function

See Analysis 2.3

3-12 months of supplementation:

There was moderate-certainty evidence from Grodstein 2007 that beta-carotene probably has no effect on executive function measured with a category fluency test after a mean of 12 months (range two months to three years) (MD 0.07, 95% CI -0.48 to 0.62, n = 1904).

> 2-5 years of supplementation:

There was low-certainty evidence from Kang 2009 of a small detrimental effect of vitamin E on category fluency (MD 0.42, 95% CI 0.06 to 0.78, n=2819) and of no effect of vitamin C (MD -0.03, 95% CI -0.39 to 0.33, n=2819) or beta-carotene (MD 0.22, 95% CI -0.14 to 0.58, n=2819).

> 5-10 years of supplementation:

Yaffe 2004 found no evidence of an effect of combined vitamin C and E and beta-carotene on category fluency (MD -0.4, 95% CI -0.99 to 0.19, n = 1100; low-certainty evidence)

Pooled data on vitamin E from Kang 2006 and Kang 2009 showed no evidence of an effect on category fluency (MD 0.11, 95% CI -0.26 to 0.48, n = 6808; low-certainty evidence). Kang 2009 also found no evidence of an effect of vitamin C (MD -0.25, 95% CI -0.7 to 0.2, n = 1583; low-certainty evidence) or beta-carotene (MD 0.01, 95% CI -0.44 to 0.46, n = 1583; low-certainty evidence). Kesse-Guyot 2011 treated participants for eight years with a combination of vitamin C, vitamin E, beta-carotene, zinc, and selenium and assessed cognition six years after the end of treatment. There was no clear evidence of a difference between groups on category fluency (MD 0.50, 95% CI -0.09 to 1.09, n = 4447, moderate-certainty evidence).

> 10 years of supplementation:

Grodstein 2007 found that beta-carotene probably has no effect on executive function measured with a category fluency test after treatment for a mean of 18 years (range 15 to 20 years) (MD 0.01, 95% CI -0.36 to 0.38, n = 4052; moderate-certainty evidence).

Speed of processing

> 5-10 years of supplementation:

Kesse-Guyot 2011 treated participants for eight years with a combination of vitamin C, vitamin E, beta-carotene, zinc, and sele-

nium and assessed cognition six years after the end of treatment. There was no difference between groups on time taken to complete a trial-making task (TMT) (MD 0.00, 95% CI -0.59 to 0.59, n = 4447, moderate-certainty evidence).

there were 65 prostate cancer cases per 1000 men taking placebo only and 76 prostate cancer cases per 1000 men taking vitamin E only.

Incidence of dementia

See Analysis 2.4

> 2-5 years of supplementation:

One study (MRC/BHF 2002) reported the number of participants who developed dementia during follow-up. All participants were at high vascular risk. There were 62 events, 31 in each group (RR 1, 95% CI 0.61 to 1.64, n = 20536; low-certainty evidence).

> 5-10 years of supplementation:

The mean duration of supplement use in Kryscio 2017 was 5.4 ± 1.2 years although follow-up continued for up to six years after supplement discontinuation. The unadjusted cumulative incidence of dementia was 3.95% in the vitamin E arm, 4.96% in the (vitamin E + selenium) arm and 4.62% in the placebo arm. A modified ITT analysis including all participants who had at least one follow-up visit and adjusted for a variety of covariates showed that neither arm had an incidence significantly different from the placebo arm: vitamin E HR 0.88 (95% CI 0.64 to 1.20, P = 0.41); (vitamin E + selenium) HR 1.00 (95% CI 0.74 to 1.35, P = 0.98). We considered this to be low-certainty evidence due to imprecision and indirectness.

All-cause mortality

See Analysis 2.5

> 2-5 years of supplementation:

In one study (MRC/BHF 2002), 1389/10267 participants died in the placebo group and 1446/10269 participants in the antioxidant group (RR 1.04, 95% CI 0.97 to 1.11, n = 20536; moderate-certainty evidence).

Serious adverse events

Kryscio 2017 was an ancillary study to SELECT, in which the primary study was the incidence of prostate cancer. SELECT began recruitment in 2002. By 2011, there was a statistically significant 17% increase in the risk of prostate cancer in the men taking vitamin E only compared to those taking placebo only. After a mean of seven years of follow-up (mean 5.5 years taking supplements)

Other outcomes

Not reported.

Vitamin D

We considered data unsuitable for pooling because of the five-fold difference in the daily dose of vitamin D3.

Overall cognitive function

See Analysis 3.1

There was moderate-certainty evidence from the small pilot study (Aspell 2017) that the change in Montreal Cognitive Assessment (MoCA) score over six months did not differ between the vitamin D3 group (4000 IU on alternate days) and the placebo group (MD 0.76 points, SD not reported, n = 60, P = 0.186).

There was moderate-certainty evidence from one study (Rossom 2012) that there was probably no effect of treatment with a daily dose of 400 IU vitamin D3 + calcium carbonate on the modified MMSE (3MS) at any time point up to a mean follow-up of 7.8 years.

3-12 months of supplementation:

MD 0, 95% CI -0.24 to 0.24, n = 4122.

> 1-2 years of supplementation:

MD 0, 95% CI -0.25 to 0.25, n = 3910.

> 2-5 years of supplementation:

MD -0.2, 95% CI -0.53 to 0.13, n = 3515.

> 5-10 years of supplementation:

MD -0.1, 95% CI -0.81 to 0.61, n = 918 (low-certainty evidence, additional downgrade due to incomplete outcome data).

Episodic memory/executive function/speed of processing

Aspell 2017 reported results for executive function using the Trail Making Task but no results for episodic memory, although the trial registry entry indicated that it was to be measured with the Wechsler Memory Scale. For change in TMT scores over six months, there was probably little or no difference between the groups (TMTB-A: MD 5.1, SD not reported, n = 60, P = 0.467).

In annual domain-specific cognitive assessments over approximately three to eight years, the authors of Rossom 2012 reported that there were "no significant differences in any domain-specific cognitive scores between treatment groups either in unadjusted analyses or after adjustment for age, smoking, baseline 3MS, and treatment assignment in the hormone or diet modification trials." Results were not fully reported. They included the California Verbal Learning Test (episodic memory), letter and category fluency (executive function) and finger tapping (speed of processing).

Incidence of MCI or dementia

In Rossom 2012, "[t]he incidence of MCI or all-cause dementia did not differ significantly between placebo and treatment groups, with 4.8% of calcium plus vitamin D subjects (62.2 per 10,000 person years) and 5.1% of placebo subjects (65.9 per 10,000 person years) developing cognitive impairment during a mean follow-up time of 7.8 years (HR 0.94, 95% CI 0.72 to 1.24; P = 0.68)". The survival curve showed that the incidence did not differ over the course of follow-up. We considered this to be low-certainty evidence, downgraded due to imprecision and indirectness (women only).

Other outcomes

Not reported.

Zinc and copper

For this comparison, the only data we were able to extract came from Yaffe 2004 which treated participants with 80 mg zinc and 2 mg copper daily for a median of 6.9 years. No usable data were available from Maylor 2006, but the authors reported finding no cognitive effects of six months of supplementation with zinc gluconate.

Overall cognitive function

See Analysis 4.1

> 5-10 years of supplementation:

One study (Yaffe 2004) found no evidence of an effect of treatment on the modified MMSE (MD 0.6, 95% CI -0.19 to 1.39, n = 1072; moderate-certainty evidence).

Episodic memory

See Analysis 4.2

> 5-10 years of supplementation:

One study (Yaffe 2004) found no evidence of an effect of treatment on a delayed recall test (MD -0.7 95% CI -1.74 to 0.34, n = 1072; moderate-certainty evidence).

Executive function

See Analysis 4.3

> 5-10 years of supplementation:

One study (Yaffe 2004) found no evidence of an effect of treatment on category fluency (MD -0.3, 95% CI -0.89 to 0.29, n = 1072; moderate-certainty evidence).

Incidence of dementia or MCI

Yaffe 2004 reported that at the time of administration of the cognitive test battery, 4.5% of participants had a 3MS score < 80, which was the authors' definition of cognitive impairment, and that "there was no significant effect of treatment group on likelihood of having cognitive impairment." We considered this to be low-certainty evidence.

Other outcomes

Not reported.

Selenium

For this comparison, the only data came from the selenium arm of Kryscio 2017. The only outcome reported was incidence of dementia. Supplements - in this case, selenium 200 mcg per day - were taken for a mean of 5.4 ± 1.2 years although follow-up continued for up to six years after supplement discontinuation. The unadjusted cumulative incidence of dementia was 4.15% in the selenium arm and 4.62% in the placebo arm. A modified ITT analysis including all participants who had at least one follow-up visit and adjusted for a variety of covariates showed no significant difference in incidence between the selenium and placebo arms (HR 0.83, 95% CI 0.61 to 1.13, P = 0.23, n = 7388). We considered this to be low-certainty evidence due to imprecision and indirectness.

Complex multivitamins (containing B vitamins and antioxidants +/- minerals)

Overall cognitive function

See Analysis 5.1

> 2-5 years of supplementation:

One study (Grodstein 2013) found no effect of a complex multivitamin on the TICS over a mean of 2.5 years of treatment (range 0.18 to 5.3 years) (MD 0.04, 95% CI -0.09 to 0.18; n = 5947; low-certainty evidence).

> 5-10 years of supplementation:

The same study found no effect of the complex multivitamin on the TICS over a mean of 8.5 years (range 0.3 to 14.2 years) (MD 0.12, 95% CI -0.14 to 0.38; n = 2324; low-certainty evidence).

Episodic memory

See Analysis 5.2

3-12 months of supplementation:

One study (Cockle 2000) found no evidence of an effect of treatment on a word scan (recognition) test (MD 33.17, 95% CI - 28.12 to 94.46; n = 124; low-certainty evidence). Another study (Wolters 2005) used a pattern recognition subtest of the Berliner Amnesie test (BAT) and reported the results as medians with 5th to 95th percentiles; the difference between groups was 'not significant' (no P value given).

> 2-5 years of supplementation:

One study (Grodstein 2013) found no evidence of an effect of treatment on a composite verbal memory score after a mean of 2.5 years (range 0.18 to 5.3 years) (MD -0.01, 95% CI -0.05 to 0.03; n = 5942; low-certainty evidence).

> 5-10 years of supplementation:

One study (Grodstein 2013) found no evidence of an effect of treatment on a composite verbal memory score after a mean of 8.5 years (range 0.3 to 14.2 years) (MD -0.01, 95% CI -0.07 to 0.05; n = 2324; low-certainty evidence).

Executive function

See Analysis 5.3

> 2-5 years of supplementation:

One study (Grodstein 2013) found no evidence of an effect of treatment on verbal fluency after a mean of 2.5 years (range 0.18 to 5.3 years) (MD -0.02, 95% CI -0.33 to 0.29; n = 5942; low-certainty evidence).

> 5-10 years of supplementation:

One study (Grodstein 2013) found no evidence of an effect of treatment on verbal fluency after a mean of 8.5 years (range 0.3 to 14.2 years) (MD -0.22, 95% CI -0.65 to 0.21; n = 2324; low-certainty evidence).

Speed of processing

See Analysis 5.4

3-12 months of supplementation:

One study (Cockle 2000) found no evidence of an effect of treatment on a total reaction time (TRT) (MD -14.71, 95% CI - 57.22 to 27.80; n = 127; low-certainty evidence). Another study (Wolters 2005) used the symbol search subtest from the WAIS-III and reported the results as medians with 5th to 95th percentiles. The vitamin group was significantly faster than the placebo group (P = 0.015) at baseline and this difference persisted at the end of the intervention (P = 0.01).

Other outcomes

Not reported.

ADDITIONAL SUMMARY OF FINDINGS [Explanation]

Antioxidants compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life

Patient or population: cognitively healthy people in mid and late life

Setting: community

Intervention: antioxidant vitamins (vitamins C, E, and beta-carotene, alone or in combination)

Comparison: placebo

Outcomes			Relative effect (95% CI)	№ of participants (studies)	Certainty of the evidence	Comments
	Risk with placebo	Risk with Antioxidants			(GRADE)	
Vitamin E supplementation Overall cognitive functioning assessed with: MMSE Scale from: 0 to 30 follow-up: 3 months	The mean MMSE score was 26.6	MD 1.4 higher (1.18 higher to 1.62 higher)	-	74 (1 RCT)	⊕○○○ VERY LOW ¹²³	Higher score indicates better cognitive function.
Vitamin E supplementation Overall cognitive functioning assessed with: TICS Scale from: 0 to 41 follow-up:range 6 years to 10 years	Two studies found no o	evidence of an effect of	-	(2 RCTs)	⊕⊕⊖⊖ LOW ⁴⁵	Range of follow-up durations was approximate. Mean in one study was 9.6 years and in the other study was 8.9 years. There were 5226 participants in one study. The number in the other study was not reported for this time point (approximately 800)

Beta-carotene supplementation Overall cognitive functioning assessed with: TICS Scale from: 0 to 41 follow-up: mean 12 months	The mean TICS score was 34.29	MD 0.14 lower (0.37 lower to 0.09 higher)	- 9	1904 (1 RCT)	⊕⊕⊕⊝ MODERATE ⁶	Higher score indicates better cognitive function.
Beta-carotene supplementation Overall cognitive functioning assessed with: TICS follow-up: mean 8.9 years	Not reported	MD 0.13 lower (0.46 lower to 0.19 higher)	9	(1 RCT)	⊕⊕○○ LOW ⁴⁷	Mean difference with 95% CI reported in paper. Absolute values in each treatment group not reported. Number in each treatment group at final follow-up not reported
Beta-carotene supplementation Overall cognitive functioning assessed with: TICS Scale from: 0 to 41 follow-up: mean 18 years	The mean TICS score was 34.23	MD 0.18 higher (0.01 higher to 0.3 higher)	- 5	4052 (1 RCT)	⊕⊕⊕⊖ MODERATE ⁶	
Vitamin C supplementation Overall cognitive functioning assessed with: TICS follow-up: mean 8.9 years	Not reported	MD 0.46 higher (0.14 higher to 0.79 higher)		(1 RCT)	⊕⊕⊖⊖ LOW ⁴⁷	Mean difference with 95% CI reported in pa- per. Absolute values in each group not re- ported. Number in each treatment group at final follow-up not reported

Supplementation with beta-carotene + vit C + vit E Overall cognitive func- tioning assessed with: TICSm Scale from: 0 to 39 follow-up: mean 5 years	The mean TICSm score was 24.02	MD 0.09 higher (0.05 lower to 0.23 higher)	-	20536 (1 RCT)	⊕⊕⊕⊝ MODERATE ⁸	
Supplementation with beta-carotene + vit C + vit E Overall cognitive func- tioning assessed with: 3MS (modified MMSE) Scale from: 0 to 100 follow-up: median 6.9 years	The mean 3MS score was 92.1	MD 0.6 higher (0.2 lower to 1.4 higher)		1100 (1 RCT)	⊕⊕⊕⊝ MODERATE ⁴	Higher score indicates better cognitive function.
dementia follow-up: mean ap-	Two studies, one using carotene, vit C and vit selenium, found no sign incidence of dementia	E and one using vit E \pm		25 993 (2 RCTs)	⊕⊕⊖⊖ LOW ⁸⁹	

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ¹ Downgraded for risk of bias due to lack of blinding (open-label study)
- ² Downgraded for indirectness (diabetic participants only)
- ³ Downgraded for imprecision due to small sample size
- ⁴ Downgraded for risk of bias due to incomplete outcome data
- ⁵ Downgraded for indirectness (both studies included only women and in one study all participants also had cardiovascular disease or 3 or more coronary risk factors)
- ⁶ Downgraded for indirectness (included men only)
- ⁷ Downgraded for indirectness (included only women with cardiovascular disease or \geq 3 coronary risk factors)
- ⁸ Downgraded for indirectness (all participants had high risk of coronary heart disease death over the next 5 years)
- ⁹ Downgraded for imprecision due to wide confidence interval

MMSE:MiniMentalStateExamination

TICS: Telephone Interview for Cognitive Status

TICSm: Modified Telephone Interview for Cognitive Status

3MS: The Modified Mini-Mental State Examination

Vitamin D3 compared to placebo for maintaining cognitive function in cognitively healthy people in mid and late life

Patient or population: cognitively healthy people in mid and late life

Setting: community

Intervention: vitamin D3 +/- calcium carbonate

Comparison: placebo

Outcomes	Outcomes Anticipated absolute effects* (95% CI)		No of participants (studies)	Certainty of the evidence	Comments	
	Risk with placebo Risk with Vitamin D3			(GRADE)		
Overall cognitive functioning assessed with: MoCA change from baseline Scale from: 0 to 30 follow-up: 6 months	There was no significant difference between groups in the change in MoCA score (P=0.186)		60 (1 RCT)	⊕⊕⊕⊝ MODERATE ¹	Experimental intervention was vitamin D3 4000 IU on alternate days Data not fully reported (conference abstract only).	
Overall cognitive functioning assessed with: 3MS Scale from: 0 to 100 follow-up: mean 7.8 years	The mean 3MS score MD 0.1 lower was 96.6 (0.81 lower to 0.61 higher)	-	918 (1 RCT)	⊕⊕⊖⊖ LOW ²³	Experimental intervention was vitamin D3 400 IU/day and calcium carbonate 1000 mg/day	
dementia or MCI	No significant difference in incidence between treatment groups. In an intention-to-treat analysis, incidence was 62.2/10 000 person-years in the vit D + Ca group and 65.9/10 000 person-years in the placebo group (HR 0.94, 95% CI 0.72 to 1.24, P = 0.68)		4143 (1 RCT)	⊕⊕⊖⊝ LOW ³⁴	Experimental intervention was vitamin D3 400 IU/day and calcium carbonate 1000 mg/day	

*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ² Downgraded for risk of bias due to incomplete outcome data
- ³ Downgraded for indirectness (study included women only)
- ⁴ Downgraded for imprecision due to wide confidence interval MCI:MildCognitiveImpairment

MOCA: Montreal cognitive assessment

3MS: The Modified Mini-Mental State Examination

¹ Downgraded for imprecision (small sample size)

Patient or population:cognitively healthy people in mid and late life

Setting: Community

Intervention: Zinc + copper oxides

Comparison: Placebo

Outcomes	Anticipated absolute eff	fects* (95% CI)	Relative effect (95% CI)	№ of participants (studies)	Certainty of the evi- Comments dence
	Risk with placebo	Risk with Zinc + copper			(GRADE)
Overall cognitive functioning assessed with: 3MS Scale from: 0 to 100 follow-up: median 6.9 years		MD 0.6 higher (0.19 lower to 1.39 higher)	-	1072 (1 RCT)	⊕⊕⊕⊖ MODERATE ¹²
or MCI	ia 4.5% of participants met the authors' definition of cognitive impairment (3MS < 80) and "there IS was no significant effect of treatment group on likelihood of having cognitive impairment."			1072 (1 RCT)	⊕⊕⊖⊝ LOW ³

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

MCI:Mildcognitiveimpairment

Downgraded for risk of bias due to unclear selective reporting and high risk of bias due to incomplete outcome data Downgraded for indirectness (Yaffe 2004 included participants with age-related macular degeneration)

³ Downgraded two levels for study limitations due to incomplete outcome data and selective reporting ³ MS:TheModifiedMini-MentalStateExamination

Complex multivitamin compared to control for maintaining cognitive function in cognitively healthy people in mid and late life

Patient or population: cognitively healthy people in mid and late life

Setting: community

Intervention: complex multivitamins (combinations of B vitamins and antioxidant vitamins +/- minerals)

Comparison: placebo

Outcomes	Anticipated absolute effects* (95% CI)		Relative effect (95% CI)	№ of participants (studies)	Certainty of the evidence	Comments
	Risk with control	Risk with Complex multivitamin			(GRADE)	
Overall cognitive functioning assessed with: TICS Scale from: 0 to 41 follow-up: mean 2.5 years	The mean TICS score was 34.3	MD 0.04 higher (0.09 lower to 0.18 higher)	-	5947 (1 RCT)	⊕⊕⊖⊖ LOW ¹²	Follow-up ranged from 0.18 to 5.3 years.
Overall cognitive functioning assessed with: TICS Scale from: 0 to 41 follow-up: mean 8.5 years	The mean TICS score was 33.1	MD 0.12 higher (0.14 lower to 0.38 higher)	-	2324 (1 RCT)	⊕⊕⊖⊝ LOW ¹²	Follow-up ranged from 0.3 to 14.2 years.
Incidence of MCI or all- cause dementia - not measured			-	-	-	

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

TICS:TelephoneInterviewforCognitiveStatus

MCI: mild cognitive impairment

¹ Downgraded for study limitations due to incomplete outcome data and wide range of follow-up durations

² Downgraded for indirectness (study included only men)

DISCUSSION

Summary of main results

B vitamins

Fourteen studies with 27,882 participants contributed data. Only two studies lasted for longer than five years, although between them these included more than 10,000 participants. We found no evidence that giving B vitamin supplements to cognitively healthy adults, mainly in their 60s and 70s, had any effect on global cognitive function or on episodic memory, executive function, or speed of processing. There were very sparse data on adverse effects or on incidence of cognitive impairment or dementia, quality of life, functional performance, or clinical global impression.

Antioxidant vitamins (vitamins C and E, and betacarotene)

Eight studies with 47,840 participants contributed data. Six studies had a mean treatment duration longer than five years. Two studies followed participants up for more than 10 years. The results of these trials were more mixed than for B vitamins. There was low-certainty evidence that vitamin C supplements may have a small beneficial effect on episodic memory after 2 to 5 years of treatment and on overall cognitive function after 5 to 10 years, although effects were not seen at other time points. There were no longer-term data on vitamin C. For vitamin E, there was lowcertainty evidence of a small detrimental effect on executive function after 2 to 5 years and no data beyond 10 years. For betacarotene, there was no evidence of any effects up to 10 years, but moderate-certainty evidence from one large study with a duration of 15 to 20 years of small benefits on overall cognitive function and episodic memory after a mean of 18 years of treatment. There were very few cognitive data from trials using combinations of antioxidant vitamins. There was no evidence of an effect of 5 to 10 years of antioxidant combination treatment in two trials, but one trial which assessed participants six years after the end of an 8-year treatment period (using vitamins C, E, beta-carotene plus zinc and selenium) reported a small benefit on episodic memory. Two studies found no effect of antioxidant vitamins on the incidence of dementia. Other review outcomes were not reported in these trials.

Vitamin D

One large trial in this comparison provided low to moderate-certainty evidence of no effect on cognitive outcomes or the incidence of MCI or dementia of a relatively low dose of vitamin D3 with calcium carbonate in any time period up to 10 years. It provided no evidence on other review outcomes. There was also a small pilot trial which used a much higher dose of vitamin D3 and found no evidence of cognitive effects over six months.

Zinc and copper

One study contributed usable data, providing low-certainty evidence of no effect on cognitive outcomes or the incidence of cognitive impairment after 5 to 10 years, but no evidence on other outcomes.

Selenium

One study found no evidence of an effect of approximately five years of selenium supplementation on the incidence of dementia, but there was no information on other outcomes.

Complex multivitamins (B vitamins and antioxidants +/- minerals)

Two small, short-term studies and one large study with a mean duration of treatment of 8.5 years at final follow-up found no effects of complex multivitamins on cognitive outcomes. There was no evidence on other review outcomes.

Overall completeness and applicability of evidence

Our principal interest was in whether or not vitamin supplementation could be of benefit to the general middle-aged or older population in terms of maintaining cognitive function or preventing dementia in late life. Some general points can be made about the evidence across all the interventions.

Many of the included studies, especially the larger and longer ones, were not designed primarily with cognitive outcomes in mind and did not assess cognition at baseline. It is therefore possible that some participants may not have been cognitively healthy at baseline, although we considered it unlikely that people with known dementia would have been recruited. In some cases, limited cognitive assessments, using measures which lack sensitivity to detect change in cognitively healthy people, were added to existing studies some years after onset. Participation in these studies was often restricted in ways which reflected the original aims, e.g. recruitment limited to one gender, or to people with particular comorbidities or risk factors, so the evidence was often not directly applicable to the older population in general. Only three studies specifically assessed the incidence of dementia. Studies which were designed to assess cognition tended to be shorter (less than two years), which may not be long enough to detect effects. If cognitive effects are detected on this timescale, it is not clear what implications this has for the incidence of dementia. There are problems inherent in combining short-term studies, in which separation of groups may only be seen if the intervention leads to cognitive improvement (or harm), and studies which are long enough to detect a change in the trajectory of age- or pathology-related decline. We considered it important to separate studies into groups of different durations, but our time divisions were inevitably somewhat arbitrary and this decision greatly limited the numbers of studies which could be combined.

There is no well developed understanding of the mechanisms by which any of these interventions might affect cognition, which may be the reason that studies used a wide variety of doses and treatment combinations. This makes it difficult to compare studies and leaves open the possibility that different doses or combinations could have different effects.

B vitamins

The body of evidence on vitamin B supplements addresses our review question only indirectly. Many studies restricted participation to those with particular comorbidities or who had some evidence of low vitamin B levels or exposure at baseline. Four of the 13 studies allowed inclusion of participants younger than 60 years, but in all cases, the mean age of participants was > 60 years so the data do not adequately address the effects of supplementation in midlife. We excluded very short-term studies (< 3 months), but the majority of included studies treated participants for 2 years or less, which may not be long enough to detect any effect. The particular vitamin B subtypes and their doses differed between studies and in several studies, particularly those with factorial designs, some participants receiving B vitamins also received other potentially active interventions. The studies were too heterogeneous to allow these various characteristics to be explored in subgroup analyses.

Antioxidant vitamins

We considered vitamin C, vitamin E, and beta-carotene in this group. The evidence on cognitive effects of antioxidant vitamins is very largely from studies which had other primary aims and to which very limited cognitive assessment was added after the studies were underway. In two of the large trials, all participants were at high vascular risk. However, the age of participants at recruitment was generally lower and the duration of the trials was generally longer than the B vitamin trials, so this body of evidence may be more informative about the potential of prolonged treatment to protect cognitive function against decline. Only two of these studies assessed incidence of dementia so most of the results have no more than indirect implications for dementia prevention. There was substantial variation in the doses of vitamins used. Again, the use of factorial designs in some studies meant the comparator groups included participants exposed to a variety of treatments.

Vitamin D

One study used a combination of vitamin D and calcium carbonate. Mean follow-up was 7.8 years. All participants were postmenopausal women, but inclusion criteria were otherwise broad. The dose of vitamin D was relatively low (400 IU/day). This was one of very few studies that used a rigorous method to assess for incident MCI or dementia. The other study used a higher dose

of vitamin D3 (4000 IU on alternate days), but was small and provided only preliminary evidence on supplementation with this dose.

Minerals (zinc and copper, selenium)

We found limited data from two studies of mineral supplements. One used a combination of zinc and copper oxides. It was designed to look for an effect on age-related macular degeneration, but there is no reason to think that this limits generalisibility of the cognitive results. Cognitive assessment was after a median of 6.9 years. Dementia incidence was not an outcome. The other was a selenium-only arm of a factorial study in which dementia incidence was the only relevant outcome reported. It found no evidence of an effect of selenium.

Complex multivitamins with or without minerals

There were three included studies in this category. One used a combination of B vitamins and antioxidant vitamins. Two used commercially available multivitamin and mineral combinations. Two studies were very small and lasted less than a year. In the large study, all participants were men, treated for a mean of 8.5 years. Dementia incidence was not an outcome.

Quality of the evidence

The large studies were generally well-conducted and we did not consider that risk of bias was a major threat to the results, although there were some gaps in reporting in all domains. Uncertainty about the risk of selection bias and concerns about attrition bias led us to downgrade the certainty of some results due to possible study limitations. We considered much of the evidence to be indirect with respect to our review question because participation in many studies was restricted, e.g. by comorbidities, baseline biochemistry, or gender. Overall, we considered the certainty of the evidence behind almost all effect estimates to be moderate or low with respect to our review question.

Potential biases in the review process

Particularly for B vitamins, bias may arise from pooling studies which differed in participants (e.g. with or without elevated homocysteine at baseline) and interventions (various constituents and doses). This 'broad brush' approach may disguise effects.

Agreements and disagreements with other studies or reviews

A comprehensive systematic review and meta-analysis of the cognitive effects of several years of treatment with B vitamins was

published by the B-Vitamin Treatment Trialists' Collaboration in 2014 (Clarke 2014). It included 11 trials for which data were available by September 2010. All the included trials were also included in this review. The authors investigated the effect of the interventions on homocysteine levels as well as on cognition. Although they found that the treatments effectively lowered homocysteine levels (by slightly more than 25%), they also found no evidence of significant effects on global ('MMSE-type') cognitive function, domain-specific cognitive function, or cognitive aging. We are not aware of similar systematic reviews of the effects of treatment with non-B vitamins or minerals.

tamins and minerals might modify these processes. Moving forward, there is a place both for investigating possible mechanisms in smaller trials, which can incorporate highly sensitive cognitive assessments and possibly biomarker outcomes, and for larger, longer, pragmatic trials in which it may be necessary to rely on simpler and less sensitive outcome measures. Considering the data in this review, the only real signals of effect were from long-term studies of antioxidant vitamins (vitamin C and beta-carotene) so these might be considered the most promising candidates for further research.

AUTHORS' CONCLUSIONS

Implications for practice

We found no evidence to support recommending any vitamin or mineral supplement to middle-aged or older adults who wish to maintain their cognitive function or prevent dementia.

Implications for research

All of the studies in this review differed in their population, interventions, and outcomes. Therefore, many questions about possible effects of different doses or treatment combinations on different participants remain unanswered. In particular, there are very few data on the long-term effects of supplementation started in midlife. Identifying the most important questions is likely to depend on a better understanding of the pathophysiology of cognitive decline and dementia and of the mechanisms by which vi-

ACKNOWLEDGEMENTS

The protocol for this review was largely based on a general template constructed for the development of a larger series of protocols and reviews covered by a National Institute for Health Research (NIHR) Systematic Reviews Programme Grant. The common protocol covered four types of intervention, for which some evidence exists that these may modify the risk of developing cognitive impairments or dementia. These include vitamin or mineral supplementation (or vitamin and mineral supplementation), exercise, cognition, and dietary interventions. The general protocol was inspired by a generic protocol approved by the Cochrane Musculoskeletal Group for another series of reviews (Da Costa 2012; Da Costa 2014; Nüesch 2009a; Reichenbach 2010; Rutjes 2009a; Rutjes 2009b; Rutjes 2010).

The authors are grateful to Professors Lisette de Groot, Simone Eussen, and Jane Armitage for responding to enquiries about their work and providing additional data.

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^{*} Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Anand 2011

Methods	2-arm, open-label RCT, with 3 months duration of treatment and follow-up
Participants	Location: Ootacamund, Tamil Nadu India. Single centre Recruitment: outpatient department, Government headquarters hospital, Ootacamund Sample size: • Number randomised: 38 in intervention, 36 in comparison • Number completed: 38 in intervention, 36 in comparison Participant baseline characteristics: • Age: mean 54.65 ± 8.44 intervention and 56.61 ± 8.17 in comparison • Total MMSE: 25.16 ± 0.66 in intervention, 25.3 ± 0.37 in comparison • Gender: male/female 15/23 intervention and 14/22 in comparison • Others: Most of the participants were from an economically deprived background, non-vegetarians. Inclusion criteria: • Type 2 diabetes with no comorbidities • Age between 35 and 65 of either sex • On oral hypoglycaemic agents (metformin, glibenclamide and combination) • Duration of disease less than or equal to 10 years • HbA1c > 9% Exclusion criteria: • Dementia • Antidepressant therapy • Type 1 diabetes • Juvenile diabetes • Juvenile diabetes • Pregnancy/lactation • Significant hepatic and renal dysfunction
Interventions	Intervention group (n = 38): vitamin E 600 mg once daily for three months Comparator group (n = 36): treatment-as-usual for three months Treatment received by both groups: oral hypoglycaemic agents (glibenclamide 5 mg, metformin 500 mg) for three months
Outcomes	Outcomes of interest in the review: Cognitive outcome: MMSE (Tamil version); MMSE 18-23 classified as MCI and 24-30 classified as no cognitive impairment
Notes	Funding Sources: "Nil". Declarations of interest: none declared. Other: population was from a government hospital clinic in India and "most of the enrolled patients were coming under economically deprived background."
Risk of bias	
Bias	Authors' judgement Support for judgement

Anand 2011 (Continued)

Random sequence generation (selection bias)	Low risk	Quote: "Enrolled patients were randomized by using computer assisted randomization procedure and assigned to control group and intervention group" Comment: adequate generation of a randomised sequence
Allocation concealment (selection bias)	Unclear risk	Comment: no description of allocation concealment. The study was open-label. Unknown whether allocation for the next patient to be enrolled could be known in advance
Blinding of participants and personnel (performance bias) All outcomes	High risk	Quote: "open-label trial" Comment: did not describe control group receiving placebo of Vitamin E. No de- scription of blinding and described as an 'open-label' study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Comment: no description of measures used, if any, to blind personnel involved in measuring outcomes. Since outcome measures depended on patient performance, risk of bias was judged to be high
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: There was no description of the completeness of outcome data for each main outcome, nor of exclusions from the analysis
Selective reporting (reporting bias)	Low risk	Comment: no evidence of selective reporting bias
Other bias	Low risk	No other risk of bias detected.

Andreeva 2011

Methods	Multicentre, randomised, placebo-controlled, 2 x 2 factorial trial An ancillary study of SU.FOL.OM3
Participants	Main study took place from 2003-2009. Quote: "As part of the SU.FOL.OM3 assessment battery, the F-TICS-m was administered during 2007-2009 only to participants who had completed 4 years of follow-up." Location: 257 participating centres in France Recruitment: recruitment of subjects done through a network of 417 cardiologists and neurologists from all over France, working either in general hospitals, independently or in rehabilitation centres. Treatment with tablets at home Sample size: 1881 participants were eligible for the cognitive study (had completed)

4 years of follow-up in main study); 1748 completed the cognitive assessment.

- · Number randomised in SU.FOL.OM3:
 - 620 in group A (B vitamins and Omega-3 PUFA)
 - 633 in group B (Omega-3 PUFA)
 - 622 in group C (B vitamins)
 - 626 in group D (placebo)

· Number completing cognitive study:

- 438 in group A (B vitamins and Omega-3 PUFA)
- 439 in group B (Omega-3 PUFA)
- 446 in group C (B vitamins)
- 425 in group D (placebo)

Group A (B Vitamins and Omega -3 PUFA) baseline characteristics:

- age (y): 61.6 ± 8.8
- gender, n (%) female: 80 (18.3)

Group B (Omega-3 PUFA) baseline characteristics:

- age: 60.1 ± 8.7
- gender, n (%) female: 88 (20.0)

Group C (B vitamins) baseline characteristics:

- age (y): 61.4±8.7
- gender, n (%) female: 93 (20.9)

Group D (Placebo) baseline characteristics:

- age (y): 60.9±8.9
- gender, n (%) female: 96 (22.6)

Inclusion criteria:

Men and women aged 45-80 y with a recent (1-12 months prior to inclusion) myocardial infarction (MI), unstable angina, or ischaemic stroke were eligible for participation There was no diagnostic criterion used to detect dementia/cognitive impairment. Instead, participants were given the Issac's Set Test at baseline and results were stated as a Mean \pm SD

Exclusion criteria:

Patients incapable of understanding the study protocol or who refused to sign the informed consent, patients with a pathology that might interfere with homocysteine or omega-3 fatty acid metabolism, in particular, those that use methotrexate for the treatment of a cancer or rheumatoid arthritis, chronic renal failure (plasma level of creatinine > 200 $\mu \rm mol/L$ or creatinine clearance < 40 mL/min), patients with a non-cardiovascular pathology with a suspected survival time less than the 5 years period of the study (solid cancer, evolved dementia, leukaemia) and patients taking B vitamins or omega-3 fatty acids

Interventions

Intervention (n = 1748): Study supplements were given as 2 capsules administered orally once per day. The supplements were provided in a package containing 53 blisters of 7×2 capsules, each corresponding to one week of both treatments. A year's supply was given at baseline and given again at each annual follow-up appointment for 5 years in total

Group A (B vitamins and Omega-3 PUFA): 5-methyltetrahydrofolate (folate, 0.56 mg) and vitamins B6 (3 mg) and B12 (0.02 mg) and eicosapentaenoic and docosahexaenoic acids (600 mg) in a 2:1 ratio

Group B (Omega-3 PUFA): eicosapentaenoic and docosahexaenoic acids (600 mg) in a 2:1 ratio

Andreeva 2011 (Continued)

	Group C (B Vitamins): 5-methyltetrahydrofolate (folate, 0.56 mg) and vitamins B6 (3 mg) and B12 (0.02 mg) Group D (placebo): 2 placebo Capsules Comparator group (n = 425): 2 placebo tablets given once daily. The placebo capsules looked and tasted identical to the active supplements (taken from SU.FOL.OM3 trial paper) Use of additional interventions (common to all treatment arms): no
Outcomes	Outcomes of interest in the review: Global cognitive functioning: French version of the modified Telephone Interview for Cognitive Status (F-TICS-m) Episodic memory: delayed recall subdomain from F-TICSm
Notes	Funding sources: The SU.FOL.OM3 trial was supported by the French Ministry of Research (grant R02010JJ), the Ministry of Health, Sodexo, Candia, Unilever, Danone, Roche Laboratories, Merck EPROVA AG, and Pierre Fabre Laboratories Declarations of interest: none declared Treatment adherence: Compliance with treatment was self-reported in the biannual questionnaires (or by telephone interview). Compliance was defined as taking at least 80% of the allocated treatment. (taken from SU.FOL.OM3 trial paper)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "We used computerized block randomisation (block size = 8) with stratification by sex, age (45-54, 55-64, and 65-80 y), prior CVD, and city of residence." Comment: adequate method of random sequence generation.
Allocation concealment (selection bias)	Low risk	Comment: no information. Centralised, computerised system, likely to be adequately concealed
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "The placebo capsules looked and tasted identical to the active supplements" Comment: probably blinded
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Comment: did not mention blinding of study personnel including outcome assessors
Incomplete outcome data (attrition bias) All outcomes	High risk	Comment: 24.8% of participants in SU. FOL.OM3 were lost to follow-up over the first 4 years and before the onset of the cognitive study. These participants appeared to

Andreeva 2011 (Continued)

		be well balanced between groups 133/1881 (7%) participants eligible for the cognitive study were excluded: 72 (3.8%) were unable to complete the cognitive assessment and 61 refused to participate. Treatment allocations of these participants were not reported Cognitive data were reported for 69.9% of the randomised participants
Selective reporting (reporting bias)	Unclear risk	Quote: "No changes to the outcomes of interest were made after launching the trial" Comment: The authors have stated no changes to the outcomes were made, however, these were not previously published in a protocol as this is an ancillary study and the main study protocol did not mention any cognitive outcomes
Other bias	Low risk	No other risks identified

Aspell 2017

Methods	2-arm, parallel group RCT
Participants	Location: one site in Ireland Recruitment: no details provided Number randomised: 60 Participant characteristics: • age: mean 68.5 years (SD 4.9 years) • gender: 53.3% female Inclusion criteria: community dwelling Exclusion criteria: low or high serum vit D (defined as < 15nmol/L or > 125nmol/L), current use of supplemental vit D ≥ 800 IU/day, screen positive for cognitive impairment using the Telephone Cognitive Screen (TCogS), hypercalcaemia, hyperparathyroidism, epilepsy, stroke, renal disease, schizophrenia, bipolar affective disorder, recurrent psychotic depression, alcohol and drug abuse within the past 5 years, anti-convulsants, anti-psychotic medications, significant hearing difficulties even when wearing hearing aids, illness that caused permanent decrease in memory or other mental function
Interventions	50 mcg/day of vitamin D3 administered as 4000 IU on alternate days or placebo
Outcomes	Overall cognitive function: Montreal Cognitive Assessment (MoCA) Executive function: (Trails Making Task A & B, B-A) Episodic memory: Wechsler Memory Scale - in trial registry entry, not reported Attention and visual reasoning: 2 validated measures (not included in review)

Aspell 2017 (Continued)

Notes Described as a pilot study. Only conference abstract available 18.3% of participants were deemed vitamin D deficient at baseline according to Institute of Medicine guidelines
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Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomised", no additional details
Allocation concealment (selection bias)	Unclear risk	No details
Blinding of participants and personnel (performance bias) All outcomes	Low risk	From trial registry entry: masking: double (participant, investigator)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	From trial registry entry: masking: double (participant, investigator)
Incomplete outcome data (attrition bias) All outcomes	Low risk	"A retention rate of 93.8% was achieved." No information on treatment assignment of missing participants
Selective reporting (reporting bias)	High risk	Trial registry entry included a measure of episodic memory which was not reported with other results in the conference abstract
Other bias	Low risk	No other risks identified

Cockle 2000

Methods	2-arm, parallel group, double-blind, placebo-controlled RCT (Intended duration was 48 weeks, but due to very high dropout after 24 weeks, no data were reported beyond 24 weeks)
Participants	Location: One site in UK Recruitment: subjects recruited by newspaper advertisement Number randomised: 69 in intervention, 70 in comparison Participant baseline characteristics: • Age: • overall: median 70 years • placebo: mean 70.2 (5.4) and median 70 (60-81) • intervention: mean 70.7 (5.6 and median 70 (60-81) • Gender: • overall: 51 males and 88 females • placebo (24 males and 46 females) • intervention (27 males and 42 females)

Cockle 2000 (Continued)

Bias	Authors' judgement Support for judgement
Risk of bias	
Notes	Quote: "the objectives of the study were to examine the relationship between vitamin status and cognitive functioning and mood in healthy, elderly volunteers and to assess the subsequent impact of multivitamin supplementation on these measures" Funding sources: none stated Declarations of interest: none stated, but second author from Roche Pharmaceutical
Outcomes	Primary Outcomes: Relationship between vitamin status and cognitive function. "At each visit, treatment compliance, blood vitamin levels and Critical Flicker Fusion, Choice ReactionTimeTask, Sternberg Memory ScanningTask, Word ScanTask, and the Profile Of Mood States questionnaire were carried out" "MMSE, Sundrom Kurtz Test, National Adult Reading Test, Alice Heim's 4, Alice Heim's 5 and the Geriatric Depression Scale were assessed at baseline and at week 48." Eligible cognitive outcomes: • episodic memory measured with Sternberg Memory Scanning Task (msec), lower values indicated benefit • episodic memory measured with Word scan task (msec), lower values indicated benefit* • speed of processing measured with Total reaction time (msec), lower values indicated benefit • speed of processing measured with Recognition reaction time (msec), lower values indicated benefit • speed of processing measured with Motor reaction time (msec), lower values indicated benefit • speed of processing measured with Motor reaction time (msec), lower values indicated benefit • speed of processing measured with Motor reaction time (msec), lower values indicated benefit • speed of processing measured with Motor reaction time (msec), lower values indicated benefit
Interventions	Intervention (n = 69): Multivitamin supplements: two gelatine capsules orally each day. Each capsule contained vitamin A palmitate 3334 IU, thiamine mononitrate 14 mg, riboflavin 16 mg, pyridoxine 22 mg, vitamin B12 0.03 mg, vitamin C 600 mg, dl-alphatocopherol acetate 100 mg, folic acid 4 mg, d-biotin 2 mg and nicotinamide 180 mg Quote: "The dosing was calculated to provide the equivalent of one US RDA (National Research Council, 1980) of vitamin A and ten US RDAs of the remaining vitamins." Capsules supply were provided at baseline and, in patients who were compliant, a second supply was given at week 12 for an intended treatment duration of 24 weeks Comparator group (n = 70): matching placebo (rape seed oil) Use of additional interventions (common to both treatment arms): not reported
	 Main diagnosis: All subjects were in good physical and mental health and had no history of significant disease or mental illness, as determined by a medical examination and health questionnaire. MMSE: intervention: 28.9 (SD:1.4) placebo: 28.6 (SD:1.5) Inclusion criteria: not reported Exclusion criteria: not reported

Cockle 2000 (Continued)

Random sequence generation (selection bias)	Unclear risk	Quote: "Subjects were allocated to each treatment group according to a predetermined randomization schedule." Comment: insufficient information about the sequence generation method
Allocation concealment (selection bias)	Unclear risk	Quote: "Subjects were allocated to each treatment group according to a predetermined randomization schedule." Comment: insufficient information about allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "The study was a double-blind, placebo-controlled, parallel group study with each subject receiving a multivitamin supplement or placebo daily for up to 24 weeks." Quote: "Treatments were given orally as two soft gelatine capsules containing either multivitamins or matching placebo (rape seed oil)" Comment: The study was reported as double-blinded with the use of matching placebo
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "at each visit, treatment compliance, blood vitamin levels and Critical Flicker Fusion, Choice Reaction Time Task,were carried out" Comment: It was not reported whether outcome assessment was blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: 'Due to circumstances not related to the study treatment, there was a significant dropout of subjects after the first six months of this study, with no participants continuing through to visit 4. Data from 139 subjects was available at baseline, from 127 subjects at visit 1 and from 90 subjects at visit 2. Due to the high dropout of subjects between visit 2 and visit 3, data from the latter visit was not included." Comment: There were 12 dropouts (8.6%) at visit 1 (12 weeks) and 49 dropouts (35. 8%) at visit 2 (24 weeks). Outcome data were completely missing from planned visits 3 and 4 (36 and 48 weeks)

Cockle 2000 (Continued)

S	Selective reporting (reporting bias)	High risk	Comment: A number of tests of cognition and functioning reported in the Methods to have been measured at baseline and 48 weeks were not reported (including MMSE)
(Other bias	Low risk	No other risks identified

Dangour 2015	
Methods	2-arm, double-blinded, multicentre, parallel group RCT, with 12 months of treatment and follow-up
Participants	Location: 7 general practices in South East England Recruitment: recruited at GP by mail, baseline appointment and 12-month follow-up at King's College Hospital, London, treatment one tablet daily at home Sample size: Number randomised: 99 in intervention, 102 in comparison Number completed: 91 in intervention, 92 in comparison Group A (B vitamins) baseline characteristics: age (mean ± SD): 79.943.5 gender (n): 46 males, 54 females main diagnosis: no diagnosis Group B (placebo) baseline characteristics: age: 80.1 ± 3.7 gender: 48 males, 54 females main diagnosis: no diagnosis (Inclusion criteriat) Age: ≥ 75 years, Participants with a Mini-Mental State Examination score > 24 (maximum score 30) were asked to provide a blood sample to assess serum vitamin B12 and haemoglobin concentration. Those with moderate vitamin B12 deficiency who did not have anaemia (serum vitamin B12 concentrations ≥ 107 and < 210 pmol/L were eligible to join the trial The diagnostic criterion used for dementia/cognitive impairment screening was the MMSE. Only those with a score > 24 were eligible to join the trial (Exclusion criteria:) Diabetes, dementia, epilepsy, alcohol addiction, pacemakers (and other implanted metallic devices where central neurophysiologic testing was contraindicated), residents of nursing homes, diagnosis of pernicious anaemia, current consumption of vitamin B12 supplements and those who had received a vitamin B12 injection within the previous 6 months. Individuals with very low vitamin B12 concentrations (< 107 pmol/L, which is a cutoff typically used for deficiency) or who were shown to have anaemia (haemoglobin concentration, 110 g/L for women and 120 g/L for men) were excluded
Interventions	Intervention (n = 97): vitamin B12 (cyanocobalamin), oral administration, 1 mg tablet/day, 12 months Comparator group (n = 97): placebo tablet administered once daily identical to intervention in shape, size, colour, smell, taste, and identically packaged

Dangour 2015 (Continued)

	Use of additional interventions (common to both treatment arms): not reported	
Outcomes	Outcomes of interest in the review: • Episodic memory measured with California Verbal Learning Test at delayed r with higher values indicating benefit • Executive function measured with verbal fluency, with higher values indicating benefit • Speed of processing measured with Choice Reaction time, with lower values indicating benefit	
Notes	Funding sources: Supported by the Food Standards Agency (N05072) and the Department of Health. National Health Service Research and Development and King's College Hospital Trust Research and Development provided service support costs Declarations of interest: none declared Treatment adherence: Adherence was measured by counting the number of tablets returned at the end of the study	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Allocation codes were obtained from a central computerized randomiza- tion service." Comment: adequate method of random se- quence generation
Allocation concealment (selection bias)	Low risk	Quote: "Allocation codes were obtained from a central computerized randomiza- tion service." Comment: adequate method of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "All study personnel were blinded to the treatment allocation." "Allocated treatment consisted of a single tablet administered daily that was identical in size, shape, color, smell, and taste for both the intervention and placebo and packaged into identical pots." Comment: As all participants were given identical tablets this ensured blinding was maintained
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "All study personnel were blinded to the treatment allocation." "Allocated treatment consisted of a single tablet administered daily that was identi-

Dangour 2015 (Continued)

		cal in size, shape, color, smell, and taste for both the intervention and placebo and packaged into identical pots." Comment: Study mentions personnel were blinded to treatment allocation and due to identical tablets this would have maintained appropriate blinding
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: Cognitive outcome data were available for 184/201 participants (92%). Dropout rates and reasons were similar in both groups
Selective reporting (reporting bias)	Low risk	Comment: followed study protocol, no reporting bias found
Other bias	Low risk	Comment: none found

Durga 2007

Methods	2-arm, placebo-controlled, randomised clinical trial, with 3-year duration of treatment
Participants	Location: Gelderland region in the Netherlands, single centre
	Recruitment: Division of Human Nutrition, Wageningen University, Wageninger
	Netherlands
	Number randomised: 406 in intervention, 413 in comparison
	Participant baseline characteristics:
	• Age at baseline (years), mean (SD):
	o placebo: 60 (6)
	o folic acid: 60 (5)
	• Male, n (%):
	o placebo: 292 (70%)
	o folic acid: 294(72%)
	• MMSE (points):
	o placebo: 29 (28-30)
	o folic acid: 29(28-30)
	Inclusion criteria:
	 Participants were men and post-menopausal women aged 50-70 years.
	More than 80% self-reported compliance during a 6-week placebo run-in period
	was required.
	Exclusion criteria:
	• homocysteine concentration less than 13 micromol/L and greater than 26
	micromol/L
	self-reported medical diagnosis of renal or thyroid disease
	self-reported use of medications that influence folate metabolism
	 participants reportedly used B vitamin supplements or drugs that could affect
	atherosclerotic progression

Durga 2007 (Continued)

Interventions	Intervention (n = 405): 800 μ g per d Comparator group (n = 413): placeb	·	
Outcomes	test, verbal fluency test, letter digit sub These 5 tests were used to construct 5	a priori defined cognitive domains using z-scores: a speed, information processing speed, and word ge of 5 domains); liate recall' $(Z_{VerbalFluencytest})$	+Z _{15-W}
Notes	for Health Research and Development University, and Wageningen Centre of Sciences is an alliance of major Dutch government. Wageningen Centre for Fithe development of new and innovated Declarations of interest: Jane Durg Lausanne, Switzerland and Petra Verhoo Research Institute in Vlaardingen, the entails examining the health benefits of However, the study reported in the cur <i>The Lancet</i> before the authors joined to	ded by a grant from the Netherlands Organisation (ZonMw, grant number 20010002), Wageningen or Food Sciences. Wageningen Centre for Food food industries, research institutes and the Dutch ood Sciences does long-term strategic research for d food with attention to health aspects a currently works at Nestle Research Center in efficurrently works at the Unilever Food and Health Netherlands. The work at both food companies a variety of food ingredients, including folic acid. The transcript was completed and submitted to the companies, when they were still employed by the Centre for Food Sciences. All authors declare	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Quote: "Patients were allocated treatment or placebo with permuted blocks of sizes	

four and six, which varied randomly. Specialised staff who were not involved in the study allocated and labelled the capsule boxes with participants' unique sequence number. Participants in the same household received the same treatment."

Comment: Method of sequence generation

was not reported.

Durga 2007 (Continued)

Allocation concealment (selection bias)	Low risk	Quote: "Specialized staff who were not involved in the study allocated and labelled the capsule boxes with participants' unique sequence number" Comment: appropriate allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "The folic acid and placebo capsules were indistinguishable in appearance". 70% of participants in the folic acid group and 71% in the placebo group thought they had been allocated to folic acid Comment: well blinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Comment: All staff, including all authors, were unaware of group assignment until completion of the trial and after data analyses Quote: outcome assessment well blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: 17 participants lost to follow-up were assigned the median test score of the total population at the end of the study Comment: Although there were imputations, the small proportion of loss to follow-up unlikely to affect results
Selective reporting (reporting bias)	Unclear risk	Quote: From clinicaltrials.gov: "Cognitive performance at year 3 (cognitive domains: simple speed, cognitive flexibility, and memory; and information processing speed and semantic memory)" Comment: unclear if the domains reported were prespecified (authors were contacted for further information)
Other bias	Low risk	No other risks detected

Eussen 2006

Methods	3-arm, parallel-group, randomised, placebo-controlled trial, 24 weeks duration
Participants	Location: the Netherlands Recruitment: free-living older persons and older persons living in care-facility homes, recruited via mailed health questionnaires Number randomised: 195 randomised, of whom 119 had CDR 0 at baseline and were included in this review "Randomization was stratified according to MMA concentration at the screening visit

(< and > 0.45 μ mol/L), age (< and > 80), sex and MMSE score (< and > 24 points)." Vitamin B12: 38 Vitamin B12 + folic acid: 38 Placebo: 43 Participant baseline characteristics: • Age, mean (SD): o vitamin B12: 80.42 (5.52) o vitamin B12 + folic acid: 80.82 (4.18) o placebo: 79.86 (4.74) • .Gender o vitamin B12: female 29/38 (76%) o vitamin B12 + folic acid: female 29/38 (76%) o placebo: female 34/43 (79%) • Baseline cognitive function - MMSE score (mean (SD)) o vitamin B12: 28.18 (1.43) o vitamin B12 + folic acid: 28.34 (1.40) o placebo: 27.93 (1.80) **Inclusion criteria:** aged ≥ 70 y. Mild vitamin B12 deficiency defined as (1) a serum vitamin B12 concentration between 100 and 200 pmol/L, or (2) a serum vitamin B12 concentration between 200 and 300 pmol/L, a plasma MMA concentration ≥ 0.32 μ mol/L, and a serum creatinine concentration $\leq 120 \mu$ mol/L. Ingested 90% or more of capsules during a 2-week placebo run-in period prior to randomisation Exclusion criteria: history of cobalamin deficiency, use of cobalamin (> 50 µg/day) or folic acid (> 200 µg/day) supplementation or injections, surgery or diseases of the stomach or small intestine, anaemia, dementia, life-threatening diseases, or severe hearing or visual problems Interventions Intervention: 1. vitamin B12: 1000 μg vitamin B12 (cyanocobalamin) per day orally for 24 weeks 2. vitamin B12 and folic acid: 1000 μg vitamin B12 (cyanocobalamin) plus 400 μg folic acid per day orally for 24 weeks Comparator group: • Placebo capsule The placebo capsules contained AVICEL PH102 (Medipulp GmbH, Aschaffenburg, Germany) as a filler Use of additional interventions (common to all treatment arms): not reported Outcomes Cognitive function Cognitive function was assessed before and after 24 wk of treatment with the use of an extensive neuropsychologic test battery that included the domains of attention, construction, sensorimotor speed, memory, and executive function Cognitive function was assessed by 6 trained and registered neuropsychologists during the run-in period (baseline) and at week 24 of the intervention during a 1.5-2-h session Neuropsychological test battery included: • finger tapping, computerised • motor planning 2 and 3, computerised • Figure of Rey copy, immediate recall and delayed recall • 15-word learning immediate recall, delayed recall and recognition • Trail-making test, part A and part B

Eussen 2006 (Continued)

	Digit span forward and backward	
	• Raven	
	• Stroop test	
	• Similarities, WAIS	
	Word fluency, letter	
	Word fluency, animals	
	Depression measured with GDS	
	Biochemical measures: vitamin B12, MMA, holoTC, homocystine, RBC folate	
	Compliance was checked by counting the number of unused capsules remaining in cap-	
	sule dispensers and by verifying pill counts in the participants' diaries. Mean compliance	
	was 99%	
Notes	Professor Simone Eussen and Professor Lisette de Groot kindly provided data separately on participants with CDR 0 at baseline for inclusion in this review	
	Study supported by grants from ZON-MW, The Hague, Netherlands; Kellogg's Benelux,	
	Zaventem, Belgium; Foundation to Promote Research into Functional Vitamin B12	
	Deficiency and the European Union BIOMED Demonstration Project; Nutricia Health	
	Foundation, Wageningen, Netherlands	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomized", no further information
Allocation concealment (selection bias)	Unclear risk	No information
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quotes: "The capsules given to the separate treatment groups were identical in appearance, smell and taste." "The study had a double-blind design." Comment: participants and personnel blind to allocation
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The study had a double-blind design." No specific mention of outcome assessors" Comment: outcome assessors probably blind to allocation
Incomplete outcome data (attrition bias) All outcomes	Low risk	Incomplete outcome data varied between cognitive tests. For included outcomes, data available for minimum of 82% of participants In study as a whole, "16% were unable to complete the trial, mostly because of illness, and the dropout rate was slightly higher in the vitamin B12 + folic acid group than in

Eussen 2006 (Continued)

Interventions

		the other groups." Dropout in whole study: 10/64 B12, 15/66 B12 + folic acid, 8/65 placebo Comment: major effect of differential dropout unlikely
Selective reporting (reporting bias)	Low risk	All outcomes mentioned in methods fully reported
Other bias	Low risk	No other biases identified
Ford 2010		
Methods	2-arm, double-blinded, parallel group RCT, with 2 years duration of treatment and 8-year follow-up The average follow-up time (time to event) was 8.3 years in both groups	
Participants	Location: Australia, number of sites not reported Recruitment: community-representative hypertensive men 75 years and older randomly selected from a large population-based study of abdominal aortic aneurysm screening (Health in Men Study) Participants were recruited from July 18, 2000, to February 6, 2002 Number randomised: 150 in intervention, 149 in comparator Participant (baseline) characteristics: • Age, mean (SD): • placebo: 78.7 (2.7) • vitamin: 79.3 (2.8) • Baseline cognitive function • ADAS-cog, mean (SD) • placebo: 11.8 (6.0) • vitamin: 11.0 (5.1) • MMSE score, mean (SD) • placebo: 27.6 (1.9) • vitamin: 27.5 (1.8) Inclusion criteria: community-representative hypertensive men 75 years and older Exclusion criteria: Beck Depression Inventory (BDI) score ≥ 18 or a Mini-Mental State Examination (MMSE) score ≤ 24, illness likely to cause severe disability or death within 12 months (for example, metastatic cancer, Parkinson disease, or a history of stroke), non-English speaking, living in residential care facilities, or patients who were already taking vitamin B supplements	

Comparator group (n = 149):

Intervention (n = 150):

Vitamins: one capsule containing 400 μg B12, 25 mg B6, 2 mg folic acid, every morning

Use of additional interventions (common to both treatment arms): not reported

Placebo, one identical oral capsule every morning for 2 years

Outcomes

Primary Outcomes: to ascertain whether treatment with vitamins B12, B6, and folic acid over 2 years would improve cognitive performance (change in the cognitive subscale of the Alzheimer's Disease Assessment Scale) compared with placebo in older men

Secondary Outcomes:

Longer-term impact of B vitamin supplementation by comparing performance on the TICS as well as a diagnosis of dementia registered in the Western Australian Data Linkage System (WADLS) until September 30, 2009

A diagnosis of dementia was recorded if the death certificate contained one of the following ICD-10 codes: F00 (dementia in AD), F01 (vascular dementia), F03 (unspecified dementia), and G30 (AD). Additionally, a dementia diagnosis was sought from records of inpatient admissions and outpatient contacts

Secondary measures of interest included also the California Verbal Learning Test, Mini-Mental State Examination, Digit Cancellation Test, Clock Drawing Test*, TICS, changes in biochemical parameters (total plasma homocysteine, vitamin B12 and quality of life (Short Form (SF)-36)

Eligible cognitive outcomes:

- global cognitive function measured with ADAS-cog at 1 and 2 years, with higher values indicating benefit
- global cognitive function measured with MMSE at 1 and 2 years, with higher values indicating benefit*
- episodic memory measured with California Verbal Learning Test 1 (CVLT 1) at 1 and 2 years, with higher values indicating benefit
- ullet episodic memory measured with California Verbal Learning Test 2 (CVLT 2) at 1 and 2 years, with higher values indicating benefit*
- semantic memory measured with Clock Drawing Test at 1 and 2 years, with higher values indicating benefit*
- speed of processing measured with Digit Cancellation Test (DCT) at 1 and 2 years, with higher values indicating benefit*

Function outcome extracted: none reported

Quality of life outcome: SF-36

Safety outcome: none reported

* outcome data used in statistical analyses

Notes

Funding sources, quote: "Supported by the National Health and Medical Research Council of Australia (139123 to L.F.). The vitamin and placebo tablets were provided free of charge by Blackmores Ltd"

Declarations of interest, quote: "Dr. Ford reports no disclosures. Dr. Flicker serves as Internal Medicine Editor for Geriatric Medicine, Associate Editor for BMC Geriatrics, on the editorial board of the Australasian Journal on Ageing, and as Editor of the Cochrane Dementia and Cognitive Improvement Group; and receives/has received research support from Pfizer Inc. and NHMRC. Dr. Alfonso and J. Thomas report no disclosures. Dr. Clarnette serves on scientific advisory boards for Lundbeck Inc., Pfizer Inc., and Novartis; serves as an editorial advisor to Geriatric Medicine in General Practice; and estimates that 10% of his practice at Fremantle Hospital consists of cognitive testing. Dr. Martins serves as Senior Editor for the Journal of Alzheimer's Disease; serves as a consultant for and holds stock in Alzhyme Ltd.; and receives research support from Commonwealth Scientific and Industrial Research Organisation (Australia). Dr. Almeida has received funding for travel from Blackmores Ltd."

Treatment adherence

Ford 2010 (Continued)

Men who consumed at least 75% of the study tablets during this 2-year trial were considered compliant. Compliance was determined by pill count and medication diaries

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Participants were given consecutive numbers and allocated to active vs placebo arms based on computer-generated random permuted blocks. Blocks consisted of 8 subjects (4 subjects allocated to each group) so as to minimize the risk of having unbalanced entry into each arm of the study during the period of recruitment." Comment: adequate method of sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "An external and independent academic controlled the randomization procedures of the trial." Comment: adequate method of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Vitamins and placebo were administered in the form of identical oral capsules." "Participants and investigators were blinded to the group membership of men in the trial until the last follow-up assessment was completed. There were no breaches of protocol" Comment: participants and personnel blinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Participants and investigators were blinded to the group membership of men in the trial until the last follow-up assessment was completed." Comment: outcome assessment blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "32 (21%) men in the intervention and 26 (17%) in the placebo group withdrew consent or were lost during the trial." Comment: For the primary outcome, an ITT analysis was presented and attrition bias was considered as low For the secondary outcome of cognitive

Ford 2010 (Continued)

		impairment or dementia at long-term fol- low-up: "Seventy-three of these men took part in a later telephone survey in 2008 and completed the Telephone Interview for Cognitive Status (TICS) questionnaire", "41 (27%) in the intervention group and 32 (21%) in the placebo group". Data on dementia diagnosis were available on all 172/299 (57.5%) participants who were alive in 2008 Comment: high risk of attrition bias for this long-term secondary outcome
Selective reporting (reporting bias)	Low risk	Comment: All outcomes specified in the Methods were reported in the Results. Protocol not available
Other bias	Low risk	No other risks identified

Grodstein 2007

Methods	Substudy of the Physicians' Health Study II, a randomised, double-blind, placebo-controlled, 2 x 2 x 2 factorial trial testing \$\mathcal{B}\$-carotene, vitamin E, ascorbic acid, and a multivitamin for their role in preventing chronic diseases including total and prostate cancer, cardiovascular disease, and age-related eye disease among 14,641 male physicians aged 50 years or older The trial was done from 1997 to 1 June 2011. The cognitive function substudy began in 1998. Those eligible for the cognitive substudy were PHS II participants older than 65 years in November 1998 The Physicians' Health Study II was a continuation of the Physicians' Health Study, which began in 1982 and had randomised male participants to low-dose aspirin and \$\mathcal{B}\$-carotene. Participants included those continuing their original \$\mathcal{B}\$-carotene assignment from the Physicians' Health Study and newer recruits randomised from 1998 onwards
Participants	Location: not reported Recruitment: The Physicians' Health Study II. Quote: "In July 1997, invitations to enrol in PHS II were mailed to eligible participants from PHS I who had been part of an earlier trial of aspirin and β-carotene among 22071 physicians aged 40 to 84 years in 1982. Second, in July 1999, invitation letters were mailed to a new group of male physicians identified from a list provided by the American Medical Association." Quote: "PHS I participants who are willing and eligible for participation in PHS II will retain their randomized β-carotene assignment from PHS I and will be also randomized to vitamin E" "new physician participants will be randomly assigned to the same interventions" Number randomised: • 2967 β-carotene (2031 continuing from PHS and 936 newer recruiters) • 2989 placebo (2021 continuing from PHS and 968 newer recruiters) Participant (baseline) characteristics (at entry PHS III): All participants were men.

Ace	mean:
1120,	mican.

- Patients from PHS at entry in PHS II
 - o ß-carotene: 70.9
 - o placebo: 71.0
- Newly recruited
 - ß-carotene: 72.9placebo: 72.8

Inclusion criteria: not reported

Exclusion criteria: history of cirrhosis or active liver disease, patients receiving anticoagulants, or reported a serious illness that may interfere with study participation Men were also required to forgo current use of multivitamins or individual supplements

containing more than 100% of the recommended daily allowance of vitamin E, vitamin C, \(\mathcal{B}\)-carotene, or vitamin A during PHS II follow-up

For those newly recruited: no history of cancer, active liver disease, current renal disease, peptic ulcer, or gout

Quote: "Only physicians who comply with the pill-taking regimen at least two-thirds of the time and remain willing and eligible to participate will be subsequently randomized"

Interventions

Intervention:

- 1. ß-carotene (Lurotin, 50 mg on alternate days, or its placebo; BASF, Florham Park, New Jersey):
- 2. vitamin E (synthetic alfa-tocopherol, 400 IU on alternate days, or its placebo; BASF);
 - 3. ascorbic acid (synthetic ascorbic acid, 500 mg daily, or its placebo; BASF);
- 4. multivitamin (Centrum Silver or its placebo daily; Pfizer, New York, New York) NB: Grodstein 2007 reported on the β-carotene and Grodstein 2013 on multivitamin supplementation.

Comparator group: placebo ß-carotene.

Exposure to study intervention:

2031 patients in ß-carotene and 2021 placebo continued from the PHS study (started in 1982) in which they were receiving ß-carotene versus placebo ß-carotene. As most of the cognitive assessments were done in 2001, some of the patients continuing from PHS had been exposed to ß-carotene (or placebo ß-carotene) for about 19 years, whereas the "new recruiters" had been exposed from 1997 to 2001, thus for about 4 years

The mean time from randomisation to cognitive assessment was 18 years (range, 15-20 years) in 4052 continuing participants from the PHS, and was 1 year in 2021 new recruits (range, 2 months-3 years)

Quote: "PHS participants, who remained blinded to beta carotene assignment."

Quote: "Treatment assignment to beta carotene or beta carotene placebo was retained from the PHS (although participants may have stopped taking beta carotene during the 18-month interval between studies), and the men were newly randomized to receive vitamin E, ascorbic acid, multivitamin, or placebo."

Use of additional interventions (common to both treatment arms): not reported

Outcomes

Primary Outcomes:

A global composite score averaging 5 tests of global cognition, verbal memory, and category fluency

The cognitive battery included the Telephone Interview for Cognitive Status (TICS); immediate and delayed recall on the East Boston Memory Test (EBMT); delayed recall

Grodstein 2007 (Continued)

of a 10-word list; and a category fluency task

The cognitive function substudy began in 1998.

The beta carotene arm of the PHS II continued until May 2003, its planned stopping date, but most of the assessments were done in 2001

Quote: "Second cognitive assessments were begun in 2002. The beta carotene arm was terminated prior to completion of second assessments (second and third assessments were planned to allow further data collection in the continuing trial of other vitamins). Nonetheless, some data from the second interview were available (n = 4074, with 88. 3% participation of those who were contacted before termination of the beta carotene study)."

Eligible cognitive outcomes:

- global cognitive function measured with TICS*
- episodic memory (verbal memory) measured using an average of z-scores* for immediate and delayed recall on the EBMT and delayed recall of a 10-word list
- executive functioning measured with category fluency: animals named during 1 minute*

Function outcome extracted: none reported

Quality of life outcome extracted: none reported

Safety outcome extracted: none reported * outcome data used in statistical analyses

Notes

Funding source: "This study was supported by grants from the National Institutes of Health (CA34944, CA40360, CA97193, HL26490, HL34595, and AG15933), and from BASF Corporation (Florham Park, New Jersey), Wyeth (New Jersey), and DMS (New Jersey). Dr Grodstein was partially supported by a New Scholars in Aging award from the Ellison Medical Foundation. Role of the Sponsor: No supporting organization had any role in the study design; conduct of the study; collection, management, analysis, and interpretation of the data; or preparation, review, or approval of the manuscript." **Declaration of interest:** "None reported".

Treatment adherence:

ß-carotene study:

"Overall, 79.3% reported taking at least 2 of 3 of their study pills; this was similar in those assigned to active treatment (79.1%) or placebo (79.5%), and in the continuing participants from the PHS (79.2%) vs new recruits (79.4%)"

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" "Randomizationusing a computer-generated list of random numbers" Comment: adequate method of random generation

Grodstein 2007 (Continued)

Allocation concealment (selection bias)	Unclear risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" Comment: method of allocation concealment not reported
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" Comment: method of blinding not reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" "An Endpoint Committee of physicians blinded to the participants' treatment assignment, will review the medical records for final confirmation of a reported diagnosis" Comment: The method of blinding of outcome assessment for the cognitive substudy was not reported. The statement about the Endpoint Committee is contained in the design paper which, however, does not mention any cognitive outcome
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "of the 7045 age-eligible patients, 271 were no longer active PHSII participants (3.8%) and 4 were deceased (0.06%). Of the remaining 6770, we were unable to reach 315 (4.7%). Of the 6455 participants contacted, 5956 (92.3%) completed a cognitive assessment: 4052 from the original PHS and 1904 new recruits" "Participation (in the cognitive substudy) was virtually identical in those assigned to betacarotene (92.0%) vs placebo (92.6%) and in continuing participants (from the PHS) (92.1%) vs new recruits (to PHS II) (92.7%)."
Selective reporting (reporting bias)	Low risk	Comment: no differences found between the protocol (clinicaltrials.gov Identifier: NCT00270647) and article. In addition, all outcomes specified in the Methods were reported in the Results

Other bias	Low risk	No other source of bias identified	
Grodstein 2013			
Methods	trolled, 2 x 2 x 2 x 2 facto multivitamin for their role cancer, cardiovascular disea- aged 50 years or older	The trial was done from 1997 to 1 June 2011. The cognitive function substudy began	
Participants	enrol in PHS II were mailed earlier trial of aspirin and be in 1982. Second, in July 1 physicians identified from a The substudy of cognitive factor of the substudy of the substudy of cognitive factor of the substudy of the substance of the substudy of the subst	Location: not reported Recruitment: The Physicians' Health Study II. Quote: "In July 1997, invitations to enrol in PHS II were mailed to eligible participants from PHS I who had been part of an earlier trial of aspirin and beta-carotene among 22,071 physicians aged 40 to 84 years in 1982. Second, in July 1999, invitation letters were mailed to a new group of male physicians identified from a list provided by the American Medical Association." The substudy of cognitive function was conducted among PHS II participants aged 65 years or older Number randomised: 2980 in multivitamin, 2967 in placebo Participant (baseline) characteristics: All participants were men. • Age, mean (SD): o multivitamin: 71.6 (6.0) placebo: 71.6 (5.9) Inclusion criteria: not reported Exclusion criteria: history of cirrhosis or active liver disease, patients receiving anticoagulants, or reported a serious illness that may interfere with study participation Men were also required to forgo current use of multivitamins or individual supplements containing more than 100% of the recommended daily allowance of vitamin E, vitamin C, ß-carotene, or vitamin A during PHS II follow-up For those newly recruited: no history of cancer, active liver disease, current renal disease,	
Interventions	New Jersey); 2. vitamin E (synthetic a BASF); 3. ascorbic acid (synthetic 4. multivitamin (Centru NB: Grodstein 2007 reportsupplementation. Comparator group: placeby	io mg on alternate days, or its placebo; BASF, Florham Park, lfa-tocopherol, 400 IU on alternate days, or its placebo; ic ascorbic acid, 500 mg daily, or its placebo; BASF); m Silver or its placebo daily; Pfizer, New York, New York) ted on the ß-carotene and Grodstein 2013 on multivitamin on multivitamin nations (common to both treatment arms): Because of the	

Grodstein 2013 (Continued)

	factorial design, some participants were also taking vitamin E and/or vitamin C Exposure to study intervention: Mean time from randomisation to initial cognitive assessment was 2.5 years (range, 0.18 to 5.3 years) and mean time from randomisation to the final cognitive assessment was 8.5 years (range, 0.3 to 14.2 years)
Outcomes	Primary Outcomes: A global composite score averaging 5 tests of global cognition, verbal memory, and category fluency The cognitive battery included the Telephone Interview for Cognitive Status (TICS); immediate and delayed recall on the East Boston Memory Test (EBMT) (verbal memory); the delayed recall of a 10-word list (verbal memory); and a category fluency task The cognitive function substudy began in 1998. Up to 4 repeated cognitive assessments by telephone interview were completed over 12 years Quote: "The beta-carotene group of the PHS II continued as planned through May 2003 Treatment and follow-up of the vitamin E and C components continued through August 2007The multivitamin intervention continued through 1 June 2011" Eligible cognitive outcomes: • global cognitive function measured with TICS • episodic memory (verbal memory) measured using an average of z-scores* for immediate and delayed recall on the EBMT and delayed recall of a 10-word list • executive functioning measured with category fluency: animals named during 1 minute Function outcome extracted: none reported
	Quality of life outcome extracted: none reported Safety outcome extracted: none reported
Notes	Funding sources, Quote: "National Institutes of Health, BASF, Pfizer, and DSM Nutritional Products." "By grants CA 097193, CA 34944, CA 40360, HL 26490, HL 34595, AG 15933, and T32-AG000158 from the National Institutes of Health, and an investigator-initiated grant from BASF. Study agents and packaging were provided by BASF and Pfizer, and study packaging was provided by DSM Nutritional Products." Declarations of interest, quote: "Disclosures can be viewed at www.acponline.org/authors/icmje/ConflictOfInterestForms.do?msNum_M13-1340." Several authors reported some conflicts of interest Treatment adherence: "83.5% of the multivitamin group and 84.2% of the placebo group reported taking at least two thirds of their study pills"

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" "Randomizationusing a computer-generated list of random numbers" Comment: adequate method of random generation

Grodstein 2013 (Continued)

Allocation concealment (selection bias)	Low risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" Comment: method of allocation concealment not reported, but likely to be adequate
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" Comment: method of blinding not reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The PHS II is a randomized, double-blind, placebo-controlled, 2x2x2x2 factorial trial" "An Endpoint Committee of physicians blinded to the participants' treatment assignment, will review the medical records for final confirmation of a reported diagnosis" Comment: The method of blinding of outcome assessment for the cognitive substudy was not reported. The statement about the Endpoint Committee is contained in the design paper which, however, does not mention any cognitive outcome
Incomplete outcome data (attrition bias) All outcomes	Low risk	There was good retention of participants from the first to the second (96%) and the third (90%) assessments Quote: "the fourth assessment was not attempted in many participants because of trial completion (that is, 2700 (45%) of the initial 5947 who completed the initial interview were invited to participate at the fourth assessment before the trial closed on 1 June 2011)" Comment: 3623 (61%) patients had none of the assessments at last visit so we considered there was a high risk of attrition bias for the final assessment only (time point > 5 to 10 years)
Selective reporting (reporting bias)	Low risk	Comment: No differences identified between the protocol ClinicalTrials.gov: NCT00270647) and the article. All outcomes specified in the Methods were re-

Grodstein 2013 (Continued)

		ported in the Results	
Other bias	Low risk	No other risks identified	d
Hankey 2013			
Methods	Prespecified substudy of the placebo-controlled trial	Prespecified substudy of the VITATOPS trial, a randomized, 2-arm, parallel group, placebo-controlled trial	
Participants	Number randomised: 8164 p TATOPS. This substudy was a defined as MMSE ≥ 24 six who had 1 or more further M 1110/2214 allocated to B vita 1104/2214 allocated to placed Participant (baseline) charace gender: 67.3% male gend	Location: international, multicentre (20 countries from 4 continents) Number randomised: 8164 participants with recent stroke or TIA randomised in VI-TATOPS. This substudy was of the 2214 participants who were cognitively unimpaired - defined as MMSE ≥ 24 six months after the qualifying cerebrovascular event - and who had 1 or more further MMSEs during follow-up 1110/2214 allocated to B vitamins 1104/2214 allocated to placebo Participant (baseline) characteristics: • gender: 67.3% male • age, mean (SD): 63.6 (11.8) • baseline cognitive function: ○ B vitamins group MMSE 28.05 (SD 1.76) ○ placebo group MMSE 28.09 (SD 1.77) Inclusion criteria: "patients presenting within 7 months of stroke (ischaemic or haemorrhagic) or TIA (eye or brain), as defined by standard criteria, are eligible" (for VI-TATOPS). For cognitive substudy, MMSE ≥ 24 six months after the qualifying cerebrovascular event and ≥ 1 further MMSE measured during follow-up Exclusion criteria: patients taking folic acid or vitamin B6 on medical advice; taking methotrexate for any reason; pregnant or at risk of pregnancy; limited life expectancy	
Interventions	Intervention (n = 1110): Folic acid 2 mg, vitamin B6 2 Comparator (n = 1104): Placebo	Folic acid 2 mg, vitamin B6 25 mg, vitamin B12 500 μ g Comparator (n = 1104):	
Outcomes	ment (MMSE score < 24 on ≥ Secondary outcomes were tHo baseline, ≥ 6 months after th	Primary outcome for this prespecified substudy was a new diagnosis of cognitive impairment (MMSE score < 24 on ≥ 2 follow-up visits ≥ 6 months after the qualifying stroke Secondary outcomes were tHcy, the mean MMSE, cognitive decline (i.e. a decline from baseline, ≥ 6 months after the qualifying stroke, of ≥ 3 points in the MMSE score on ≥ 2 follow-up visits), composite of cognitive impairment and decline	
Notes	Council, the Biomedical Resea of Australia, the Royal Perth	Funded by National Health and Research Council of Australia, the UK Medical Research Council, the Biomedical Research Council of Singapore, the National Heart Foundation of Australia, the Royal Perth Hospital Medical Research Foundation, and the Health Department of Western Australia	
Risk of bias			
Bias	Authors' judgement	Support for judgemen	t

Hankey 2013 (Continued)

Random sequence generation (selection bias)	Low risk	Quote: "After baseline data have been provided by the investigator and it is confirmed that the patient meets the study eligibility criteria, a central 24-hour telephone service or an interactive website uses random permuted blocks stratified by hospital to allocate a treatment pack number."
Allocation concealment (selection bias)	Low risk	Quote: "After baseline data have been provided by the investigator and it is confirmed that the patient meets the study eligibility criteria, a central 24-hour telephone service or an interactive website uses random permuted blocks stratified by hospital to allocate a treatment pack number."
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "the tablets being either vitamin supplements or matching placebo"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	No specific information about site investigators. Likely blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "The 2214 cognitively unimpaired participants who underwent subsequent cognitive testing were more likely to be white, have a qualifying TIA and lacunar syndrome at baseline, and less likely to have experienced an early recurrent stroke within 1 year of randomization compared with the 394 cognitively unimpaired participants who did not proceed to further cognitive testing." "there was no significant difference in the prevalence of baseline characteristics among the 2214 cognitively unimpaired participants allocated B vitamins (n = 1110) compared with placebo (n = 1104), with the exception of atrial fibrillation (9. 1% B vitamins versus 6.5% placebo; P = 0. 022)."
Selective reporting (reporting bias)	Low risk	Prespecifed cognitive outcomes fully reported
Other bias	Low risk	No other risks identified

Kang 2006

Methods	Cognitive substudy of the Women's Health Study, a randomized, placebo-controlled, 2 x 2 factorial trial of vitamin E and low-dose aspirin with follow-up to approximately 10 years
Participants	Location: USA, number of sites not reported Recruitment: female health professionals Number randomised: 3184 in intervention, 3193 in comparator Participant (baseline) characteristics: • Age at randomisation: • vitamin E group: 66.2 (± 4)(range 60.4-89.9) • placebo: 66.3 (± 4.1)(range 60.4-87.1) • Age at initial cognitive assessment: • vitamin E group: 71.8 (± 4)(range 66.1-95.5) • placebo: 71.9 (± 4.1)(range 66.0-92.8) • · Gender: women 100% • · Baseline cognitive function: not assessed Inclusion criteria: Women 65 years or older participating in the Women's Health Study. Inclusion criteria for the main study: "Women were eligible if they were 45 years of age or older; had no history of coronary heart disease, cerebrovascular disease, cancer (except non-melanoma skin cancer), or other major chronic illness; had no history of side effects to any of the study medications; were not taking aspirin or nonsteroidal antiinflammatory medications (NSAIDs) more than once a week (or were willing to forego their use during the trial); were not taking anticoagulants or corticosteroids; and were not taking individual supplements of vitamin A, vitamin E, or beta carotene more than once a week." Exclusion criteria: not reported for the cognitive substudy
Interventions	Intervention (n = 3184): Vitamin E supplementation (600 IU α -tocopherol acetate; Natural Source Vitamin E Association, La Grange Ill) on alternate days Comparator group (n = 3193): Placebo Participants also randomised to aspirin 100 mg on alternate days in 2 x 2 factorial design Use of additional interventions (common to both treatment arms): not reported
Outcomes	The first cognitive assessment was a mean of 5.6 years (range 4.4 to 6.8 years) after randomisation and the final assessment was a mean of 4 years (range 2.6 to 5.7 years) later. We considered that all assessments fell into our duration category of > 5-10 years of treatment and, following the protocol, used the latest, i.e. the final, assessment only Primary Outcomes: global composite score averaging performance on general cognition, verbal memory and category fluency Tests used: • Telephone Interview of cognitive status (TICS) • East Boston Memory test immediate and delayed recall • Delayed recall of the TICS 10-word list Secondary Outcomes: composite score of verbal memory, averaging performance across 4 measures of verbal memory Cognitive assessments of general cognition, verbal memory, and category fluency were administered by telephone at 2-year intervals by trained nurses

Kang 2006 (Continued)

Time from randomisation to final cognitive assessment: approximately 10 years Eligible cognitive outcomes: • global cognitive function measured at approximately 10 years, with higher values

- indicating benefit
- executive functioning measured at approximately 10 years, with higher values indicating benefit
- episodic memory measured with TICS 10-word test delayed recall at approximately 10 years, with higher values indicating benefit

Function outcome extracted: none reported Quality of life outcome extracted: none reported Safety outcome extracted: none reported

Notes

Primary aim of study was to investigate effects of interventions on the primary prevention of cardiovascular disease and cancer. At the time of initiation of the cognitive substudy, participants had been treated for a mean of 5.6 years since randomisation

Funding sources, quote: "This work was supported by grants CA47988 and AG15933 from the National Institute of Health", "The funding agency did not play any role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; and preparation, review, or approval of the manuscript"

Declarations of interest, For the sub-study: none reported. For the main study, quote: "Dr. Ridker reports having received grant support from Bayer. Dr. Cook reports having served as a consultant to Bayer. Dr. Gaziano reports having served as a consultant to, and receiving grant support from, Bayer and McNeil. Dr. Hennekens reports having served as a consultant to Bayer and McNeil and receiving grant support from Bayer"

Treatment adherence

Quote: "compliance was comparable between the 2 groups: as of the final cognitive assessment, the percentage that reported taking at least two thirds of the assigned pill was 75.4% for the vitamin E group and 76.9% for the placebo group"

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Comment: method of random sequence generation not reported. Probably adequate
Allocation concealment (selection bias)	Unclear risk	Comment: method of allocation concealment not reported. Probably adequate
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "The WHS is a double-blind, RCT."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Comment: for the substudy on cognitive function, blinding of outcome assessment was not reported

Kang 2006 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: 18% of patients randomised and included in the substudy were not included in the analysis at last assessment
Selective reporting (reporting bias)	Unclear risk	Comment: All outcomes in the Methods were reported in the results. Protocol not available
Other bias	Low risk	No other risks identified

Other bias	20W Tisk
Kang 2008	
Methods	WAFACS study. A 2 x 2 x 2 x 2 factorial, randomised, placebo-controlled trial of 3
Wictious	antioxidants (vitamin E, vitamin C, and \(\beta\)-carotene) and B vitamins with 6.6 years o
	treatment and 5.4 years of follow-up of cognitive function
	The WAFACS study was started in 1998 and continued the WACS study (Kang 2009
	with the introduction of a fourth arm treated with folic acid, vitamin B6, and vitamin
	B12
	Surviving participants of the WACS were invited to participate in the WAFACS
	When the folic acid, vitamin B6, and B12 arm was added, this combination or its placeb
	was combined in a retained factorial design with the vitamin C tablet or placebo
Participants	Location, Country: United States of America, number of sites not reported. Inclusion
1	from December 1998 to July 2000
	Recruitment: female health professionals
	Number randomised: 1071 in intervention, 1093 in comparison, but 1002 and 100
	considered at baseline because had available baseline cognitive assessment
	Participant (baseline) characteristics:
	 Mean age ± SD (range) at randomisation:
	o vitamin: 71.3 ± 4.2 (65.3-89.9)
	o placebo:71.3 ± 4.2 (65.3-90.0)
	 Mean age ± SD (range) at initial cognitive assessment: vitamin: 72.5 ± 4.2 (66.1-90.9)
	o placebo: 72.5 ± 4.2 (66.1-90.2)
	• · Gender: women 100%
	 • Main diagnosis: female health professionals aged ≥ 40y with cardiovascular
	disease or ≥ 3 coronary risk factors
	Inclusion criteria:
	This was a substudy of cognitive function among 2009 participants aged \geq 65 years from
	the Women's Antioxidant and Folic Acid Cardiovascular Study, a randomised placebo
	controlled trial designed to test the effect of a combination of B vitamins on secondar
	prevention of cardiovascular disease in female health professionals aged ≥ 40 years wit
	cardiovascular disease or ≥ 3 coronary risk factors
	The substudy was initiated a mean of 1.2 years after randomisation

Exclusion criteria: not reported for the substudy

Exclusion criteria in the main study, quote: "Women were excluded if they had a history of cancer (excluding non-melanoma skin cancer) within the past ten years, any serious non-CVD illness, or were currently using warfarin or other anticoagulants. To be eligible for

Kang 2008 (Continued)

	the folic acid/vitamin B6/B12 component, potential participants in the ongoing eight- arm trial had to be additionally willing to forgo individual supplements of folic acid, vitamin B6, and vitamin B12 at levels beyond the US recommended daily allowance"
Interventions	Intervention (n = 1002): 2.5 mg folic acid/day, 50 mg vitamin B6/day, and 1 mg vitamin B12/day Quote: "5442 women were assigned to 16 treatment assignments that were combinations of the 3 antioxidants" and 1 B vitamin supplement Quote: "Every 12 mo, the women were sent a year's supply of monthly calendar packs containing active agents or placebo." Comparator group (n = 1007): placebo. Use of additional interventions (common to both treatment arms): not reported.
Outcomes	Outcomes of interest in the review: Primary Outcomes: global composite score averaging all 5 test results measuring general cognition, verbal memory, and category fluency using z scores Quote: "We assessed cognitive function using a telephone cognitive battery with 5 tests measuring general cognition, verbal memory, and category fluency." The following tests were used: • For general cognition, Telephone Interview of Cognitive Status (TICS); • For verbal memory, delayed recall of the TICS 10-word list and the immediate and delayed recalls of the East Boston Memory Test; • For category fluency, women were asked to name as many animals as possible in 1 min. Secondary Outcomes: verbal memory composite score, calculated by averaging 4 measures of verbal memory (the immediate and delayed recalls of both the East Boston Memory Test and the 10-word list) Telephone cognitive function testing was administered up to 4 times over 5.4 years The average time from randomisation to the initial cognitive assessment was 1.2 years (range: 0.7-1.7 years), and from randomisation to the last assessment was 6.6 years (range: 6.3-6.9 years). Eligible cognitive outcomes: • global cognitive function measured with TICS at the initial cognitive assessment (1.2 years); the second cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (6.6 years follow-up), with higher values indicating benefit* • episodic memory: composite(delayed East Boston memory, immediate East Boston memory, delayed TICS-10) at the initial cognitive assessment (1.2 years); the second cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (around 3 years of follow-up) and the fourth cognitive assessment (6.6 years follow-up), with highe

Kang 2008 (Continued)

Notes	E. J
Notes	Funding sources, quote: "Folic acid, vitamin B-6, vitamin B-12, and placebos were
	provided by the BASF Corporation (Mount Olive, NJ)"
	"Supported by grants AG15933 and HL47959 from the National Institutes of Health"
	Declarations of interest, quote: "None of the authors declared a conflict of interest."
	"Other Financial Disclosures: Dr Gaziano has received investigator initiated study sup-
	port in the form of vitamin pills and packaging from Wyeth. Dr Buring has received
	investigator initiated study support in the form of vitamin pills and packaging from
	Natural Source Vitamin E Association. Dr. Manson has received investigator-initiated
	study support in the form of vitamin pills and packaging from Cognis and BASF."
	Treatment adherence
	Quote: "Compliance was comparable (83%) in the 2 treatment groups". This statement,
	however, seems to have referred to the main study and not to this substudy on cognitive
	function,

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: no information Comment: method of random sequence generation not reported
Allocation concealment (selection bias)	Unclear risk	Quote: no information Comment: method of allocation conceal- ment not reported
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: Protocol: "Participants and investigators will be blinded to treatment assignment." Comment: method of blinding not reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The telephone cognitive interviews were administered by trained interviewers" From the main study: "An endpoints committee of physicians who were blinded to randomized treatment assignment adjudicated all primary and secondary cardiovascular outcome events. Study medications and end point ascertainment were continued in a blinded fashion until the scheduled end of the trial, July 31, 2005." Comment: It was not specifically reported whether cognitive assessors were blinded, but we considered this to be highly likely

Kang 2008 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Participants received 3 follow-up cognitive assessments approximately every 2 y 94% completed at least one follow-up assessment, and 83% completed ≥ 3 of 4 assessments. In the fourth assessment, 24% of participants were not contacted for their assessment because only a short intervial had passed between their third interview and the end of the trial in July 2005." Comment: Although the authors provided some explanation for not seeing some patients at last visit (visit 4), the number of patients not undergoing cognitive evaluation was considerable: 956/2009 (47.5%) at visit 4 and 314/2009 (15.6%) at visit 3. RIsk of bias related to incomplete outcome data considered high for visit 4
Selective reporting (reporting bias)	Low risk	Comment: All outcomes reported in the Methods were presented in the Results. In addition, the outcomes reported in the protocol were reported as well in the article
Other bias	Low risk	Quote: "155 women (7%) were unreachable, had declined participation, or had died; thus, 2009 (93%) women completed the initial telephone cognitive assessment." Comment: inclusion was probably not consecutive, however, reasons for not including in the substudy were given and the number excluded appeared balanced between the 2 groups

Kang 2009

Methods	$2 \times 2 \times 2$ factorial, randomised, placebo-controlled trial of 3 antioxidants (vitamin E, vitamin C, and β -carotene) with 8.9 years (range 7.8-9.6) of follow-up
Participants	Location: Country: USA and Puerto Rico, no of sites etc. not reported. From December 1998 to July 2000 Quote: "From November 1994 to October 1996 an introductory letter, informed consent form, and enrolment questionnaire were mailed to female health professionals who were originally identified through state licensing boards and professional organizations in the continental United States, Alaska and the Commonwealth of Puerto Rico" "Potential candidates were also drawn from a cohort of women with prevalent CVD participating in the long-running Nurses' Health Study" Recruitment: female health professionals

Number randomised:

2824 in cognitive substudy

- overall, vit E: 1428;
- overall, vit C: 1406;
- overall, betaCarot: 1406;
- placebo vit E, C, betaCarot: 353;
- placebo vit E: 1396;
- placebo vit C: 1418;
- placebo betaCarot: 1418.

Participant (baseline) characteristics:

Mean age (± SD, range) at randomisation:

- overall, vit E: 69.1 (4.3; 62.6-87.4)
- placebo vit E: 69.0 (4.2; 62.6-87.9)
- overall, vit C: 69.0 (4.2; 62.6-86.9)
- placebo vit C: 69.1 (4.3; 62.6-87.9)
- overall, betaCarot: 69.1 (4.3; 62.6-87.9)
- placebo betaCarot: 69.0 (4.2; 62.6-87.6)

Mean age (± SD, range) at initial cognitive assessment:

- overall, vit E: 72.6 (4.3; 66.1-90.9)
- placebo vit E: 72.5 (4.2; 66.1-91.3)
- overall, vit C: 72.5 (4.2; 66.1-90.5)
- placebo vit C: 72.6 (4.3; 66.1-91.3)
- overall, betaCarot: 72.6 (4.3; 66.2-91.3)
- placebo betaCarot: 72.5 (4.2; 66.1-91.2)

Gender: women 100%

Main diagnosis: female health professionals aged \geq 40y with cardiovascular disease or \geq 3 coronary risk factors

Inclusion criteria: This was a substudy of cognitive function among 2824 participants aged ≥ 65 years from the Women's Antioxidant Cardiovascular Study, a randomised placebo-controlled trial designed to test the effect of a antioxidants on secondary prevention of cardiovascular disease in female health professionals aged ≥ 40 years with cardiovascular disease or ≥ 3 coronary risk factors

Exclusion criteria: not reported for the substudy

Exclusion criteria in the main study, quote: "Women were excluded if they had a history of cancer (excluding non-melanoma skin cancer) within the past ten years, any serious non-CVD illness, or were currently using warfarin or other anticoagulants. To be eligible for the folic acid/vitamin B6/B12 component, potential participants in the ongoing eightarm trial had to be additionally willing to forgo individual supplements of folic acid, vitamin B6, and vitamin B12 at levels beyond the US recommended daily allowance" Only women who were compliant (taking at least two-thirds of the pill) during the 12-week pre-randomisation run-in period were randomised

Interventions

Intervention (overall, vitamin E: 1428; overall, vitamin C: 1406; overall, betaCarot: 1406)

Vitamin E at a dose of 402~mg (600~IU) every other day, ß-carotene (50~mg every other day) and vitamin C (500~mg daily)

Every 12 months, the women were sent a year's supply of monthly calendar packs containing active agents or placebo

Comparator group: placebo (placebo vitamin E, vitamin C, ß-carotene: 353; placebo

Kang 2009 (Continued)

	vit E: 1396; placebo vitamin C: 1418; plac Use of additional interventions (common	
Outcomes	Primary Outcomes: global composite score averaging all 5 test results measuring general cognition, verbal memory, and category fluency using z scores Quote: "We assessed cognitive function by telephone and administered 5 tests measuring general cognition, verbal memory and category fluency." The following tests were used: • Telephone Interview of Cognitive Status (TICS) • delayed recall of the TICS 10-word list and the immediate and delayed recalls of the East Boston Memory Test (data not extractable per instrument) • category (animal) fluency Secondary Outcomes: 'verbal memory' composite score, calculated by averaging 4 measures of the immediate and delayed recalls of both the East Boston Memory Test and the TICS 10-word list Telephone cognitive function testing was administered up to 4 times over 5.4 years The substudy was initiated a mean of 3.5 years after randomisation. Quote: "The average time from randomization to the initial cognitive assessment was 3.5 years (range 3.1-4.7), and from randomization to the last assessment was 8.9 years (range 7.8-9.6)" Eligible cognitive outcomes: • global cognitive function measured with TICS • executive function measured with category fluency	
Notes	Also known as the WACS study. Funding sources, quote: "Vitamin E and its placebo were provided by Cognis Corporation (LaGrange, IL); all other agents and their placebos were provided by BASF Corporation (Mount Olive, NJ)." "This work is supported by grants AG15933, HL046959 from the National Institutes of Health." Declarations of interest, quote: "none." "Other Financial Disclosures: Dr Gaziano has received investigator initiated study support in the form of vitamin pills and packaging from Wyeth. Dr Buring has received investigator initiated study support in the form of vitamin pills and packaging from Natural Source Vitamin E Association. Dr. Manson has received investigator-initiated study support in the form of vitamin pills and packaging from Cognis and BASF." Treatment adherence "average compliance (defined as taking at least two-thirds of assigned study medications) during follow-up was 83% and did not differ significantly between the two groups". This statement, however, seemed to have referred to the main study and not to this substudy on cognitive function "At the end of the study, compliance (defined as taking at least two-thirds of study pills) was comparable across all groups (range 64-68%)."	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomized in a $2 \times 2 \times 2$ factorial design." Comment: method of random sequence

Kang 2009 (Continued)

		generation not reported. Probably adequate
Allocation concealment (selection bias)	Unclear risk	Comment: method of allocation concealment not reported
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "matching placebo" Comment: same study as Kang 2008 (reported vitamin B arm) in which more detailed description of the methods stated that "Participants and investigators will be blinded to treatment assignment."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The telephone cognitive interviews were administered by trained interviewers, who were masked to the participants' randomized treatment assignment." From the main study: "An endpoints committee of physicians who were blinded to randomized treatment assignment adjudicated all primary and secondary cardiovascular outcome events. Study medications and end point ascertainment were continued in a blinded fashion until the scheduled end of the trial, July 31, 2005." Comment: Physicians were blinded to treatment assignment in the main study. The nurses were the assessors in this cognitive substudy and were apparently not aware of treatment assignment
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	In this review, we used outcome data from the first cognitive assessment (after a mean of 3.5 years) - this included 2824 of 3170 eligible participants (89%) - and the fourth cognitive assessment (after a mean of 8. 9 years) - this included 1586 participants (50%). No information was given about the treatment assignments of missing participants. We considered there to be an unclear risk of attrition bias for the first assessment and a high risk for the final assessment
Selective reporting (reporting bias)	Low risk	Comment: All outcomes reported in the Methods were presented in the Results. In addition, the outcomes specified in the protocol were reported in the article

Other bias	Low risk	No other risks identified	
Kesse-Guyot 2011			
Methods	Substudy of the SU.VI.M./	Substudy of the SU.VI.MAX, a randomised, double-blind, placebo-controlled, primary prevention trial	
Participants	Recruitment: selected Fre SU.VI.MAX study (1994-for a planned follow-up of subjects who agreed to pain the SU.VI.MAX 2 stud quality of aging." Sample size:		
	had agreed to participate cognitive evaluation, but t aged 45-60 years at the on	(see Inclusion criteria below*)	
	 Age at baseline: place ± 4.5 Age at cognitive evalue 2376): 65.5 ± 4.5 Male sex: placebo: 52 *Inclusion criteria: Quote current analysis we selected 	too (n = 2071): 52.1 ± 4.6 , supplementation (n = 2376): 52.1 mation: placebo (n = 2071): 65.5 ± 4.6 , supplementation (n = .2% (n = 1080), supplementation: 51.7% (n = 1229): "Of the 6850 participants in the SU.VI.MAX 2 study, for the participants who were $45-60$ y old at baseline, had undergone had available baseline data for serum antioxidants, smoking aption."	
Interventions	mg), selenium (100 μg), at Comparison (n = 2071): p N.B. Intervention period v	Intervention (n = 2376): daily vitamin C (120 mg), beta-carotene (6 mg), vitamin E (30 mg), selenium (100 µg), and zinc (20 mg) in combination Comparison (n = 2071): placebo N.B. Intervention period was 8 years (1994-2002). Cognitive outcomes in SU.VI.MAX 2 were measured in 2007-9, a mean of 5.9 (± 0.4) years after the end of the intervention in both groups	
Outcomes	 Episodic memory: RI Executive function: c 	Outcomes of interest in the review: 1. Episodic memory: RI-48 test 2. Executive function: category fluency 3. Speed of processing: time taken to complete the Delis-Kaplan trail-making test (TMT)	
Notes	PNRA- 010), Direction C	Funding source: "Supported by the Agence Nationale de Recherche (no. ANR-05-PNRA- 010), Direction Générale de la Santé (Ministry of Health), Mederic, Sodexo, Ipsen, the Mutuelle Générale de l'Education Nationale (MGEN), and Pierre Fabre.	

Kesse-Guyot 2011 (Continued)

Mederic and MGEN are French health insurance organizations that are complementary to the national health insurance system. Ipsen and Pierre Fabre are private pharmaceutical companies. They provided financial support for the overall implementation of the research project. Sodexo, a catering company, provided financial support for several teamwork events with researchers and study participants. Because French public research is independent, the sponsors were not involved in the analysis or interpretation of the findings."

Declarations of interest: "None of the authors had a conflict of interest".

Treatment adherence: "At the end of the trial phase (2002), 84% of the participants reported having taken at least three-fourths of the capsules independent of the allocated supplementation group"

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Random treatment allocation was performed by block-sequence generation stratified by sex and age group." Comment: Should be adequate, since team included statisticians and computing staff who handled the data
Allocation concealment (selection bias)	Low risk	Quote: "Randomization was concealed from subjects and all investigators except for the few who were in charge of capsule labelling." Comment: Should be adequate.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Capsules were prepared in 52 weekly packages of 7 capsules and delivered each year in a box labelled with the participant's number and a 10-digit lot number" "a matching placebo". "The absence of an easy way to distinguish antioxidant from placebo capsules was tested in a pilot study", "double-blind" Comment: Blinding of participants and personnel seemed adequate
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Once a possible event is suspected, all relevant records,are collected from the hospitals, laboratories, or institutions or from the participants by the SU.VI.MAX medical investigators and are examined at the study coordinating centre" Comment: It was not reported if outcome assessment was blinded for adverse events. However, the blinding should be adequate

Kesse-Guyot 2011 (Continued)

		for cognitive function since the patients and investigators were well blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Comment: 1136 of 5583 (20%) of the eligible patients were not included in the analysis because of missing cognitive data
Selective reporting (reporting bias)	Unclear risk	Comment: Cognitive outcomes were not mentioned in the protocol of the main study
Other bias	High risk	Participants in this study were those who responded to an invitation 6 years after the end of the parent study. The authors described this subsample of the original trial population as "more compliant and health conscious, as shown by the comparison between included and excluded subjects." This may work to reduce any difference between groups

Kryscio 2017

Methods	PREADVISE (Prevention of Alzheimer's Disease by Vitamin E and Selenium Trial) Ancillary study to SELECT (primary aim prevention of prostate cancer) Began as a double-blind RCT in 2002, transformed into a cohort study from 2009-15 Randomised, double-blind, 2 x 2 factorial design
Participants	Location: 130 SELECT clinical sites in the US, Canada and Puerto Rico Recruitment: 7540 non-dementia men already enrolled in SELECT were recruited into PREADVISE between 2002 and 2008. 7338 had at least one follow-up visit. 4271 con- sented to continue annual telephone follow-up after the RCT was closed and supple- ments stopped in 2009/10. Of these, 3786 were screened by telephone at least once Number randomised: 7540 Participant characteristics: Age: Mean (SD) 67.5 (5.3) at PREADVISE baseline Gender: all participants were male Inclusion criteria: SELECT participants aged 62 years and over (age 60 years if black) Exclusion criteria: dementia, active neurologic and/or neuropsychiatric conditions that affect cognition, as well as history of serious head injury (> 30-minute loss of conscious- ness within the last five years prior to enrolment) and substance abuse
Interventions	Supplements were taken for 5.4 ± 1.2 years 2 x 2 factorial design: vitamin E 400 IU/day, selenium 200 mcg/day, placebo
Outcomes	Dementia incidence was the primary outcome of PREADVISE. It was determined by one of two methods. First tier screening was the Memory Impairment Screen (MIS). A score $\leq 5/8$ triggered a second tier screen which was the expanded

Kryscio 2017 (Continued)

Consortium to Establish a Registry in Alzheimer's Disease (CERADe) battery during the RCT and the modified Telephone Interview for Cognitive Status (TICSm) during the cohort study

Method 1: Participants with T score \leq 35 on CERADe or total score \leq 35 on TICSm were encouraged to obtain a memory work-up and share medical records with PREAD-VISE. Medical records were reviewed by a team of 2-3 expert neurologists and 2-3 expert neuropsychologists to determine a consensus diagnosis

Method 2: Participants who did not obtain a memory work-up were assessed by additional measures collected during the study: AD8 Dementia Screening Interview, self-reported medical history, self-reported medication use, and cognitive test scores including the MIS, CERAD T Score, NYU Paragraph Delayed Recall, and TICSm. An AD8 \geq 1 (at any time during follow-up) plus a self-reported dementia diagnosis, use of a memory enhancing prescription drug (i.e. donepezil, rivastigmine, galantamine, memantine), or cognitive score \geq 1.5 standard deviations below expected performance yielded a dementia diagnosis. The diagnosis date was assigned to the earliest event

Notes

"PREADVISE was ancillary to SELECT, a randomized controlled trial of the same antioxidant supplements for preventing prostate cancer. SELECT closed in 2009 due to a futility analysis." "The RCT was scheduled to continue supplements until 2012 but in September 2008, the SELECT Data and Safety Monitoring Committee recommended that supplements be discontinued due to lack of efficacy on prostate cancer incidence. Study sites closed over the next two years, during which time both PREADVISE and SELECT transitioned into cohort studies. RCT participants were asked to continue in the cohort study.."

PREADVISE had intended to recruit 10,400 men to achieve 80% power to detect a hazard ratio of 0.6

Power was reduced by under-recruitment, loss of sites and participants when RCT closed, and difficulties with case ascertainment

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomized". No details. Large multisite cancer prevention trial. Likely adequate
Allocation concealment (selection bias)	Low risk	No details. Large multisite cancer prevention trial. Likely adequate
Blinding of participants and personnel (performance bias) All outcomes	Low risk	"Double-blind". No details. Likely adequate
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Double-blind". No details. Likely adequate
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The transition from RCT to cohort study was associated with loss of approximately

Kryscio 2017 (Continued)

		half the participants to follow-up. "Fortunately the major cost was sample size (reduction in person-years of follow-up) since no new large selection biases were introduced." "All PREADVISE participants who completed at least one follow-up visit were included in the current analyses (intent-to-treat analysis, ITT), whether they participated in just the RCT or both the RCT and cohort studies."
Selective reporting (reporting bias)	Low risk	
Other bias	Low risk	" despite randomization occurring at SE-LECT baseline rather than PREADVISE enrolment, there were no perceivable differences between study arms in terms of medical history, APOE ϵ 4 genotype, or initial MIS."

Lewerin 2005

Lewerin 2005	
Methods	2-arm, double-blinded, parallel group RCT, 4 months treatment and follow-up
Participants	Location: one centre, Goteborg, Sweden Recruitment: outpatient clinic Number randomised: 126 in intervention, 69 in comparison. Participant (baseline) characteristics in Vitamin (n = 126) and placebo (n = 69) Age: • vitamin 75.7 (± 4.7); • placebo 75.6 (± 4.0) • whole group (n = 209) mean age: 76 years and 5 months. Median 76 years (range 70-93) Gender (males %): • vitamin: 38% (48/126) • placebo: 44% (30/69) Inclusion criteria: community-dwelling subjects. Inclusion criteria not reported Exclusion criteria: quote: "Those who had taken any vitamin supplements during the last 3 months or pharmacological doses of vitamin B12, folic acid and/or vitamin B6 during the last 3 years were not allowed to enter the study"
Interventions	Intervention (n = 126): Daily tablet containing 500 μg cyanocobalamin, 800 μg folic acid, and 3 mg vitamin B6 hydrochloride (manufactured and supplied by Recip AB, Årsta, Sweden) Comparator group (n = 69): daily oral placebo The duration of treatment was 4 months. Use of additional interventions (common to both treatment arms): Use of cardiovascular medication:

 placebo: 49% Use of antiepileptics, neuroleptics, or antidepressants (%) vitamin: 22% placebo: 23%

Outcomes

Outcomes evaluated: postural-Locomotor-Manual test and cognitive tests Postural-Locomotor-Manual test:

- Movement time
- Postural phase
- Locomotor phase
- Manual phase
- Simultaneity index

Cognitive tests:

- Digit span forward
- Digit span backward
- Identical forms
- Visual reproduction
- Synonyms
- Block design
- Digit symbol
- Thurstone's picture memory test
- Figure classification

No clear distinction between primary and secondary outcomes

In Lewerin 2003, cognitive and functional tests not reported in the Methods Cognitive testing was conducted by the same psychologist at baseline and after 4 months Follow-up: 4 months (measurements before and after treatment). Mean or median follow-up not reported

Eligible cognitive outcomes:

- executive functioning measured with synonyms at 4 months, on a scale from not reported to 30 with higher values indicating benefit*
- episodic memory measured with Thurstone's picture memory test at 4 months, on a scale from not reported to 28 with higher values indicating benefit
- episodic memory measured with identical forms (Dureman 1959) at 4 months, on a scale from not reported to 60 with higher values indicating benefit
- episodic memory measured with Wechsler visual reproduction at 4 months, on a scale from not reported to 14 with higher values indicating benefit*
- working memory measured with Wechsler digit span forward at 4 months, on a scale from not reported to 9 with higher values indicating benefit
- working memory measured with Wechsler digit span backward at 4 months, on a scale from not reported to 8 with higher values indicating benefit*
- speed of processing measured with Wechsler digit symbol at 4 months, on a scale from 0 to 90 with higher values indicating benefit*
- speed of processing measured with Thurstone's figure classification at 4 months, on a scale from not reported to 30 with higher values indicating benefit

Function outcome extracted: none reported

Quality of life outcome extracted: none reported

Safety outcome extracted: none reported

* outcome data used in statistical analyses

Notes	Funding sources, quote: "Supported by grants from the Hjalmar Svensson Foundation,		
	the Göteborg Medical Society, the Medical Faculty at Göteborg University, the Wilhelm		
	and Martina Lundgren Foundation, and the Magnus Strandqvist Foundation. Recip AB supported the study and provided the vitamin and placebo tablets" Declarations of interest, quote: "None of the authors had a personal or financial conflict of interest with respect to this study" Treatment adherence: quote "To ensure compliance, all subjects received a specified blinded number of tablets, and at the end of the study, the number of remaining tablets		
	was compared with the initial number and planned intake during the study"		
	Data on treatment adherence were not clearly reported in the results		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Of these 195 subjects, 126 were randomly assigned to receive vitamin therapy and 69 to receive placebo." Comment: method of random sequence generation not reported
Allocation concealment (selection bias)	Unclear risk	Quote: "Of these 195 subjects, 126 were randomly assigned to receive vitamin therapy and 69 to receive placebo." Comment: method of allocation concealment not reported
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "and all subjects in the placebo group received an identical (other than the vitamin content) placebo tablet." To ensure compliance, all subjects received a specified blinded number of tablets" "The tablet was identical in shape and composition to the placebo tablet apart from the vitamin content"; "double-blind" study Comment: Both participants and physicians were apparently blinded
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "Cognitive testing was conducted by the same psychologist (GS) at baseline and after 4 mo." Comment: Blinding of outcome assessors was not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote: "From flow diagram, 30/209 (14%) did not provide cognitive outcome data." Comment: no information about group al-

Lewerin 2005 (Continued)

		location of excluded participants
Selective reporting (reporting bias)	Unclear risk	Comment: protocol not available
Other bias	Low risk	Comment: no other risks identified

Methods	3-arm, parallel group, randomised, placebo-controlled trial with 6 months duration of treatment and follow-up
Participants	Location: Two centres (Coleraine, UK and Clermont-Ferrand, France) recruited younge participants (55-70 years) and another two centres (Rome, Italy and Grenoble, France recruited older participants (70-87 years)
	Recruitment: Participants were recruited through posters, leaflets, local television and radio, and community groups and organizations serving older individuals living independently
	Sample size:
	Number randomised: 201 younger participants (55-70 years), 232 older participant (70-87 years)
	Number completed study with compliance > 80% and included in analysis: 18
	younger, 199 older Participant baseline characteristics (baseline characteristics provided only for stud
	completers):
	Age in years, mean (SD):
	• 15 mg/d: younger 61.4 (4.55), older 74.3 (3.91)
	• 30 mg/d: younger 61.7 (4.57), older 74.6 (3.89)
	• placebo: younger 62.3 (4.09), older 74.2 (3.58)
	Gender, males/females:
	• 15 mg/d: younger 30/30, older 34 /32
	• 30 mg/d: younger 31/34, older 34/32
	• placebo: younger 32/31, older 35 /32
	Inclusion criteria:
	1. men and women aged between 55 and 87 years;
	2. BMI between 20 and 30 kg/m2
	3. good health;
	4. MMSE score greater than 23;
	5. geriatric depression scale score less than 6.
	Exclusion criteria:
	"Participants were excluded from further involvement in the study according to the following criteria: (I) tobacco consumption of more than 10 g/day; (2) alcohol consumption of more than 10 g/day; (2) alcohol consumption of more than 10 g/day; (3) alcohol consumption of more than 10 g/day; (4) alcohol consumption of more than 10 g/day; (5) alcohol consumption of more than 10 g/day; (6) alcohol consumption of more than 10 g/day; (7) alcohol consumption of more than 10 g/day; (8) alcohol consumption of more than 10 g/day; (9) alcohol consumption of more than 10 g/day; (10) alcohol consumption of more
	tion of more than 30 (men) or 20 (women) g/day; {3) unconventional dietary habi
	(for example, vegetarians, vegans); (4) use of a mineral supplement during the precedir
	3 months; (5) use of more than three (55-70 years) or four (70-87 years) prescription
	drugs per day; (6) use of antidepressants, laxatives, or hormone replacement therap
	(7) I I I I I I I I I I I I I I I I I I I

(7) pathological diseases, including cancer and diabetes. For participants satisfying the inclusion criteria, a biochemistry profile was performed, which included a full blood profile and tests of kidney and liver function. On the basis of these data, participants

Maylor 2006 (Continued)

	were excluded if there was insufficient renal and hepatic performance, malabsorption or inflammatory chronic pathologies, and were included if there was negative serology for the HIV and hepatitis C viruses." Quote: "49 % of the initial 842 volunteers being excluded from the study at screening on the basis of cognitive impairment, depression, pathological conditions, medications, and so on"
Interventions	Intervention
incer rentions	Intervention A: Zn supplementation as zinc gluconate 15 mg/day, two tablets at the same time each day (usually after breakfast) for 6 months Intervention B: Zn supplementation as zinc gluconate 30 mg/day, two tablets at the same time each day (usually after breakfast) for 6 months
	Comparator group: placebo/zinc gluconate 0 mg/day, two tablets at the same time each day (usually after breakfast) for 6 months
	Use of additional interventions (common to both treatment arms): not reported
Outcomes	Primary Outcomes: Cognitive measures (Cambridge Automated Neuropsychological Test Battery, CANTAB) Visual memory was tested by pattern recognition memory; working memory was tested by spatial span and spatial working memory (SWM); and attention was tested by reaction time and matching to sample visual search (MTS) Quote: "Cognitive Function was assessed in the laboratory at baseline (before supplementation) and after 3 and 6 months of supplementation, using different (parallel) versions of the CANTAB tests on each occasion." Secondary Outcomes: serum and urinary zinc Quote: "Participants fasted overnight for 12h before blood and urine samples were taken on each of these occasions to determine Zn levels."
Notes	Funding sources, quote: "European Commission 'Quality of Life and Management of Living Resources' Fifth Framework Programme. contract no. QLKI-CT-2001-00168." Declarations of interest: not reported Treatment adherence "Compliance was less than 80% for three older participants and so they were also excluded." Outcome data were not reported separately for the Zinc and placebo groups
Risk of bias	
Nisk of ours	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Participants were assigned to one of three levels of Zn supplementation (placebo or 0 mg/d, 15 mg/d and 30 mg/d) according to the same standardised random order in each centre." Comment: not described how sequence generation was done. The possibility that this was quasi-randomised could not be ruled out

Maylor 2006 (Continued)

Allocation concealment (selection bias)	Unclear risk	As above
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: "Zn supplementation (0, 15 or 30 mg/d) was administered as zinc gluconate, participants taking two tablets at the same time each day (usually after breakfast) for 6 months. (Tablets were identified by a code so that neither the experimenter nor the participants knew the dose. The code was not broken until the study had been completed and all the data had been entered into computer files ready for analysis)" Comment: Although a placebo was mentioned, there was no description of its appearance or whether it could be distinguished from the zinc supplements
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Comment: as above. Unclear whether blinding was adequate
Incomplete outcome data (attrition bias) All outcomes	High risk	Comment: There was no description of the completeness of outcome data for each main outcome, including attrition and ex- clusions from the analysis
Selective reporting (reporting bias)	High risk	Comment: CANTAB data for the different interventional groups was shown graphically only. Number of participants for each outcome was not reported
Other bias	Unclear risk	Baseline characteristics were shown only for participants who completed the trial

McMahon 2006

Methods	2-arms, placebo-controlled, randomised clinical trial with intervention of 2 years and follow-up duration of 2 years. No mean or median (range) follow-up reported
Participants	Location: Dunedin, New Zealand, number of centres involved unclear Recruitment: Department of Human Nutrition, University of Otago Number randomised: 138 in intervention, 138 in comparison Participant (baseline) characteristics: Age, years, mean (SD) • placebo (n=126): 73.4 ± 5.7 • vitamin (n=127): 73.6 ± 5.8 Female sex, n (%):

- placebo (n=126): 65 (52)
- vitamin (n=127): 47 (37)

Inclusion criteria: age of 65 years or older. Those with a fasting homocysteine concentration of at least 13 μ mol/L and a normal plasma creatinine concentration (133 μ mol/L (1.5 mg/dL) in men and 115 μ mol/L (1.3 mg/dL) in women) were invited to participate in the trial

Exclusion criteria:

- ineligible if they had suspected dementia;
- were taking medications known to interfere with folate metabolism (e.g. oral hypoglycaemic agents or antiepileptic agents);
- were taking vitamin supplements containing folic acid, vitamin B12, or vitamin B6;
 - were being treated for depression;
 - had diabetes or had a history of stroke or transient ischaemic attacks.

Quote: "Five participants (three in the placebo group and two in the vitamin group) had a baseline MMSE score of less than 26."

Interventions

Intervention (n=138):

Treatment capsules contained the filler plus 1000 μ g of folate (the calcium salt of l-5-methyltetrahydrofolate), 500 μ g of vitamin B12 (cobalamin), and 10 mg of vitamin B6 (pyridoxine) (Merck Eprova)

Comparator group (n=133): placebo.

The placebo capsules contained a blend of magnesium stearate and microcrystalline cellulose as a filler

All capsules were gelatin-coated, identical in colour and shape, and packaged in blister packs

Outcomes

Primary outcomes: scores on tests of cognition including Mini-Mental State Examination, Wechsler Paragraph Recall test, Category Word Fluency test, Rey Auditory Verbal Learning Test, trials I-V, Rey Auditory Verbal Learning Test, trial VII, Raven's Progressive Matrices, Controlled Oral Word Association Test, Part B of the Reitan Trail Making Test, the National Adult Reading Test (at baseline)

Secondary Outcomes: biochemical measurements, including plasma concentrations of homocysteine, folate, and vitamin B12

Eligible cognitive outcomes:

- global cognitive function measured with MMSE at 1 and 2 years, with higher values indicating benefit
- executive functioning measured with Raven's Progressive Matrices at 1 and 2 years, with higher values indicating benefit
- executive functioning measured with Part B of the Reitan Trail Making at 1 and 2 years, with lower values indicating benefit
- executive function measured with Category Word Fluency test (total no. of words generated in three 1-min tests) at 1 and 2 years, with higher values indicating benefit
- episodic memory measured with Rey Auditory Verbal Learning, Trials I-V at 1 and 2 years, with higher values indicating benefit*
- episodic memory measured with Wechsler Paragraph Recall test at 1 and 2 years, with higher values indicating benefit
- episodic memory measured with Rey Auditory Verbal Learning, Trial VII at 1 and 2 years, with higher values indicating benefit

Notes	Funding sources: supported by an Otago Research Grant, Merck Eprova for providing the vitamin and placebo capsules	
	Declarations of interest: No potential conflict of interest relevant to this article was	
	reported	
	Treatment adherence:	
	Compliance was assessed by counting returned capsules. Data on compliance were not	
	reported	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Before randomization, all eligible participants were stratified according to the median values for age and homocysteine concentration in the screening population. Random decimals between 0 and 1 were generated for each person in each of the four strata. Those below the median of the random numbers in each stratum were assigned to the vitamin group, and the remainder were assigned to the placebo group." Comment: adequate method of sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "Random decimals between 0 and 1 were generated for each person in each of the four strata. Those below the median of the random numbers in each stratum were assigned to the vitamin group, and the remainder were assigned to the placebo group." Comment: unclear method of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "We conducted a two-year, double-blind, placebo-controlled, randomized clinical trial" "All capsules were gelatin-coated, identical in color and shape, and packaged in blister packs." Comment: Participants and personnel were blinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "One of the authors administered all the cognitive tests." Comment: unclear whether outcome assessment was blinded

McMahon 2006 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: < 10% of randomised patients were not included in the analysis
Selective reporting (reporting bias)	Low risk	Comment: All the primary and secondary outcomes mentioned in the Methods and protocol were reported
Other bias	Low risk	Comment: none identified

Methods	2×2 factorial randomised placebo-controlled study of antioxidant vitamins and simvastatin. Mean duration of follow-up 5 years
Participants	Location: 69 Hospitals in the UK
	Recruitment: Medical collaborators from 69 UK hospitals appointed senior nurses to
	run special clinics for the study. With the permission of their relevant consultant col-
	leagues, records of patient discharges and of special wards or clinics were used to identify potentially eligible candidates for the study. The coordinating centre used this informa-
	tion to seek agreement, in the name of the local collaborator, from general practitioners
	to invite patients to the local study clinic
	Sample size:
	• Number randomised: 10,269 in intervention, 10,267 in comparison
	• Number completed: 10,241 in intervention, 10,228 in comparison
	Participant (baseline) characteristics:
	Inclusion criteria:
	1. Male or female aged 40 to 80 years;
	2. High risk of coronary heart disease death over the next 5 years:
	o (i) coronary disease: definite or probable clinical diagnosis of myocardial infrared and probable continuous parties. PTCA and CARC.
	infarction, unstable angina, stable angina, PTCA or CABG; o (ii) occlusive disease of non-coronary arteries: clinical, angiographic, or
	ultrasound diagnosis of carotid artery stenosis (e.g. transient ischaemic attack or non-
	disabling stroke not thought to be haemorrhagic), carotid endarterectomy, leg artery
	stenosis (e.g. intermittent claudication) or surgery;
	o (iii) diabetes mellitus: clinical diagnosis of insulin-dependent or non-insulin-
	dependent diabetes;
	or (iv) treated hypertension: use of antihypertensive drug therapy in male
	aged 65 years or over (in order to be at sufficiently high risk of vascular events);
	3. No clear indications for the study treatments: the patient is not already taking
	HMG CoA reductase inhibitors or high-dose vitamin E supplements, and neither the
	patient nor the patient's doctor considers there to be any definite need to do so;
	4. No clear contraindications to the study treatments:
	 (i) baseline plasma cholesterol: < 3.5mmol.L⁻¹; (ii) chronic liver disease (i.e. cirrhosis or hepatitis) or abnormal liver function
	(i.e. alanine transaminase > 1.5 x upper limit of normal);
	 (iii) severe renal disease or evidence of renal impairment (i.e. creatinine > 2 x
	upper limit of normal);

	 (iv) inflammatory muscle disease (such as dermatomyositis or polymyositis) or creatine kinase > 3 x upper limit of normal; (v) concurrent treatment with cyclosporin (or a condition likely to result in organ transplantation and the need for cyclosporin); (vi) concurrent treatment with fibrates or high-dose niacin. (N.B. patients on cholesterol-lowering diets or drugs - other than HMG CoA reductase inhibitors, fibrates or high-dose niacin - could still be entered in the study); or (vii) child-bearing potential (i.e. premenopausal woman who is not sterilised or using a reliable method of contraception); No other predominant medical problem: (i) severe heart failure or some importantly life-threatening condition other than vascular disease (such as very severe chronic airways disease or any cancer other than non-melanoma skin cancer); (ii) psychiatric disorder, senility or physical disability (such as severely disabling stroke); Or (iii) recent history of alcohol or drug abuse; Eligible patients who completed a pre-randomisation run-in phase on active treatment were randomly allocated to receive simvastatin (40 mg daily) or matching placebo tablets and, in a '2 x 2 factorial' design, antioxidant vitamins (600 mg vitamin E, 250 mg vitamin C, and 20 mg beta-carotene daily) or matching placebo capsule. Exclusion criteria: People were ineligible if they had life-threatening conditions, such as chronic liver disease, severe renal disease, severe heart failure, severe chronic airways disease, or diagnosed cancer (other than non-melanoma skin cancer). Those taking high-dose vitamin E supplements were not randomised
Interventions	Intervention (n=10,241): 600 mg synthetic vitamin E, 250 mg vitamin C and 20 mg β-carotene administered orally once per day for an average of 5 years Comparator group (n=10,228): "Matching placebo in specially prepared calendar packs" Use of additional interventions (common to both treatment arms): factorial design also included simvastatin
Outcomes	Outcomes of interest in the review: Global cognitive function measured with TICSm (The main comparisons for antioxidant vitamin supplementation were analyses of total coronary heart disease events and of fatal coronary heart disease events during the scheduled treatment period. Cognitive function was a secondary outcome. Quote: "It had been suggested that antioxidant vitamins (in particular, vitamin E) might slow cognitive decline, so the modified Telephone Interview for Cognitive Status (TICS-m) questionnaire was administered to participants during their final follow-up")
Notes	Funding sources: "The study was funded by the UK Medical Research Council, the British Heart Foundation, Merck & Co (manufacturers of simvastatin: J Tobert, R Tomiak, J Young, A Tate, E John, F Walker, G Warner) and Roche Vitamins Ltd (manufacturers of the vitamins: R Salkeld, E Stöcklin, M Wahl)." Declarations of interest: "P Sleight has received honoraria and costs for participating in meetings." Treatment adherence: Compliance with study treatment was assessed at each follow-up

MRC/BHF 2002 (Continued)

by reviewing the calendar packed capsules remaining and for those who had stopped,
the reasons for doing so were sought

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The central telephone randomisation system used a minimisation algorithm to balance the treatment groups with respect to eligibility criteria and other major prognostic factors." Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Comment: "central telephone randomisation system"
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Comment: "identical tablets"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Comment: Outcome assessors at study centres not aware of treatment allocation
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: Relatively low dropout rate and high number of participants completed follow-up at 5 years
Selective reporting (reporting bias)	Low risk	Comment: single cognitive outcome measure reported
Other bias	Low risk	Comment: no other risks identified

Rossom 2012

Methods	Women's Health Initiative Memory Study Post hoc analysis of a 2-arm randomised double-blind placebo-controlled trial	
Participants	Location: US multicentre Recruitment: Forty Women's Health Initiative (WHI) clinical centres across the United States. Participants were enrolled between 1994 and 1999 Number randomised: 2034 in intervention, 2109 in comparison	
	Participant (baseline) characteristics:	
	• Age: mean 71 years (65-80)	
	o vitamin D/calcium: mean 70.7 years	
	o placebo: mean 70.9 years	

- Gender: women 100%
- Main diagnosis: postmenopausal women aged 65 and older without probable dementia or cognitive impairment at baseline

Inclusion criteria: women aged 65 and older without probable dementia or cognitive impairment at baseline who participated in the WHI Calcium and Vitamin D Trial Other inclusion criteria specified in the main study: Eligible women were 50 to 79 years of age at the initial screening and had no evidence of a medical condition associated with a predicted survival of less than three years and no safety, adherence, or retention risks **Exclusion criteria:** diseases associated with significant risk of mortality (invasive cancer in the previous 10 years; any history of breast cancer or a suspicion of breast cancer at the time of screening; acute myocardial infarction, stroke, or transient ischaemic attack in the previous 6 months; known chronic active hepatitis or severe cirrhosis), safety (blood cell count indicative of disease, severe hypertension, current use of oral corticosteroids) , and adherence or retention (unwillingness or inability to complete baseline study requirements)

Interventions

Intervention (n=2034):

1000 mg of calcium carbonate and 400 IU of vitamin D3 Two tablets per day in divided doses and with meals

Comparator group (n=2109): placebo

Two tablets per day in divided doses and with meals

Supplements or placebo were provided by GlaxoSmithKline.

Active tablets, chewable or swallowable (after July 1997), contained 500 mg of elemental calcium (as calcium carbonate) and 200 IU of vitamin D

Use of additional interventions (common to both treatment arms):

Use of personal supplemental calcium up to 1000 mg/day and vitamin D up to 600 IU/day in addition to the study tablets was allowed

Calcium supplement use, n (%):

- vitamin D/calcium: 1061 (52.2)
- placebo 1196 (56.7)

Vitamin D supplement use, n (%):

- vitamin D/calcium: 929 (45.7)
- placebo 1047 (49.6)

See also Notes below

Outcomes

Outcomes of interest to the review:

Quote: "The Women's Health Initiative Calcium and Vitamin D trial (CaD) was designed to determine whether calcium plus vitamin D3 supplementation prevents hip fracture and colorectal cancer"

WHIMS (the Women's Health Initiative Memory Study) was an ancillary study

Primary outcomes of WHIMS: classifications of probable dementia, mild cognitive impairment and cognitively normal

Quote: "A diagnosis of dementia was based on Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition, criteria."

Quote: "A diagnosis of MCI was based on accepted criteria at the time of WHIMS initiation, operationally defined as performance in the 10th or lower percentile on at least one Consortium to Establish a Registry for Alzheimer's Disease (CERAD) test based on CERAD norms, a report from the designated informant of functional impairment on the part of the participant that was not severe enough to interfere with basic activities

Rossom 2012 (Continued)

of daily living, no evidence of a psychiatric or medical condition that could account for the cognitive decline, and absence of adjudicated dementia."

Secondary outcomes: global cognitive function as assessed annually using the Modified Mini Mental State Examination (3MS) and domain-specific cognitive functions as assessed annually in a subset of 1420 CaD participants using the WHISCA cognitive battery

The WHISCA cognitive battery included the Digit Span Forward and Backward Test, the Primary Mental Abilities Vocabulary test, the Card Rotations Test, letter and semantic fluency tests, the California Verbal Learning Test, the Benton Visual Retention Test, and the Finger Tapping Test

Depressive symptoms were measured using the Burnham screening algorithm Probable dementia or MCI were assessed during a mean follow-up of 7.8 years Domain-specific measures of cognitive function were obtained in the 1420 women enrolled in the CaD trial and WHISCA seen for an average of 4.9 (treatment) and 5.0 (placebo) annual visits through 2007

Eligible cognitive outcomes:

- global cognitive function measured with Modified MMSE (3MS)
- episodic memory measured with Benton Visual Retention Test
- episodic memory measured with California Verbal Learning Test
- speed of processing measured with finger tapping
- executive function measured with letter and category fluency at 5 years

Notes

"All WHIMS participants ... were also participants in the WHI Hormone Therapy trials. Some were also participants in a third trial of dietary modification to compare a low-fat eating pattern to a usual ad-libitum diet."

Funding sources, quote: "The WHI program is funded by the National Heart, Lung, and Blood Institute, National Institutes of Health, U.S. Department of Health and Human Services through Contracts N01WH22110, 24152, 32100-2, 32105-6, 32108-9, 32111-13, 32115, 32118-32119, 32122, 42107-26, 42129-32, and 44221."

"Sponsor's Role: The sponsors did not play any role in the design or conduct of the study; collection, management, analysis, or interpretation of the data; or preparation, review, or approval of the manuscript"

Declarations of interest, quote: "Dr. Rossom's work on this manuscript was supported in part by the Veterans Health Administration through Clinical and Health Services Research and Development Grants 04SCRCOE-001 and HFP-98-001. Dr. JoAnn Manson and colleagues at Brigham and Women's Hospital, Harvard Medical School, are recipients of funding from the National Institutes of Health to conduct the VITamin D and OmegA-3 TriaL, a large-scale randomized trial of vitamin D and omega-3s in the prevention of cancer and cardiovascular disease"

Treatment adherence

"Adherence was assessed annually in a structured interview and by weighing remaining pills."

Adherence (defined as > 80%):

- vitamin D/calcium: 1446 (72.3%)
- placebo: 1566 (75.5%)

Rossom 2012 (Continued)

Random sequence generation (selection bias)	Low risk	Quote: "Eligible women were randomly assigned in a double-blind fashion to receive supplements or placebo (provided by GlaxoSmithKline) in equal proportions with use of a permuted-block algorithm stratified according to clinical center and age" Comment: method of random sequence generation not reported, probably adequate
Allocation concealment (selection bias)	Unclear risk	Quote: "Eligible women were randomly assigned in a double-blind fashion to receive supplements or placebo (provided by GlaxoSmithKline) in equal proportions with use of a permuted-block algorithm stratified according to clinical center and age" Comment: method of allocation concealment not reported
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "1,000 mg of calcium carbonate and 400 IU of vitamin D or identical placebos.", "Eligible women were randomly assigned in a double-blind fashion" Comment: Participants and personnel were apparently blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "A central adjudication panel of experts at the WHIMS Clinical Coordinating Center (Wake Forest School of Medicine, Winston-Salem, NC) made classifications." Comment: not reported but assessors were probably blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "All primary outcomes were analyzed on a time to-event basis according to the intention-to-treat principle." Comment: All patients included in the post hoc analysis were considered for the main outcome whereas only a subset was analysed for cognitive tests
Selective reporting (reporting bias)	Low risk	Comment: All outcomes described in the methods or protocol (ClinicalTrials.gov number, NCT00000611) were reported in the results
Other bias	Low risk	No other risks identified

SEARCH 2010

Methods	2 x 2 factorial placebo-controlled randomised trial
Participants	Location: UK multicentre Recruitment: potentially eligible patients identified from hospital discharge records of 88 UK hospitals. Randomisation took place between September 1998 and October 2001 Number randomised: 6033 B vitamins, 6031 placebo Participant (baseline) characteristics: • Age: mean age 64.2 (8.9) years • B vitamins: 1885 < 60y; 2414 ≥ 60y and < 70y; 1734 ≥ 70y • placebo: 1880 < 60y; 2414 ≥ 60y and < 70y; 1737 ≥ 70y • Gender: • B vitamins: 83% male • placebo: 83% male • Main diagnosis: myocardial infarction Inclusion criteria: aged 18-80 years, myocardial infarction > 3 months before screening, taking a statin or clear indication for statin therapy Exclusion criteria: clear indication for folic acid, contraindications to study treatments, other predominant medical problem which might limit compliance with 5 years of study treatment
Interventions	Participants randomised in 2 x 2 factorial design to (folic acid 2mg + vitamin B12 1mg) or placebo and to simvastatin 20 mg or 80 mg
Outcomes	Outcomes relevant to this review: TICSm, verbal fluency test, diagnosis of dementia. All cognitive outcomes measured at final follow-up, after a mean of 6.7 (1.5) person-years
Notes	"The study was funded by Merck (manufacturers of simvastatin and suppliers of the vitamins)." "The funders had no role in the design and conduct of the study; collection, management, analysis and interpretation of the data; and preparation, review and approval of the manuscript."

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The central telephone randomization system used a minimization algorithm to balance the treatment groups with respect to major prognostic factors" Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "central telephone randomisation system" Comment: adequate method of allocation concealment
Blinding of participants and personnel (performance bias)	Unclear risk	Comment: no information regarding participants and personnel blinding

SEARCH 2010 (Continued)

All outcomes		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "All such information is reviewed by coordinating center clinical staff, blind to study treatment allocation, and events are coded according to prespecified criteria" Comment: outcome assessors were blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Comment: For 3173/12064 (26%) of the eligible patients, TICSm was not available
Selective reporting (reporting bias)	Low risk	Comment: no cognitive outcomes were included in the original protocol published in 1998. Additional information was obtained from the authors: "The cognitive assessment was added during the study (before any unblinding) and included in the Data Anaysis Plan but had not been specified in the original Protocol." We considered there to be a low risk of this introducing any bias
Other bias	Low risk	Comment: no other risks identified

Stott 2005

Methods	Factorial 2 X 2 x 2, randomised, placebo-controlled, double-blind study with 3 active treatments with 12-week treatment and 1-year follow-up
Participants	Location: United Kingdom and the Netherlands, 2 centres Recruitment: hospital-based Number randomised: 185 Participant (baseline) characteristics: • Age, mean ± SD: ○ placebo: 72.8 ± 5.4 ○ folic acid plus vitamin B12: 72.9 ± 6.0 ○ riboflavin (vitamin B2): 74.6 ± 5.3 ○ vitamin B6: 74.7 ± 6.1 ○ folic acid plus vitamin B12 plus vitamin B2: 76.5 ± 8.0 ○ folic acid plus vitamin B12 plus vitamin B6: 72.6 ± 6.4 ○ vitamin B6 plus vitamin B12 plus vitamin B2 plus vitamin B6: 74.0 ± 6.5 • Gender (male/female): ○ placebo: 14/10 ○ folic acid plus vitamin B12: 12/11 ○ riboflavin (vitamin B2): 10/13 ○ vitamin B6: 11/12 ○ folic acid plus vitamin B12 plus vitamin B2: 10/13

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	 o folic acid plus vitamin B12 plus vitamin B6: 12/11 o vitamin B6 plus vitamin B2: 10/13 o folic acid plus vitamin B12 plus vitamin B2 plus vitamin B6: 9/14 ◆ Main diagnosis: patients aged "65 y with ischemic vascular disease" Inclusion criteria: age ≥ 65 y and Ischaemic vascular disease, defined as one or more of the following: history of angina pectoris, previous acute myocardial infarction, evidence of major ischaemia or previous acute myocardial infarction on the basis of a 12-lead electrocardiogram, ischaemic stroke, transient ischaemic attack, intermittent claudication, or surgery for peripheral arterial disease Exclusion criteria: acute vascular event < 1 wk previously; major surgery < 1 mo previously; any other major acute illness < 1 mo previously; severe renal impairment (serum creatinine > 400 μmol/L); severe hepatic impairment; malignancy within the previous year (excluding local skin cancer); severe congestive heart failure (New York Heart Association class IV); total anterior cerebral infarct with major residual disability; malabsorption; inability to give informed consent (e.g. due to dementia or dysphasia); major cognitive impairment (Mini-Mental State Examination score < 19); existing treatment with riboflavin, vitamin B6, vitamin B12, or folic acid preparations; haemoglobin concentration < 10 g/dL; and mean cell volume > 100 fL plus either a low red blood cell folate concentration (< 280 ng/mL) or a low serum vitamin B12 concentration (< 250 pg/mL)
Interventions	Intervention: Folic acid 2.5 mg plus vitamin B12 400 mcg; vitamin B6 25 mg; riboflavin (vitamin B2) 25 mg; placebo in a 2 x 2 x 2 factorial design 1. folic acid plus vitamin B12: 23 2. vitamin B6: 23 3. riboflavin (vitamin B2): 23 4. folic acid plus vitamin B12 plus vitamin B2: 23 5. folic acid plus vitamin B12 plus vitamin B6: 23 6. vitamin B6 plus vitamin B12 plus vitamin B2 plus vitamin B6: 23 Comparator: placebo: 24 Intervention or placebo were continued for 12 weeks. The daily dose was provided in a total of 2 capsules (1 red and 1 white), irrespective of patient group Quote: "The patients initially entered a single-blind, placebo, run-in phase lasting between 2 and 4 weeks. Patients who successfully completed the run-in phase were randomly allocated" Use of additional interventions (common to both treatment arms): not reported
Outcomes	Overall cognitive function was assessed by using the Telephone Interview for Cognitive status (TICSm) at both baseline visits and at 6 and 12 months; attention and speed of information processing were assessed by using the Letter Digit Coding Test (at both baseline visits and 12 months) Serious adverse events, including incident vascular events, were recorded at each review
Notes	Funding sources , quote: "Supported by a grant from the Healthcare Foundation (reference 112/57)." Declarations of interest , quote: "None of the authors had any conflicting or competing interests, and the funder of the study had no role in data collection, analysis, or inter-

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pretation	of the data	a or in the	writing (of the	report.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Allocation was determined at a site remote from the clinical study (Robertson Centre for Biostatistics) in randomized permuted blocks of 8, stratified by hospital center." Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "Allocation was determined at a site remote from the clinical study (Robertson Centre for Biostatistics) in randomized permuted blocks of 8, stratified by hospital center." Comment: adequate method of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Treatment allocation was concealed from the patients and the investigators (double-blind)." Comment: adequate method of participants and personnel blinding
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Treatment allocation was concealed from the patients and the investigators (double-blind)." Comment: adequate method of outcome assessment blinding
Incomplete outcome data (attrition bias) All outcomes	High risk	Comment: Numbers of participants missing outcome data calculated from tables. 18/185 (10%) missing data for TICSm at 1 year. 39/185 (21%) missing data for LDCT at 1 year. Dropouts not mentioned in paper. No reasons for missing data given. Presumed completers analysis
Selective reporting (reporting bias)	High risk	Quote: from ISRCTN registry "Retrospectively registered" "Primary outcome measures: 1. Serum HCY, plasma von Willebrand factor, vitamin levels (red cell folate, serum vitamin B12, B2, and B6) 2. Piloting of telephone follow-up of cog-

Stott 2005 (Continued)

		nition (Telephone Interview for Cognitive Status (TICSm)) and of disability (Barthel Index and short Instrumental Activities for Daily Living (ADL) scale). Secondary outcome measures: not provided at time of registration." Comment: Disability and IADL (relevant to this review) not reported in paper	
Other bias	Low risk	Comment: no other risks identified	
Toole 2004			
Methods			
Participants	Recruitment: university-affilia practices, and Veterans Affairs Number randomised: 3680 (1) min group) Participant (baseline) charact • Age, mean (SD): • low-dose: 66.2 (10.8 • high-dose: 66.4 (10.9 • Gender: • low-dose: 37.2% fer • high-dose: 37.7% fer • baseline cognitive function • low-dose: 26.9 (3.3) • high-dose: 26.9 (3.4) Inclusion criteria: non-disablic with onset ≤ 120 days before rather for North American stroke portion dose vitamins for 1 month before Exclusion criteria: potential expectancy < 2 years, untreated pressure > 185 mmHg or dias severe cognitive impairment, all medications that affect total horizontal affect tot	 Participant (baseline) characteristics: Age, mean (SD): low-dose: 66.2 (10.8) high-dose: 66.4 (10.8) Gender: low-dose: 37.2% female high-dose: 37.7% female Baseline cognitive function, MMSE mean (SD): low-dose: 26.9 (3.3) high-dose: 26.9 (3.4) Inclusion criteria: non-disabling ischaemic stroke (Modified Rankin Stroke Scale ≤ 3) with onset ≤ 120 days before randomisation; total homocysteine level ≥ 25th percentile for North American stroke population; age ≥ 35 years, compliance ≥ 75% with low-dose vitamins for 1 month before randomisation Exclusion criteria: potential sources of emboli, other major neurological illness, life expectancy < 2 years, untreated anaemia or untreated B12 deficiency, systolic blood pressure > 185 mmHg or diastolic blood pressure > 105 mmHg, refractor depression, severe cognitive impairment, alcoholism or other substance abuse, use within 30 days of medications that affect total homocysteine level, childbearing potential, participation in another trial with active intervention, general anaesthesia or hospital stay of ≥ 3 days, 	
Interventions	by the US Food and Drug Ad	alation contained the reference daily intakes recommended dministration for vitamins except for its content of folic B12. Folic acid 2.5 mg, vitamin B6 25 mg, vitamin B12	

Toole 2004 (Continued)

	Low-dose multivitamin formulation contained the reference daily intakes recommended by the US Food and Drug Administration for vitamins (contained folic acid 20 μg , vitamin B6 0.2 mg, vitamin B12 6 μg)
Outcomes	Outcomes relevant to review: mean MMSE at 1-year follow-up
Notes	"All eligible participants were given the low-dose vitamin formulation for 1 month to determine compliance, assessed by pill counts. Only persons taking at least 75% of the vitamins during the run-in phase were eligible to be randomized."

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Participants were randomized to the high-dose or low-dose vitamin groups within strata defined by clinic, sex and age (≥ 70 years, < 70 years). Permuted block randomization (with block size randomly selected as 4 or 6) was used."
Allocation concealment (selection bias)	Low risk	Quote: "The allocation of participants was programmed by the statistical coordinating center, encrypted, and entered into a data entry program installed on a study computer at each site. After computer verification that all eligibility criteria had been met, participants were randomly assigned 1 of 20 medication codes. Allocation information was accessible only to the drug distribution center, which bottled and distributed the vitamins to clinics, and to selected coordinating center personnel who could assist with randomization in case of computer failure."
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Both pill formulations were manufactured (Magno-Humphries Laboratories, Tigard, Ore) to be indistinguishable by external colour, weight or dissolution in water. No request was ever made to break the blind."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Both pill formulations were manufactured (Magno-Humphries Laboratories, Tigard, Ore) to be indistinguishable by external colour, weight or dissolution in water. No request was ever made to break the

Toole 2004 (Continued)

		blind." No information specifically on site investigators. Blind probably maintained
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Mean MMSE score available for 3097/3680 (84.2%) participants at 1-year follow-up. No information on missing participants, including group allocation
Selective reporting (reporting bias)	High risk	Protocol-specified MMSE at 24 months not reported. Number of participants with MMSE score in each treatment group at 12 months not reported
Other bias	Low risk	No other risks identified

Methods	Multicentre, double-blind, randomised, placebo-controlled trial, with 2-year duration of treatment and 2-year duration of follow-up
Participants	Location: 3 research centres in the Netherlands: VU University Medical Center (Ams terdam), Erasmus Medical Center (Rotterdam), and Wageningen University (WU, Wageningen) Recruitment: Dutch elderly men and women who participated in the B-PROOF (B Vitamins for the Prevention of Osteoporotic Fractures) study which assessed the efficacy of oral supplementation with 400 µg folic acid and 500 µg vitamin B in the prevention
	of fractures, with cognitive performance as secondary outcome Most participants were recruited via registries of municipalities in surroundings of the research centres
	The study was conducted between October 2008 and March 2013. Number randomised: 1516 in intervention, 1511 in comparison Participant (baseline) characteristics:
	• Age (mean, SD):
	o total population (n=2919): 74.1 years (6.5)
	o cognitive sub-sample (n=856):
	♦ B vitamin (n=425): 72.6 years (5.7)
	♦ placebo (n=431): 72.6 years (5.8)
	• Gender (males, n %):
	o total population (n=2919): 1459 (50)
	o cognitive sub-sample (n=856):
	♦ B vitamin (n=425): 244 (57)
	♦ placebo (n=431): 255 (59)
	Main diagnosis: not reported Indian principles of 6 more and day having an almost diagnosis have a market and almost diagnosis.
	Inclusion criteria: age 65 years or older, having an elevated plasma homocysteine leve (12-50 μmol/L), being competent to make own decisions, and having a compliant table intake (≥ 85%) in the run-in period
	Exclusion criteria: cancer diagnosis within the last 5 years except for basal cell carcinoma
	and squamous cell carcinoma, bedridden, serum creatinine level > 150 µmol/L, curren

Van der Zwaluw 2014 (Continued)

	or recent (< 4 months) use of intramuscular injections of vitamin B12 or folic acid supplementations (> 300 μmol), and participation in other intervention studies 6% of the study population had a MMSE < 25.
Interventions	Intervention (n=1516): B vitamin group: 400 μg folic acid and 500 μg vitamin B12 daily Comparator group (n=1511): placebo. Treatment was continued for 2 years. Use of additional interventions (common to both treatment arms): Both tablets contained 15 μg (600 IU) vitamin D3.
Outcomes	Outcomes of interest in review: Quote: "The primary goal of the B-PROOF study was [to] assess the efficacy of oral supplementation with folic acid and vitamin B12 in the prevention of fractures, with cognitive performance as a secondary outcome." The 'primary' outcome of the cognitive study was "the difference between treatment groups in performance on episodic memory after 2 years of supplementation." 'Secondary' cognitive outcomes were "differences in performance at follow-up on attention and working memory, information processing speed, executive function, and single test scores." Quote: "Cognitive performance was assessed at baseline and at the end of the intervention period by well-trained research assistants following a standard protocol. We used the MiniMental State Examination (MMSE) and an extensive, sensitive, and validated neuropsychological test battery"that "consisted of the Rey Auditory Verbal Learning Test (RAVLT), Digit Span forward and backward, Trail Making Test (TMT) parts A and B, Stroop Color-Word Test, Symbol Digit Modalities Test, and Letter Fluency (3 letters)" Quote: "To compare results of the individual cognitive tests and to limit the number of dependent variables, crude test scores were clustered into compound scores for the 4 neuropsychological domains" "For all 4 cognitive domains, predefined interaction terms of treatment with sex, baseline age (< 80, ≥ 80 years), plasma HCY (< 14.4, ≥ 14.4 µmol/L), and APOE genotype (APOE e4 or not) were tested" Cognitive outcomes: • global cognitive function measured with MMSE at 2 years, with higher values indicating benefit • executive functioning measured with Stroop Interference (part 3 - (part 1 + part 2/2/)) at 2 years, with lower values indicating benefit • working memory measured with Digit span forward at 2 years, with higher values indicating benefit • working memory measured with RAVLT-Decay (delayed recall trial 5) at 2 years, with higher values indicating benefit • episodic memory measured with RAVLT- trials I-V, imm

- episodic memory measured with RAVLT- trial 9 Recognition max 30 words at 2 years, with higher values indicating benefit
- speed of processing measured with Trail making part A at 2 years, with lower values indicating benefit
- speed of processing measured with Stroop 1 and 2 mean at 2 years, with lower values indicating benefit
- speed of processing measured with Symbol digit modalities test at 2 years, with higher values indicating benefit
- verbal fluency measured with Verbal Fluency total no at 2 years, with higher values indicating benefit

Notes

Funding sources, quote: "B-PROOF was supported and funded by The Netherlands Organization for Health Research and Development (ZonMw, grant 6130.0031), the Hague; unrestricted grant from NZO (Dutch Dairy Association), Zoetermeer; MCO Health, Almere; NCHA (Netherlands Consortium Healthy Ageing) Leiden/Rotterdam; Ministry of Economic Affairs, Agriculture and Innovation (project KB-15-004-003), the Hague; Wageningen University, Wageningen; VU University Medical Center, Amsterdam; and Erasmus Medical Center, Rotterdam. All organizations are based in the Netherlands. The sponsors had no role in the design or implementation of the study, data collection, data management, data analysis, data interpretation, or in the preparation, review, or approval of the manuscript."

Declarations of interest, quote: "The authors report no disclosures relevant to the manuscript."

Treatment adherence

"Average compliance to treatment was 90%, and 84% of all participants had an overall compliance ≥ 80%. Compliance was equal for both treatment groups"

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was done by an independent person by means of computergenerated randomization numbers in stratified permuted blocks of size 4, stratified by study center, sex, age (65-79 years, \geq 80 years), and HCY levels (12-17 mmol, \geq 18 mmol)" Comment: computer-generated sequence
Allocation concealment (selection bias)	Low risk	Quote: "Randomization was done by an independent person". Comment: allocation probably concealed
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Participants and employees of the study were blinded until data analyses were finished." Comment: participants and personnel blinded

Van der Zwaluw 2014 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Participants and employees of the study were blinded until data analyses were finished." Comment: outcome assessors blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: from CONSORT flow diagram, 22-24% attrition from each group Comment: significant loss of participants, but reasons given and appeared balanced between groups
Selective reporting (reporting bias)	Low risk	Comment: No differences identified in the outcomes reported in the protocol (clinicaltrials.gov: NCT00696514) and the article. All outcomes specified in the Methods were reported in the Results
Other bias	Low risk	Comment: no other apparent sources of bias

Walker 2012

Methods	$2 \times 2 \times 2$ factorial design, parallel group, double-blinded, multicentre, duration of treatment 24 months, duration of follow-up 24 months
Participants	 Location: Australia. 3 sites: 2 urban and 1 rural Recruitment: The population-based sample was recruited by a direct mailing of a screening survey and consent form to 105,000 randomly selected adults aged from 60 to 74 years Number randomised: 453 in intervention, 456 in comparison Participant (baseline) characteristics: Age, mean (±SD): overall: 65.94 ± 4.24 folic acid-vitamin B12: 65.92 ± 4.30 placebo: 65.97 ± 4.18 Gender, male, n (%): overall: 358 (39.8) folic acid-vitamin B12: 181 (40.5) placebo: 177 (39.1) Main diagnosis: elevated psychological distress (Kessler Distress 10 Scale > 15) Inclusion criteria: cohort of community-dwelling older 60-74 y adults with elevated psychological distress (Kessler Distress 10-Scale; scores > 15); did not engage in physical activity at public health-recommended levels as indicated by International Physical Activity Questionnaire scores; did not take folic acid, vitamin B12, or vitamin B complex supplements; had no history of dementia, bipolar disorder, or current suicide risk; had competent literacy skills; and did not have a medical condition that would contraindicate exercise or folic acid use Exclusion criteria: High likelihood of a depressive disorder with K10 scores of ≥ 30

Walker 2012 (Continued)

	were excluded. Those individuals with low concentrations of red blood cell folate (< 250 nmol/L) and vitamin B12 (<130 nmol/L), and abnormal thyroid stimulating hormone concentrations (0.35-5.0 mU/L) were excluded because participation may have led to potentially adverse outcomes	
Interventions	Intervention: 1) Daily oral dose of one tablet consisting of 400 μg folic acid + 100 μg vitamin B12 supplementation (Matchland Pty Ltd, t/a New Products Development -ABN57052101176, Brisbane, Australia) for 24 months The protocol changed to 2 daily oral doses (200 μg folic acid + 50 μg vitamin B12 each) from July 2007 Comparator group: Drug: placebo. Placebo tablets were manufactured by the same producers of the folic acid + vitamin B12 tablets and were identical except for the omission of the active substances under investigation Use of additional interventions (common to both treatment arms): Due to the factorial design, in both vitamin and placebo groups, approximately half the participants received an intervention to promote physical activity and approximately half info about nutrition for older adults; and approximately half received a mental health literacy intervention and approximately half received arthritis and pain management info	
Outcomes	Outcomes of interest in the review: Primary outcomes: • TICSm total score Secondary outcomes: • TICSm subscores: 1) orientation; 2) registration, recent memory, and delayed recall (immediate and delayed recall reported separately); 3) attention/calculation; and 4) semantic memory, comprehension, and repetition • Processing speed measured with BTACT • IQCODE	
Notes	Funding sources, quote: "Supported by beyond blue: the national depression initiative and the Australian Government Department of Health and Ageing. HC was supported by a National Health and Medical Research Council (NHMRC) fellowship no. 525411. JGW was supported by NHMRC Capacity Building Grant 418020" "The sponsors had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; or preparation, review, or approval of the manuscript." Declarations of interest, quote: "None of the authors had any conflicts of interest." Treatment adherence "Adherence was monitored by telephone assessment at 14 time points and by blood assay at baseline and at 12- and 24-mo assessments."	
Risk of bias		

Walker 2012 (Continued)

Random sequence generation (selection bias)	Low risk	Quote: "Randomisation followed the screening assessment with block size fixed at eight and using strata comprising location, gender and high (19) and low (16-18) K10 depression scores (conducted by A.J.M.) using an automated computerised system." Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "using an automated computerised system". Comment: adequate method of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Participants, interviewers, investigators and the survey administrators were masked to active intervention and folic placebo (<i>sic</i>) allocation." Comment: Participants and personnel were blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Participants, interviewers, investigators and the survey administrators were masked to active intervention and folic placebo (<i>sic</i>) allocation." Comment: Outcome evaluators were blinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Of those who were recruited into the trial, the dropout rate was low, with only 123 (13.5%) participants withdrawing from the time of randomisation to the 24-mo assessment" Comment: Reasons for withdrawal balanced across groups
Selective reporting (reporting bias)	High risk	Comment: All cognitive results were reported, but not all were reported in sufficient detail for inclusion. Results of this study were therefore missing from some analyses
Other bias	Low risk	Comment: no other important risks identified

Wolters 2005

Methods	2-arm, parallel group, placebo controlled, double-blind RCT. Duration of treatment 6 months, duration of follow-up 6 months
Participants	Location: Germany, single centre Recruitment: recruited by newspaper advertisement in Hannover region. Institute of Food Science, University of Hanover Number randomised: not reported. Baseline measures were taken from 241 women. Data are reported for 220 who completed the study Participant (baseline) characteristics (provided for those who completed): • Age, median (5 th -95 th percentile) • overall: 63 years (60-74) • vitamin: 63 years (60-74) • placebo: 64 years (60-75) • Gender: female 100% • Main diagnosis: healthy volunteers Inclusion criteria: healthy females 60 years or older; no vitamin supplementation for two months before intervention; no medications that affect vitamin absorption on status at the time of enrolment Exclusion criteria: not reported
Interventions	Intervention (n=111): Vitamin capsule (Nobilin Q10) contained 150 mg vitamin C (calcium ascorbate), 50 mg magnesium (magnesium carbonate), 36 mg vitamin E (D- alfa-tocopherol acetate), 34 mg niacin (nicotinamide), 16 mg pantothenic acid, 9 mg β-carotene, 3.4 mg pyridoxine (pyridoxine hydrochloride), 3.2 mg riboflavin, 2.4 mg thiamine (thiamine mononitrate), 400 μg folic acid (pteroyl glutamic acid), 200 μg biotin, 60 μg selenium (selenium-enriched yeast), and 9 μg cobalamin (cyanocobalamin) One capsule/day at breakfast for 6 months Comparator group (n=109): placebo One capsule/day at breakfast for 6 months Use of additional interventions: 4.5% women received statin medication.
Outcomes	Primary Outcomes: Wechsler Adult Intelligence Scale-Revised (WAIS III) Intelligence assessed by the Kurztest fuer Allgemeine Intelligenz (KAI) Mild up to severe memory disorders assessed by a subtest of the Berliner Amnesie Test (BAT) Secondary outcomes: levels of homocysteine and methylmalonic acid at baseline and end of treatment Eligible cognitive outcomes: • episodic memory measured with Pattern recognition subtest of Berliner Amnesie Test (BAT) (higher values indicate benefit) • speed of processing measured with WAIS-III revised symbol search test (higher values indicate benefit) Function outcome extracted: none reported Quality of life outcome extracted: none reported Safety outcome extracted: none reported *: outcome data included in statistical analyses

Wolters 2005 (Continued)

Notes	Funding sources: Medicom Pharma. Vitamin capsules (Nobilin Q10R) and placebo capsules were donated by Medicom Pharma AG, Springe (Germany) Declarations of interest: no information provided	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Of these, 109 women had been randomly assigned to the placebo group and 111 women to the vitamin group by a lottery." Comment: unclear method of sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "Of these, 109 women had been randomly assigned to the placebo group and 111 women to the vitamin group by a lottery." Comment: unclear method of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Both capsules, the multivitamin and the placebo, were soft gelatine capsules filled with soy oil. The oil in the placebo capsule was colored for identical appearance." Comment: probably blinded
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The cognitive tests were performed on the day of blood sampling after the women had had breakfast." Comment: blinding of outcome assessment not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	8.7% (21/241 assessed at baseline) dropped out, but it is unclear from which group. Two participants were excluded by the experimenters for non-compliance. The total number of patients randomised was unclear. Only completers' data reported
Selective reporting (reporting bias)	Low risk	All primary and secondary outcomes specified in the Methods were reported
Other bias	Unclear risk	There were baseline imbalances in educational level (higher in vitamin group) and

Symbol Search (better in vitamin group)

Methods	4-arm, double-blinded, multicentre parallel group RCT, with up to 8 years of treatmen
Participants	Location: 11 clinical centres in the US
1 articipants	Recruitment: recruited through 11 clinical centres in the US
	Sample size: Quote "Of the 3640 AMD trial participants, 2166 (60%) completed the
	AREDS cognitive battery near the end of the trial."
	• Number randomised: 2737 in intervention, 903 in comparison (total 3640)
	• Number completed cognitive battery: 1632 in intervention, 534 in comparison
	(total 2166)
	Group A (Antioxidants) baseline characteristics:
	• Age (y), %:
	o 55-59: 4
	0 60-64: 16
	o 65-69: 34
	0 70-74: 29
	o 75-80: 16
	• Median age, y: 69
	• Gender, % female: 55
	Group B (Zinc) baseline characteristics:
	• Age (y), %:
	o 55-59: 2
	o 60-64: 18
	o 65-69: 29
	0 70-74: 34
	o 75-80: 18
	Median age, y: 70
	• Gender, % female: 57
	Group C (Antioxidants and Zinc) Baseline characteristics:
	• Age (y), %:
	o 55-59: 3
	0 60-64: 17
	o 65-69: 33
	o 70-74: 30
	o 75-80: 17
	Median age, y: 69
	• Gender, % female: 56
	Group D (Placebo) Baseline characteristics:
	• Age (y), %:
	o 55-59: 3
	0 60-64: 16
	o 65-69: 32
	0 70-74: 31
	o 75-80: 18
	Median age, y: 69
	• Gender, % female: 56
	Inclusion criteria:
	Aged 55 to 80 years old
	Individuals who passed the screening evaluation and were interested in participating in
	AREDS were provided with a one-month "run-in" supply of placebo to assess potentia

Yaffe 2004 (Continued)

Bias	Authors' judgement	Support for judgement
Risk of bias		
Notes	This was a cognitive substudy of AREDS, the Age-Related Eye Disease Study. Funding sources: "Supported by contracts from the National Eye Institute and the National Institute on Aging, National Institutes of Health, Department of Health and Human Services, Bethesda, MD, with additional support from Bausch & Lomb Inc, Rochester, NY" Declarations of interest: none declared. "The Writing Team for this report and the AREDS investigators have no commercial or proprietary interest in the supplements used in this study" Treatment adherence: At each visit, participants returned their used study medication bottles and any unused tablets, and received new bottles of their study medication. "Estimated pill counts showed that most patients took 75% or more of the assigned medications and adherence was balanced by treatment"	
Outcomes	Outcomes of interest in the review: The primary purpose of the study was to investigate the effects of the interventions on the progression of age-related macular degeneration The AREDS cognitive function battery included six validated cognitive tests with eight components: 1) The Modified Mini-Mental State Examination (3MS); 2) Animal Category; 3) Letter Fluency; 4) Logical Memory Part I and 5) Logical Memory Part II, Wechsler Memory Scale Revised; 6) Immediate Recall and 7) Word List Mean, Bushke Selective Reminding Test; and 8) Digits Backwards	
Interventions	morning and 2 each evening to meet the to Group A (Antioxidants): 500 mg Vitamin C daily Group B (Zinc and Copper): 80 mg zinc or Group C (Antioxidants and Zinc and Copp 15 mg beta carotene, 80 mg zinc oxide, 2 m	2, 400 IU Vitamin E and 15 mg beta carotene xide and 2 mg cupric oxide daily ber): 500 mg Vitamin C, 400 IU Vitamin E, ng cupric oxide daily n orally that was identical in external appearant taste acebo group were included in this review
	They were asked to take two tablets twice a than 75% of the prescribed tablets were ine good compliance with the run-in medication of disqualifying lesions noted on photogra and who signed a second consent form were of randomisation Exclusion criteria:	and compliance with the treatment regimen. It day for 1 month. Participants taking fewer eligible for enrolment. Participants who had son, who had adequate pupillary dilation and phs sent to the Reading Center for grading, the stratified into AMD categories at the time of that would make long-term follow-up or yor difficult

Yaffe 2004 (Continued)

Random sequence generation (selection bias)	Low risk	Quote: "Eligibility verification and random treatment assignment were performed by the Coordinating Center using the on-site computers, with procedures to protect the integrity of randomization and to provide randomization backup in case of hardware malfunction. Multiple levels of data encryption ensure the integrity of the treatment assignment files." "Each treatment assignment database residing on the hard drives at each Clinical Center is encrypted and includes check numbers to insure tamper free operation and proper sequential treatment assignments. There were no cases of database corruption during randomization." Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "Eligibility verification and random treatment assignment were performed by the Coordinating Center using the on-site computers, with procedures to protect the integrity of randomization and to provide randomization backup in case of hardware malfunction. Multiple levels of data encryption ensure the integrity of the treatment assignment files." "Each treatment assignment database residing on the hard drives at each Clinical Center is encrypted and includes check numbers to insure tamper free operation and proper sequential treatment assignments. There were no cases of database corruption during randomization." Comment: adequate allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "Four participants (0.1%) were reported to have been unmasked during the trial." "AREDS is a double-masked study. The AREDS participants, investigators and Reading Center personnel are masked to study-wide outcome data and treatment assignments." Comment: low risk for personnel and participants. Although 4 participants were "unmasked", this is likely to have been for

Yaffe 2004 (Continued)

		safety reasons
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Comment: "AREDS is a double-masked study. The AREDS participants, investi- gators and Reading Center personnel are masked to study-wide outcome data and treatment assignments."
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "Another limitation is that not all participants in the trial had cognitive testing." "Of the 3,640 AMD trial participants, 2, 166 (60%) completed the AREDS cognitive battery near the end of the trial. Of the 1,474 participants without cognitive testing, 330 died before the implementation of the cognitive study, and 1,144 refused or were otherwise unable to participate. Participants without cognitive testing were more likely to be older, have more ocular abnormalities, and be less educated compared with those participating in the ancillary study." No information was given about the treatment allocation of participants and non-participants in the cognitive sub-study Comment: high risk of differential non-participation
Selective reporting (reporting bias)	Unclear risk	Comment: This was an ancillary study and did not have a protocol of its own. The AREDS study design did not mention any cognitive outcomes
Other bias	Low risk	Comment: no other risks identified

AD:Alzheimer'sdisease

AD8: Ascertain Dementia 8-Item Informant Questionnaire (AD8) Dementia Screening Interview

ADAS-cog: Alzheimer's Disease Assessment Scale - cognitive

ADL: Activities of Daily Living

AMD: Age-related Macular Degeneration

APOE $\epsilon 4$: Apolipoprotein-E gene, $\epsilon 4$ allele

AREDS: The Age-Related Eye Disease Study

BASF: BASF is a registered chemical company, BASF originally stood for Badische Anilin- und Soda-Fabrik.

BAT: Berliner Amnesie Test BDI: Beck Depression Inventory

betaCarot: ß-carotene

B-PROOF: B-Vitamins for the Prevention of Osteoporotic Fractures study

BTACT: Brief Test of Adult Cognition by Telephone

CaD: Calcium and Vitamin D

CANTAB: Cambridge Automated Neuropsychological Test Battery

CDR: Clinical Dementia Rating scale

CERAD: Consortium to Establish a Registry for Alzheimer's Disease

CERAD e: expanded Consortium to Establish a Registry in Alzheimer's Disease

CERAD T: CERAD battery total score

CoA: coenzyme A reductase

CONSORT: Consolidated Standards of Reporting Trials

CVD: CardioVascular Disease

CVLT: California Verbal Learning Test

DCT: Digit Cancellation Test

DSM-IV: Diagnostic and Statistical Manual of Mental Disorders, 4th Edition

EBMT: East Boston Memory Test

FA: Folic acid

F00: ICD-10 code for dementia in Alzheimer's disease

F01: ICD-10 code for vascular dementia F03:ICD-10code for unspeci fieddementia

F-TICSm: modified Telephone Interview for Cognitive Status

G30: ICD-10 code for Alzheimer's disease

GDS: Geriatric Depression Scale

HbA1c: Hemoglobin A1c

HCY: homocysteine

HMG-COA: 3-hydroxy-3-methyl-glutaryl-coenzyme A reductase

holoTC: Holotranscobalamin

IADL: Instrumental Activities of Daily Living

IQCODE: Informant Questionnaire on Cognitive Decline in the Elderly

IQR: Interquartile range

ISRCTN: International Standard Randomised Controlled Trial Number

KAI: Kurztest fuer Allgemeine Intelligenz

K10: Kessler Distress 10-Scale LDCT: Letter Digit Coding Test. MCI: Mild cognitive impairment

MI: Memory Impairment

MIS: Memory Impairment Screen

MMA: Methylmalonic acid MMSE: Mini-Mental State Examination MoCA: Montreal Cognitive Assessment

MRI: Magnetic resonance imaging

msec: millisecond

MTS: Matching To Sample visual search test

n: number

NYU: New York University PHS: Physicians' Health Study

RAVLT: Rey Auditory Verbal Learning Test

RBC: Red blood cell

RDA: recommended daily allowances

RI-48: rappel indicé-48 items (cued-recall test-48 items)

SD: standard deviation

SF-36: 36-item Short Form Survey

SU.FOL.OM3: SUpplementation with FOlate, vitamin B6 and B12 and/or OMega-3 fatty acids trial

RCT: randomised clinical trial SWM: spatial working memory

TCogS: Telephone Cognitive Screen TIA: Transient Ischaemic Attack

TICS: Telephone Interview for Cognitive Status

TICSm: Modified Telephone Interview for Cognitive Status

TMT: Trail Making Test

VISP: Vitamin Intervention for stroke prevention trial

vit: vitamin

Vit D: vitamin D supplementation

WACS: Women's Antioxidant and Cardiovascular Study WADLS: Western Australian Data Linkage System

WAFACS: Women's Antioxidant and Folic Acid Cardiovascular Study

WAIS: Wechsler Adult Intelligence Scale

WHI: Women's Health Initiative

WHIMS: Women's Health Initiative Memory Study

WHISCA: Women's Health Initiative study of Cognitive Aging

wk: week y: year Zn: zinc

3MS: The Modified Mini-Mental State Examination

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Abbasi 2013	No cognitive outcomes
ACTRN12607000321448	Ineligible population: participants had MCI
Alavi 2014	Ineligible population. Additional detail: included in related Cochrane review focussing on the MCI study population
Almeida 2010	Subpopulation of included study focussing on depression outcome. Participants also included in Hankey 2013
Anonymous 2003	Ineligible population
Arwert 2003	Ineligible intervention
Benton 1995	Ineligible population
Bryan 2002	Ineligible intervention: duration of intervention 5 weeks
Chan 2010	Ineligible experimental intervention: combined experimental intervention
Cheng 2016	Wrong design: non-randomised

Chew 2015	Wrong design: RCT with 2x2 factorial design (omega-3 fatty acids; lutein and zeaxanthin; the combination; double placebo) with a second randomisation to multivitamin supplements with or without beta carotene and with or without high dose of zinc. The analyses of effects of beta carotene and zinc were not eligible, as no tests for interaction were performed for omega-3, lutein and zeaxanthin
Corless 1987	Wrong population
De Jager 2012	Ineligible population: participants had MCI
Dhesi 2004	Ineligible intervention: vitamin D delivered parenterally
Flicker 2011	Wrong population: all had vitamin D deficiency
Ford 2008	Ineligible outcomes
Harris 2015	Ineligible intervention: included plant extracts and probiotics as well as vitamins and minerals
Hvas 2004	Ineligible population: 56% cognitively impaired at baseline
Imaoka 2016	Ineligible population: participants cognitively impaired at baseline
Krikorian 2010	Ineligible population: participants had CDR (Clinical Dementia Rating scale) classification consistent with MCI
Kwok 2017	Ineligible population: We were unable to obtain separate data for participants with mild cognitive impairment (CDR 0.5) and normal cognition (CDR 0) at baseline
Li 2015	Ineligible population. Baseline cognitive status unclear. Data presented pictorially only (bar charts). Baseline mean MMSE < 23
Liu 2015	Study withdrawn prior to participant enrolment (ClinicalTrials.gov Identifier: NCT01317849)
Loriaux 1985	Ineligible experimental intervention
Macpherson 2012	Ineligible population: participants had subjective cognitive decline
Maniam 2004	Other reason. Additional details: potentially eligible RCT described only in an abstract published in 2004. No related full report has been identified and the likelihood such report will be published in future was judged to be minimal
McNeill 2007	No eligible outcomes
Miller 2016	No eligible outcomes
Murray-Kolb 2011	Ineligible population: age range 19-55 years
NCT00903695	Ineligible population: participants had MCI

(Continued)

NCT01095211	Ineligible population: duration of intervention 45 days
NCT01708005	Ineligible intervention: combined vitamins, endophospholipids, and omega-3 fatty acids
NCT02185222	Ineligible population: participants had memory complaint
NCT02467153	No eligible outcomes
Ng 2017	Ineligible intervention: combined protein, calorie, and vitamin supplements
Ong 2016	Ineligible intervention: no placebo control
Pase 2015	Ineligible intervention: combined experimental intervention
Pathansali 2006	Ineligible intervention: duration of intervention 4 weeks
Petersen 2005	Ineligible population. Additional detail: included in related Cochrane review focussing on population with MCI
Pettersen 2017	Ineligible intervention: no placebo control
Quick 2013	Ineligible design
Rietsema 2014	Ineligible design
Schietzel 2016	Ineligible intervention: no placebo control
Smith 1999	Ineligible population: Excluded if MMSE < 18, but no information on mean MMSE; likely to have included participants with dementia or MCI
Summers 2010	Ineligible experimental intervention: combined experimental intervention
Van Uffelen 2007	Ineligible population. Additional detail: included in related Cochrane review focussing on population with MCI
Wouters-Wesseling 2005	Ineligible experimental intervention: combined experimental intervention

MC1:Mildcognitiveimpairment CDR:Clinical DementiaRatingscale MMSE:MiniMentalStateExamination

Characteristics of studies awaiting assessment [ordered by study ID]

Sanchez 2011

Methods	Cluster double-blind, placebo controlled trial with 18 month follow-up
Participants	Location: metropolitan Santiago, Chile, 15 health centres, 20 participants selected from each health centre Setting of recruitment and treatment: GP practices Number randomised: not known Participant (baseline) characteristics: not known Inclusion criteria: Potential study subjects will be identified from the registry of older people participating in the PACAM in 15 health centres 1. men and women aged 70 to 79 years 2. participating in the National Health Service Nutrition Program for Older People (PACAM) Exclusion criteria: 1. diabetes (fasting blood glucose greater than or equal to 126 mg/dL) 2. hypothyroidism (thyroid stimulating hormone (TSH) greater than 6 mlU/L) 3. serum creatinine greater than or equal to 30 mg/mL 4. serum vitamin B12 level less than 120 or greater than 700 pmol/L or vitamin B12 supplemented 5. cerebrovascular disease 6. previous gastrointestinal surgery 7. unexplained weight lost greater than or equal to 3 kg in the previous 3 months 8. low cognitive development (Mini-mental test less than 20, and Pfeffer test greater than 5)
Interventions	
Outcomes	Primary Outcomes: 1. serum vitamin B12 status, assessed at initial visit and 3, 9, and 18 months 2. haematological and biochemical parameters, assessed at initial visit and 3, 9, and 18 months 3. peripheral nerve conduction, assessed at initial visit, 3, and 18 months 4. compound muscle action potential (CMAP) amplitude, distal motor latency, and F-wave latency (a measure of conduction time from the distal stimulation site to the spinal cord), assessed at initial visit, 3, and 18 months 5. Mini-mental score: Mini Mental State Examination (MMSE), assessed at initial visit and 18 months Secondary Outcomes: 1. serum folate, homocysteine, methyl malonic acid, holotranscobalamin, assessed at initial visit and 3, 9, and 18 months

Sanchez 2011 (Continued)

	 neurological battery evaluation includes sensory pathways of conduction of afferent sensory pathways of small diameter, quantitative thermal and somatosensory responses, specific neurosensory pathways, and visual evoked potential. Assessed at initial visit, 3, and 18 months. cognitive function: Test Wechsler, Consortium to Establish a Registry for Alzheimer's Disease (CERAD), Test Hamilton and Trail Making Test, assessed at initial visit and 18 months Compliance with the nutrition intervention is defined for this study as collecting more than 1 kg per month of the fortified food as documented by the health centres. Compliance for the pill intervention will be monitored based on collection of the pills supplement from the health centre, 12 out of 18 months will be considered compliant A subsample of 120 subjects (8 per cluster, 40 per arm) will undergo neurophysiological tests
Notes	Funding sources: Chilean National Science and Technology Research Fund (FONDECYT 1070592) (Fondo de Desarrollo Científico y Tecnológico (Chile) (ref: FONDECYT Nº 1070592)), Ministry of Health (Chile) Declarations of interest, quote: "The authors declare that they have no competing interests." Reasons for classification as "study awaiting classification": This study is marked as "completed" in ISRCTN registry (last edited 11-10-2011), but no results are available. We made several attempts to contact authors of the protocol, but received no replies

CERAD: Consortium to Establish a Registry for Alzheimer's Disease

CMAP: compound muscle action potential

PACAM: National Health Service Nutrition Program for Older People

DATA AND ANALYSES

Comparison 1. B vitamins versus control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Overall cognitive functioning	10		Std. Mean Difference (Random, 95% CI)	Subtotals only
1.1 3-12 months of supplementation (final measurement data)	3	2556	Std. Mean Difference (Random, 95% CI)	0.00 [-0.08, 0.08]
1.2 3-12 months of supplementation (change from baseline data)	2	317	Std. Mean Difference (Random, 95% CI)	-0.03 [-0.26, 0.19]
1.3 > 1-2 years of supplementation (final measurement data)	5	6956	Std. Mean Difference (Random, 95% CI)	-0.01 [-0.08, 0.06]
1.4 > 1-2 years of supplementation (change from baseline data)	1	241	Std. Mean Difference (Random, 95% CI)	0.06 [-0.20, 0.31]
1.5 > 2-5 years of supplementation (final measurement data)	4	3985	Std. Mean Difference (Random, 95% CI)	-0.01 [-0.11, 0.09]
1.6 > 5-10 years of supplementation (final measurement data)	3	10249	Std. Mean Difference (Random, 95% CI)	-0.01 [-0.09, 0.07]
2 Episodic memory	10		Std. Mean Difference (Random, 95% CI)	Subtotals only
2.1 3-12 months of supplementation (final measurement data)	4	1442	Std. Mean Difference (Random, 95% CI)	0.02 [-0.08, 0.12]
2.2 3-12 months of supplementation (change from baseline data)	2	430	Std. Mean Difference (Random, 95% CI)	0.04 [-0.15, 0.23]
2.3 > 1-2 years of supplementation (final measurement data)	4	5625	Std. Mean Difference (Random, 95% CI)	0.05 [-0.03, 0.13]
2.4 > 1-2 years of supplementation (change from baseline data)	1	241	Std. Mean Difference (Random, 95% CI)	0.06 [-0.19, 0.32]
2.5 > 2-5 years of supplementation (final measurement data)	3	3523	Std. Mean Difference (Random, 95% CI)	-0.05 [-0.15, 0.05]
2.6 > 5-10 years of supplementation (final measurement data)	1	1053	Std. Mean Difference (Random, 95% CI)	-0.10 [-0.22, 0.02]
3 Executive function	7		Std. Mean Difference (Random, 95% CI)	Subtotals only
3.1 3-12 months of supplementation (final measurement data)	3	536	Std. Mean Difference (Random, 95% CI)	0.14 [-0.18, 0.46]

3.2 3-12 months of supplementation (change from baseline data)	1	171	Std. Mean Difference (Random, 95% CI)	0.39 [0.07, 0.71]
3.3 > 1-2 years of supplementation (final measurement data)	3	2976	Std. Mean Difference (Random, 95% CI)	-0.01 [-0.15, 0.13]
3.4 > 2-5 years of supplementation (final measurement data)	2	2648	Std. Mean Difference (Random, 95% CI)	-0.06 [-0.13, 0.02]
3.5 > 5-10 years of supplementation (final measurement data)	1	1049	Std. Mean Difference (Random, 95% CI)	0.02 [-0.10, 0.15]
4 Speed of processing	7		Std. Mean Difference (Random, 95% CI)	Subtotals only
4.1 3-12 months of supplementation (final measurement data)	2	289	Std. Mean Difference (Random, 95% CI)	-0.04 [-0.28, 0.19]
4.2 3-12 months of supplementation (change from baseline data)	3	487	Std. Mean Difference (Random, 95% CI)	0.12 [-0.34, 0.58]
4.3 > 1-2 years of supplementation (final measurement data)	2	1030	Std. Mean Difference (Random, 95% CI)	0.03 [-0.09, 0.15]
4.4 > 1-2 years of supplementation (change from baseline data)	1	241	Std. Mean Difference (Random, 95% CI)	-0.04 [-0.29, 0.21]
4.5 > 2-5 years of supplementation (final measurement data)	1	818	Std. Mean Difference (Random, 95% CI)	-0.08 [-0.22, 0.06]
5 Incidence of MCI or dementia	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
5.1 > 2-5 years of	1	1803	Risk Ratio (M-H, Random, 95% CI)	1.01 [0.69, 1.48]
supplementation				
6 All-cause mortality	4		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
6.1 3-12 months of supplementation	1	201	Risk Ratio (M-H, Random, 95% CI)	0.34 [0.01, 8.33]
6.2 >1 to 2 years of supplementation	1	3680	Risk Ratio (M-H, Random, 95% CI)	0.86 [0.66, 1.11]
6.3 >2-5 years of supplementation	1	1127	Risk Ratio (M-H, Random, 95% CI)	1.98 [1.22, 3.21]
6.4 >5-10 years of supplementation	1	12064	Risk Ratio (M-H, Random, 95% CI)	1.03 [0.95, 1.12]

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Overall cognitive functioning	5		Mean Difference (Random, 95% CI)	Subtotals only
1.1 3-12 months Vitamin E	1		Mean Difference (Random, 95% CI)	1.4 [1.18, 1.62]
supplementation				
1.2 3-12 months Beta	1		Mean Difference (Random, 95% CI)	-0.14 [-0.37, 0.09]
Carotene supplementation				
1.3 > 2-5 years antioxidants (Vitamin E + Vitamin C + Beta Carotene) supplementation	1		Mean Difference (Random, 95% CI)	0.09 [-0.05, 0.23]
1.4 > 2-5 years Vitamin E supplementation	1		Mean Difference (Random, 95% CI)	0.01 [-0.23, 0.25]
1.5 > 2-5 years Vitamin C supplementation	1		Mean Difference (Random, 95% CI)	0.16 [-0.08, 0.40]
1.6 > 2-5 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	-0.18 [-0.42, 0.06]
1.7 > 5-10 years antioxidants (Vitamin E + Vitamin C + Beta carotene) supplementation	1		Mean Difference (Random, 95% CI)	0.6 [-0.20, 1.40]
1.8 > 5-10 years Vitamin C supplementation	1		Mean Difference (Random, 95% CI)	0.46 [0.14, 0.78]
1.9 > 5-10 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	-0.13 [-0.46, 0.20]
1.10 > 10 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	0.18 [0.01, 0.35]
2 Episodic memory	4		Mean Difference (Random, 95% CI)	Subtotals only
2.1 3-12 months Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	0.02 [-0.05, 0.08]
2.2 > 2-5 years Vitamin E supplementation	1		Mean Difference (Random, 95% CI)	-0.02 [-0.07, 0.03]
2.3 > 2-5 years Vitamin C supplementation	1		Mean Difference (Random, 95% CI)	-0.05 [-0.10, -0.00]
2.4 > 2-5 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	0.01 [-0.05, 0.07]
2.5 > 5-10 years antioxidants (Vitamin E + Vitamin C + Beta Carotene) supplementation	1		Mean Difference (Random, 95% CI)	-0.3 [-1.30, 0.70]
2.6 > 5-10 years Vitamin E supplementation	2		Mean Difference (Random, 95% CI)	0.02 [-0.05, 0.09]
2.7 > 5-10 years Vitamin C supplementation	1		Mean Difference (Random, 95% CI)	-0.14 [-0.22, -0.06]
2.8 > 5-10 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	0.02 [-0.05, 0.09]
2.9 > 10 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	-0.06 [-0.11, -0.02]
3 Executive function	4		Mean Difference (Random, 95% CI)	Subtotals only

3.1 3-12 months Beta	1		Mean Difference (Random, 95% CI)	0.07 [-0.48, 0.62]
Carotene supplementation				
3.2 > 2-5 years Vitamin E	1		Mean Difference (Random, 95% CI)	0.42 [0.06, 0.78]
supplementation				
3.3 > 2-5 years Vitamin C	1		Mean Difference (Random, 95% CI)	-0.03 [-0.39, 0.33]
supplementation				
3.4 > 2-5 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	0.22 [-0.14, 0.58]
3.5 > 5-10 years antioxidants	1		Mean Difference (Random, 95% CI)	-0.4 [-0.99, 0.19]
(Vitamin E + Vitamin C + Beta				
Carotene) supplementation				
3.6 > 5-10 years Vitamin E	2		Mean Difference (Random, 95% CI)	0.11 [-0.26, 0.48]
supplementation				
3.7 > 5-10 years Vitamin C	1		Mean Difference (Random, 95% CI)	-0.25 [-0.70, 0.20]
supplementation				
3.8 > 5-10 years Beta Carotene	1		Mean Difference (Random, 95% CI)	0.01 [-0.44, 0.46]
supplementation				
3.9 > 10 years Beta Carotene supplementation	1		Mean Difference (Random, 95% CI)	0.01 [-0.36, 0.38]
4 Incidence of dementia	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
4.1 > 2-5 years of	1	20536	Risk Ratio (M-H, Random, 95% CI)	1.00 [0.61, 1.64]
supplementation				
5 All-cause mortality	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
5.1 > 2-5 years antioxidants	1	20536	Risk Ratio (M-H, Random, 95% CI)	1.04 [0.97, 1.11]
(Vitamin E + Vitamin				
C + + Beta Carotene)				
supplementation				

Comparison 3. Vitamin D3 versus placebo

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Overall cognitive functioning	1		Mean Difference (Random, 95% CI)	Subtotals only
1.1 3-12 months of supplementation	1	4122	Mean Difference (Random, 95% CI)	0.0 [-0.24, 0.24]
1.2 > 1-2 years of supplementation	1	3910	Mean Difference (Random, 95% CI)	0.0 [-0.25, 0.25]
1.3 > 2-5 years of supplementation	1	3515	Mean Difference (Random, 95% CI)	-0.2 [-0.53, 0.13]
1.4 > 5-10 years of supplementation	1	918	Mean Difference (Random, 95% CI)	-0.1 [-0.81, 0.61]
2 Episodic memory	1		Mean Difference (Random, 95% CI)	Subtotals only
2.1 > 5-10 years of supplementation	1	1420	Mean Difference (Random, 95% CI)	0.05 [-0.03, 0.13]
3 Executive function	1		Mean Difference (Random, 95% CI)	Subtotals only
3.1 > 5-10 years of supplementation	1	1420	Mean Difference (Random, 95% CI)	0.0 [-0.08, 0.08]
4 Speed of processing	1		Mean Difference (Random, 95% CI)	Subtotals only

4.1 > 5-10 years of	1	1420	Mean Difference (Random, 95% CI)	0.03 [-0.05, 0.11]
supplementation				
5 Incident MCI	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
5.1 > 5-10 years of	1	4143	Risk Ratio (M-H, Random, 95% CI)	0.94 [0.72, 1.23]
supplementation				
6 Incident all-cause dementia	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
6.1 > 5-10 years	1	4143	Risk Ratio (M-H, Random, 95% CI)	1.09 [0.70, 1.71]
supplementation				

Comparison 4. Zinc + copper versus placebo

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Overall cognitive functioning	1		Mean Difference (Random, 95% CI)	Subtotals only
1.1 > 5-10 years of supplementation	1	1072	Mean Difference (Random, 95% CI)	0.6 [-0.19, 1.39]
2 Episodic memory	1		Mean Difference (Random, 95% CI)	Subtotals only
2.1 > 5-10 years of supplementation	1	1072	Mean Difference (Random, 95% CI)	-0.7 [-1.74, 0.34]
3 Executive function	1		Mean Difference (Random, 95% CI)	Subtotals only
3.1 > 5-10 years of supplementation	1	1072	Mean Difference (Random, 95% CI)	-0.3 [-0.89, 0.29]

Comparison 5. Complex multivitamin versus control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Overall cognitive functioning	1		Mean Difference (Random, 95% CI)	Subtotals only
1.1 > 2-5 years of supplementation	1	5947	Mean Difference (Random, 95% CI)	0.04 [-0.09, 0.17]
1.2 > 5-10 years of supplementation	1	2324	Mean Difference (Random, 95% CI)	0.12 [-0.14, 0.38]
2 Episodic memory	2		Mean Difference (Random, 95% CI)	Subtotals only
2.1 3-12 months of supplementation	1	124	Mean Difference (Random, 95% CI)	33.17 [-28.12, 94. 46]
2.2 > 2-5 years of supplementation	1	5942	Mean Difference (Random, 95% CI)	-0.01 [-0.05, 0.03]
2.3 > 5-10 years of supplementation	1	2324	Mean Difference (Random, 95% CI)	-0.01 [-0.07, 0.05]
3 Executive function	1		Mean Difference (Random, 95% CI)	Subtotals only
3.1 > 2-5 years of supplementation	1	5942	Mean Difference (Random, 95% CI)	-0.02 [-0.33, 0.29]
3.2 > 5-10 years of supplementation	1	2324	Mean Difference (Random, 95% CI)	-0.22 [-0.65, 0.21]

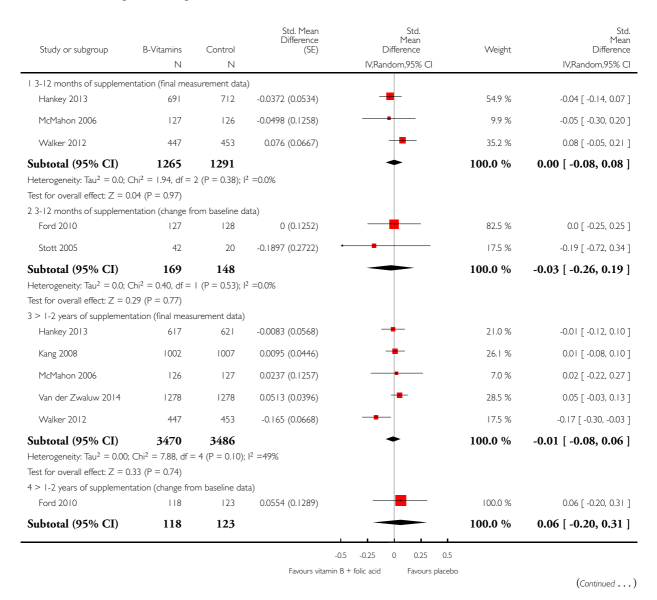
4 Speed of processing	1		Mean Difference (Random, 95% CI)	Subtotals only
4.1 3-12 months of	1	127	Mean Difference (Random, 95% CI)	-14.71 [-57.22, 27.
supplementation				80]

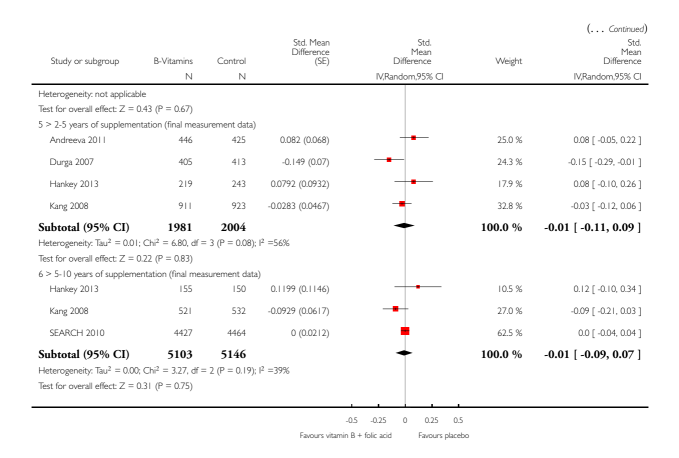
Analysis I.I. Comparison I B vitamins versus control, Outcome I Overall cognitive functioning.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: I B vitamins versus control

Outcome: I Overall cognitive functioning



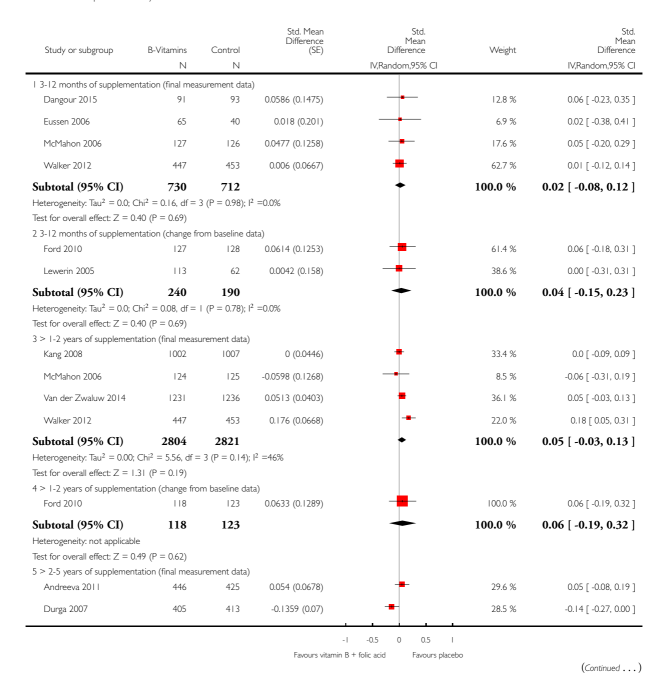


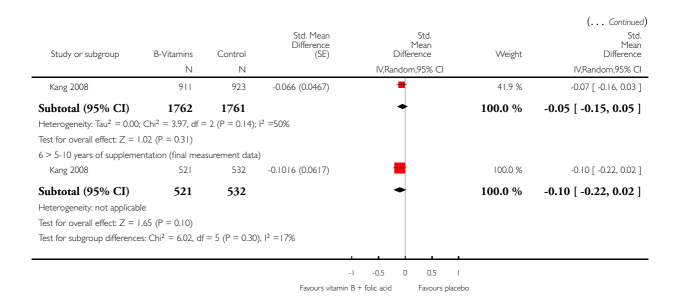
Analysis I.2. Comparison I B vitamins versus control, Outcome 2 Episodic memory.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: I B vitamins versus control

Outcome: 2 Episodic memory



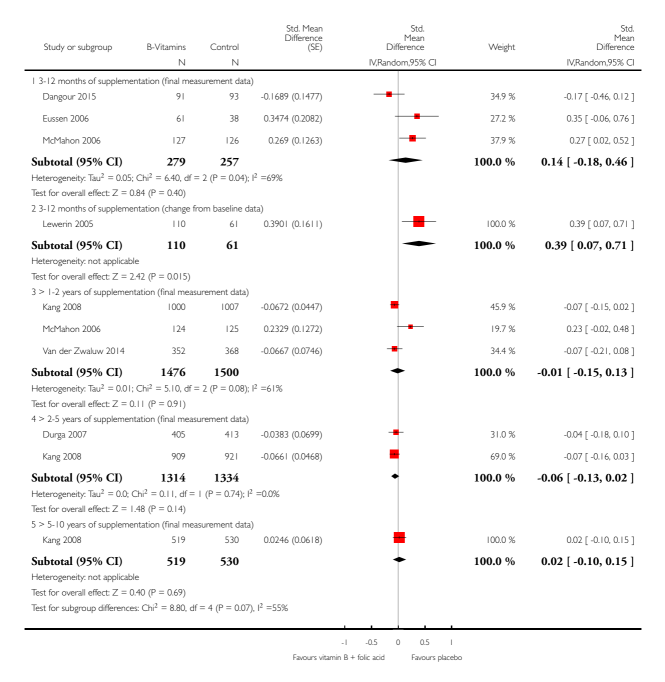


Analysis I.3. Comparison I B vitamins versus control, Outcome 3 Executive function.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: I B vitamins versus control

Outcome: 3 Executive function

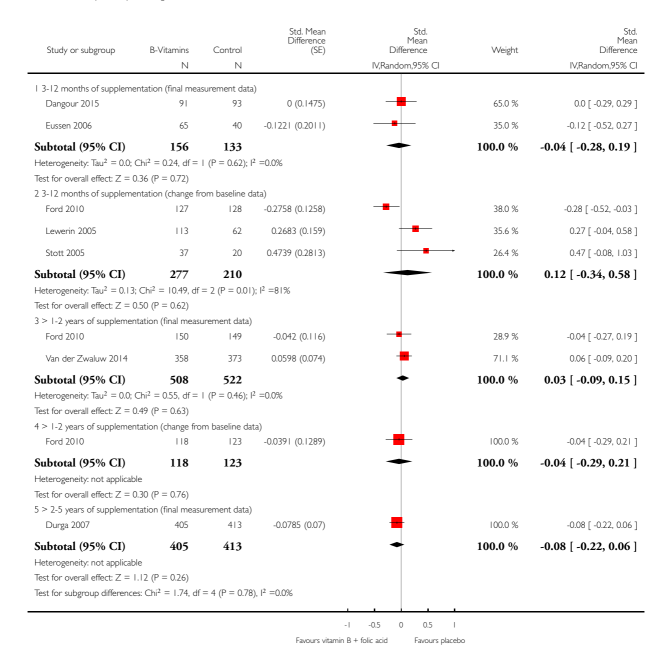


Analysis I.4. Comparison I B vitamins versus control, Outcome 4 Speed of processing.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: I B vitamins versus control

Outcome: 4 Speed of processing

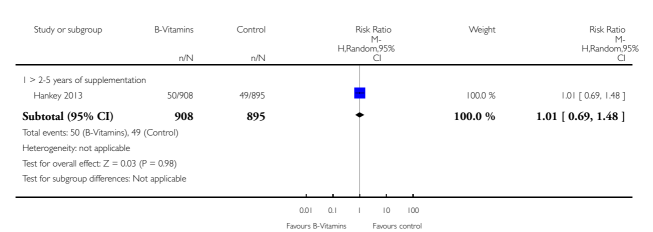


Analysis I.5. Comparison I B vitamins versus control, Outcome 5 Incidence of MCI or dementia.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: I B vitamins versus control

Outcome: 5 Incidence of MCI or dementia

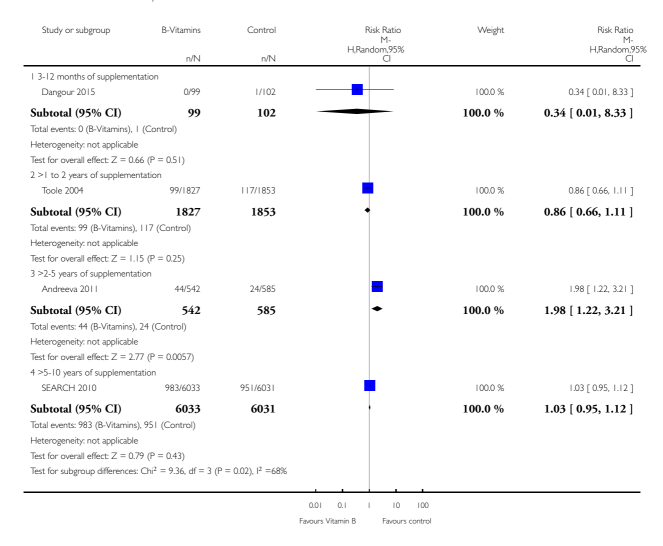


Analysis I.6. Comparison I B vitamins versus control, Outcome 6 All-cause mortality.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: I B vitamins versus control

Outcome: 6 All-cause mortality

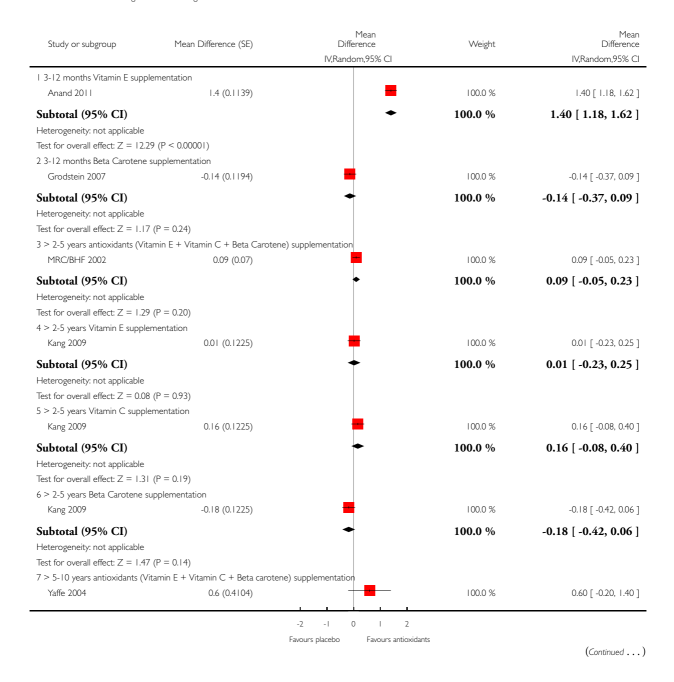


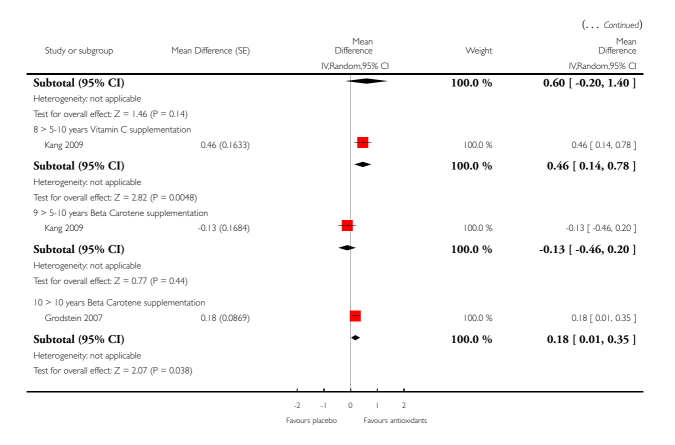
Analysis 2.1. Comparison 2 Antioxidants versus control, Outcome I Overall cognitive functioning.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 2 Antioxidants versus control

Outcome: I Overall cognitive functioning



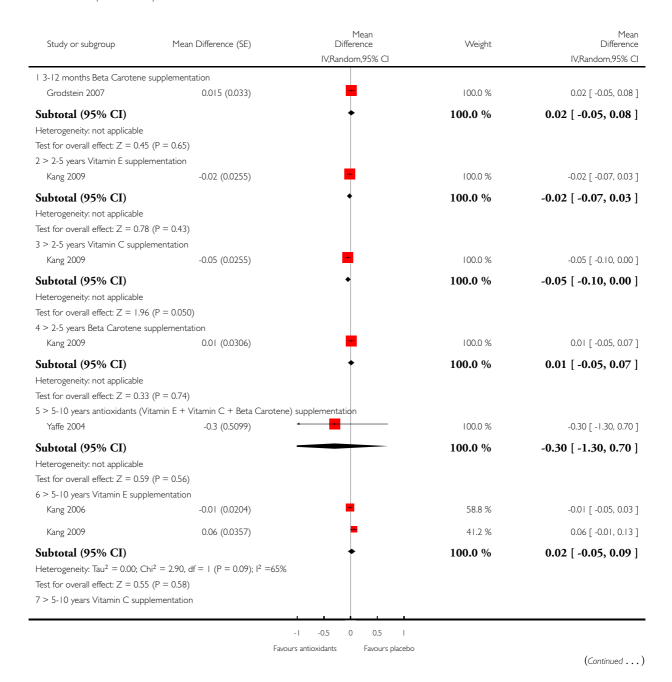


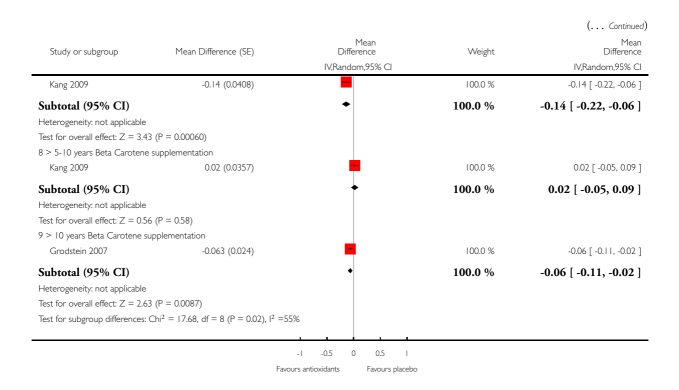
Analysis 2.2. Comparison 2 Antioxidants versus control, Outcome 2 Episodic memory.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 2 Antioxidants versus control

Outcome: 2 Episodic memory



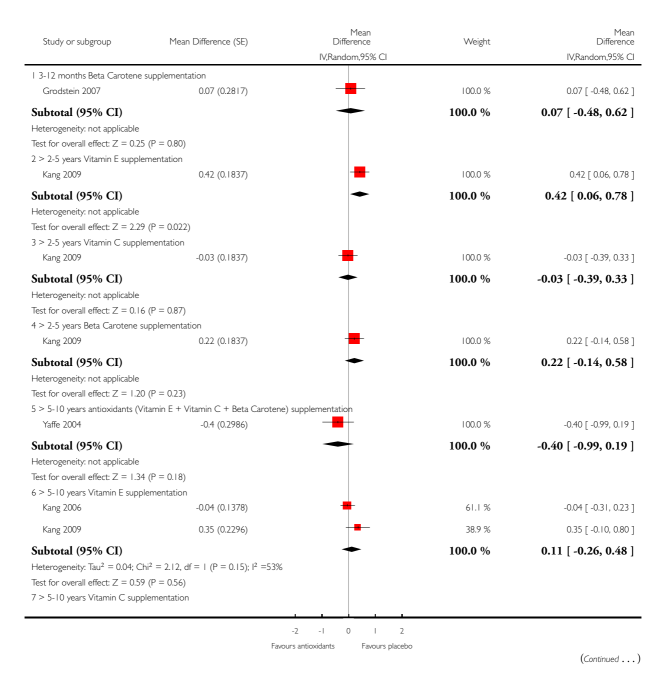


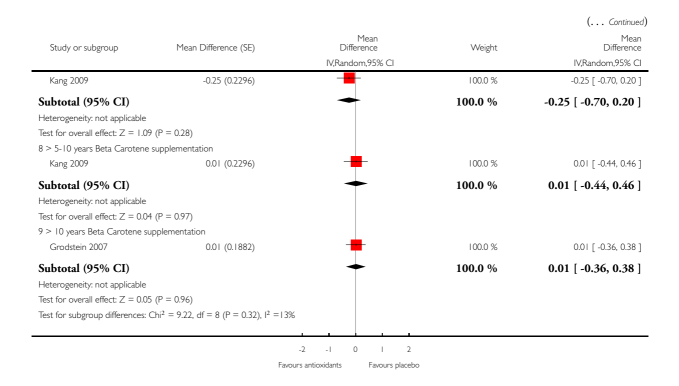
Analysis 2.3. Comparison 2 Antioxidants versus control, Outcome 3 Executive function.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 2 Antioxidants versus control

Outcome: 3 Executive function



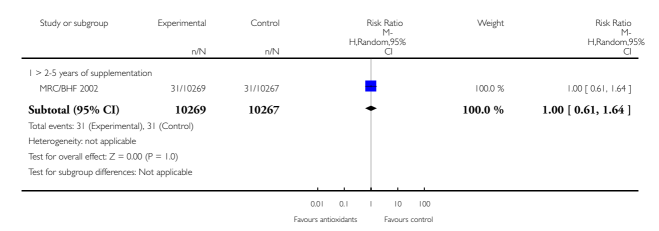


Analysis 2.4. Comparison 2 Antioxidants versus control, Outcome 4 Incidence of dementia.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 2 Antioxidants versus control

Outcome: 4 Incidence of dementia

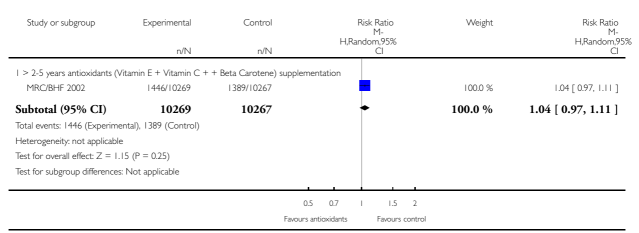


Analysis 2.5. Comparison 2 Antioxidants versus control, Outcome 5 All-cause mortality.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 2 Antioxidants versus control

Outcome: 5 All-cause mortality

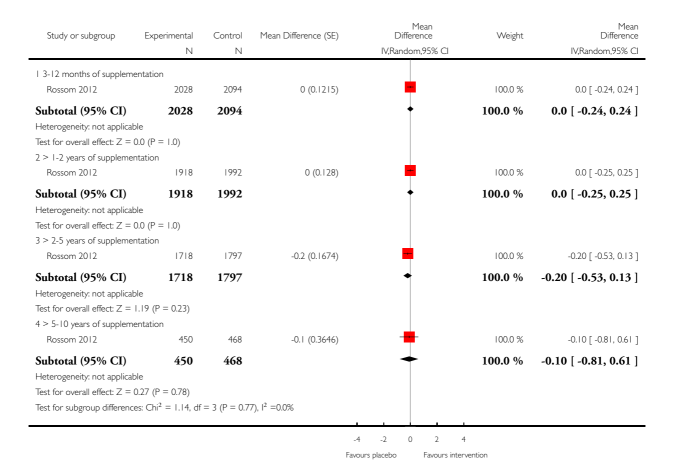


Analysis 3.1. Comparison 3 Vitamin D3 versus placebo, Outcome I Overall cognitive functioning.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 3 Vitamin D3 versus placebo

Outcome: | Overall cognitive functioning

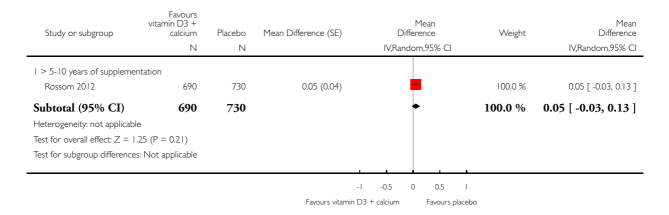


Analysis 3.2. Comparison 3 Vitamin D3 versus placebo, Outcome 2 Episodic memory.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 3 Vitamin D3 versus placebo

Outcome: 2 Episodic memory

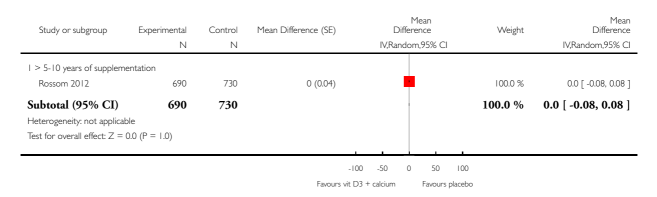


Analysis 3.3. Comparison 3 Vitamin D3 versus placebo, Outcome 3 Executive function.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 3 Vitamin D3 versus placebo

Outcome: 3 Executive function

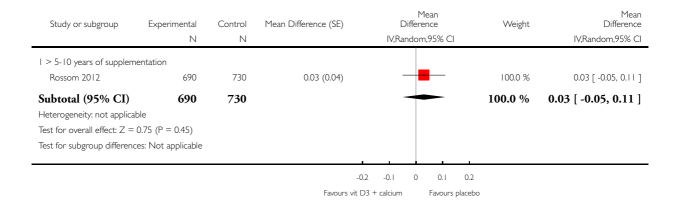


Analysis 3.4. Comparison 3 Vitamin D3 versus placebo, Outcome 4 Speed of processing.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 3 Vitamin D3 versus placebo

Outcome: 4 Speed of processing

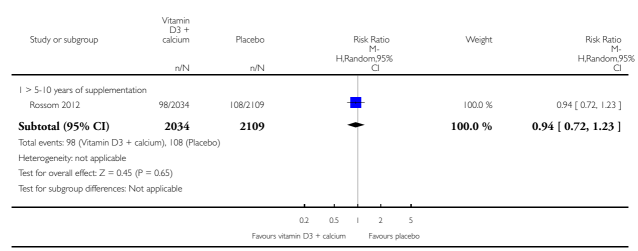


Analysis 3.5. Comparison 3 Vitamin D3 versus placebo, Outcome 5 Incident MCI.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 3 Vitamin D3 versus placebo

Outcome: 5 Incident MCI

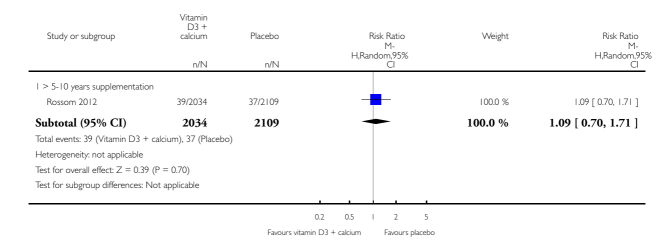


Analysis 3.6. Comparison 3 Vitamin D3 versus placebo, Outcome 6 Incident all-cause dementia.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 3 Vitamin D3 versus placebo

Outcome: 6 Incident all-cause dementia

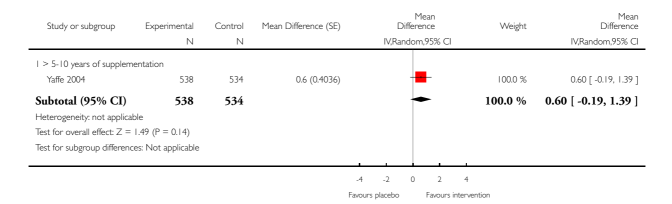


Analysis 4.1. Comparison 4 Zinc + copper versus placebo, Outcome I Overall cognitive functioning.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 4 Zinc + copper versus placebo

Outcome: | Overall cognitive functioning

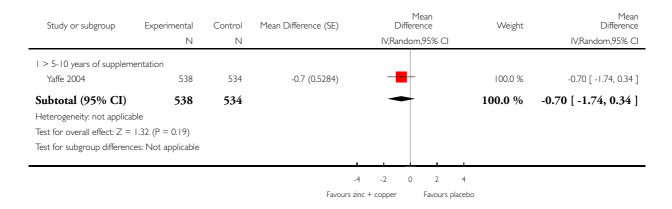


Analysis 4.2. Comparison 4 Zinc + copper versus placebo, Outcome 2 Episodic memory.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 4 Zinc + copper versus placebo

Outcome: 2 Episodic memory

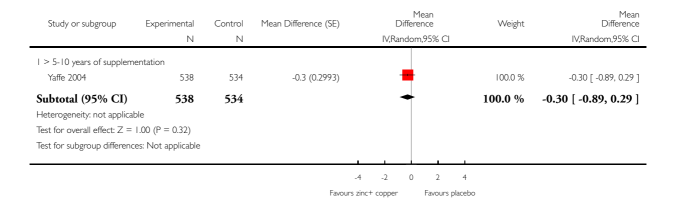


Analysis 4.3. Comparison 4 Zinc + copper versus placebo, Outcome 3 Executive function.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 4 Zinc + copper versus placebo

Outcome: 3 Executive function

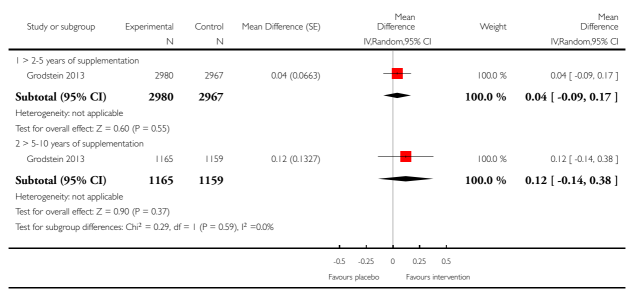


Analysis 5.1. Comparison 5 Complex multivitamin versus control, Outcome I Overall cognitive functioning.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 5 Complex multivitamin versus control

Outcome: I Overall cognitive functioning

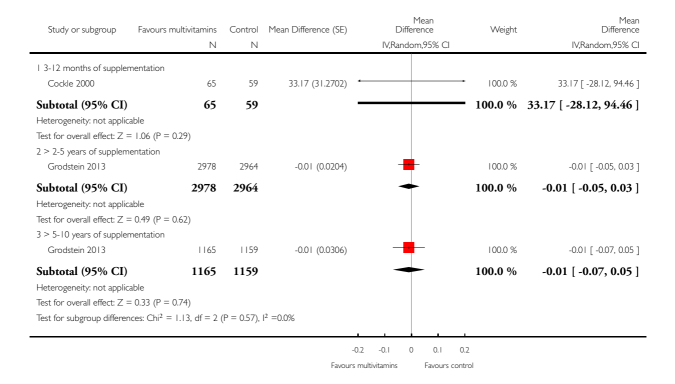


Analysis 5.2. Comparison 5 Complex multivitamin versus control, Outcome 2 Episodic memory.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 5 Complex multivitamin versus control

Outcome: 2 Episodic memory

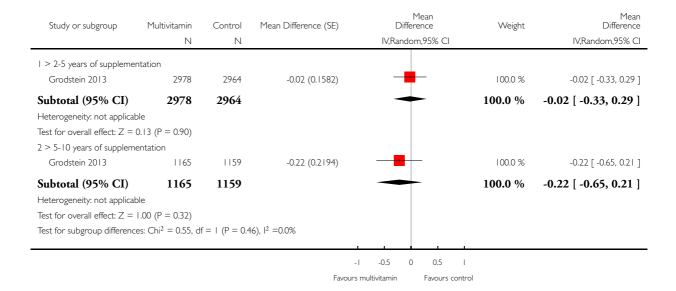


Analysis 5.3. Comparison 5 Complex multivitamin versus control, Outcome 3 Executive function.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 5 Complex multivitamin versus control

Outcome: 3 Executive function

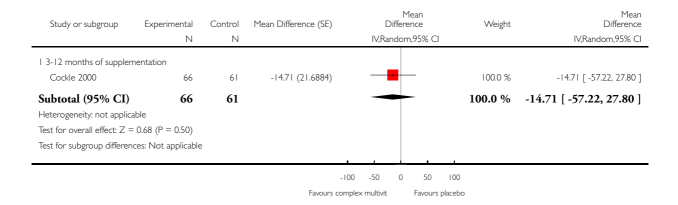


Analysis 5.4. Comparison 5 Complex multivitamin versus control, Outcome 4 Speed of processing.

Review: Vitamin and mineral supplementation for maintaining cognitive function in cognitively healthy people in mid and late life

Comparison: 5 Complex multivitamin versus control

Outcome: 4 Speed of processing



APPENDICES

Appendix I. Biological plausibility of vitamins and minerals

Supplement*	Route	Biological plausibility
Vitamins		
Vitamin A	antioxidant; anti-inflammatory; anti-cholinesterase; beta-amyloid inhibition	Carboxylic form of Vitamin A known as alltrans retinoic acid has been shown to have memory restorative function and it may be attributed to its anti-cholinesterase, anti-oxidative and anti-inflammatory potential (Sodhi 2013). Vitamin A and ß-carotene may also inhibit the formation, extension and destabilising effects of beta-amyloid fibrins. Plasma or cerebrospinal fluid concentrations of vitamin A and beta-carotene have been reported to be lower in AD patients, and increased Vit A/beta carotene

		concentrations have been clinically shown to slow the progression of dementia (Ono 2012).
Vitamin D	neuronal activity	Vitamin D receptor (VDR) and 1, alpha-hydroxylase, the terminal calcitriol-activating enzyme, are distributed throughout both the foetal and adult brain. This is thought to play a role in brain development and critical brain functions (McCann 2008). Significant correlation between serum 25(OH)D levels and cognitive scores were reported in DeLuca 1975 and Przybelski 2007.
Vitamin E	antioxidant; beta-amyloid inhibition	Vitamin E consists of a group of tocopherols and tocotrienols. Apart from lipid antioxidant activity, other functions include membrane stabilisation by forming complexes with the products of lipid hydrolysis (Wang 2000). It has been shown that the antioxidant and free radical scavenging activity of Vitamin E inhibits amyloid beta protein induced neuronal cell death and may have implication in prevention and treatment of Alzheimer's dementia (Behl 1992).
Vitamin K	neuronal activity	Vitamin K participates in the synthesis of sphingolipids. Sphingolipids participate in important cellular events such as proliferation, differentiation, senescence, and cell-cell interactions. Sphingolipid metabolism has been linked to age-related cognitive decline and neurodegenerative diseases such as Alzheimer's disease (Ferland 2012). A cross-sectional study found correlations between higher serum phylloquinone concentration and better cognitive scores in tests evaluating episodic verbal memory among healthy older adults (Ferland 2013)
Thiamine (Vitamin B1)	neuronal activity	Thiamine is required as a cofactor in the cellular production of energy and enhances normal neuronal activities (Osiezagha 2013). Rats with an episode of induced thiamine deficiency had cognitive, learning, and memory impairments (Langlais 1995)

Riboflavin (Vitamin B2)	neuronal activity	Riboflavin (7,8-dimethyl-10-ribityl-isoal-loxazine) is water-soluble. Symptoms of neurodegeneration and peripheral neuropathy in riboflavin deficiency have been documented in animal studies, but not observed in humans. Subclinical riboflavin deficiency may contribute to increased concentrations of plasma homocysteine and may be associated with increased risk of cardiovascular disease and impaired handling of iron (Powers 2003).
Niacin (Vitamin B3)	vascular: anti-inflammatory	Niacin is a water-soluble precursor cofactor essential for the formation of dozens of enzymes. Niacin decreases atherosclerosis development mainly by reducing LDL cholesterol. It also has modest HDL-cholesterol-raising and anti-inflammatory effects (Kühnast 2013). Niacin deficiency causes pellagra. Its neuropsychiatric symptoms are similar to those in Alzheimer's disease or vascular dementia (Amanullah 2010).
Vitamin B6 (Pyridoxine, pyridoxal, pyridoxamine, Pyridoxal 5' phosphate (PLP) and pyridoxamine 5' phosphate (PMP), Pyridoxine 5'-phosphate (PNP))	homocysteine; neuronal activity	Vitamin B6 is a group of water-soluble compounds (vitamers). Pyridoxal 5' phosphate (PLP) and pyridoxamine 5' phosphate (PMP) are the active coenzyme forms of vitamin B6 (ODS 2014). Vitamin B6 has many important brain functions such as biosynthesis of neurotransmitters (GABA, dopamine, noradrenaline, serotonin), receptor binding, macronutrient metabolism, and gene expression. In a study looking at low plasma B6 levels predicting cognitive decline and depression in at-risk individuals, low PLP status was seen as a risk factor for cognitive decline and depression in at-risk populations (Scott 2013).
Folic Acid (Vitamin B9)	antioxidant; homocysteine; neuronal activity	Folate is a cofactor and promotes the remethylation of homocysteine an amino acid that can induce DNA strand breakage, oxidative stress, and apoptosis. Folate is required for normal development of the nervous system, playing important roles reg-

		ulating neurogenesis and programmed cell death. Folate deficiency and its resultant increase in homocysteine levels has been linked to several neurodegenerative conditions, including stroke, Alzheimer's disease, and Parkinson's disease (Mattson 2003).
Vitamin B12 (cobalamins: cyanocobalamin, hydroxo- cobalamin, methylcobalamin, hydroxo- cobalamin)	homocysteine; neuronal activity	Vitamin B12 acts as a coenzyme in metabolism of amino acids and fatty acids required for the synthesis of nucleic acids, erythrocytes, and in the maintenance of myelin (Pawlak 2014). Lower vitamin B12 status has been associated with increased rates of cognitive decline and dementia (Clarke 2007b; O'Leary 2012).
Pantothenic Acid (Vitamin B5)	energy; metabolism	Pantothenic acid (PA) is a component of coenzyme A, an essential cofactor in fatty acid oxidation, lipid elongation, and fatty acid synthesis (Kelly 2011). This may have an indirect effect in cognition.
Biotin (Vitamin H)	energy; metabolism	Biotin is also known as Vitamin H and is part of the B complex group of vitamins. They act as cofactors in carboxylase enzymes, fatty acid, and amino acid metabolism. This may have an indirect effect in cognition
Vitamin C	antioxidant	Vitamin C has antioxidant functions and is required for the synthesis of noradrenaline from dopamine. It has been reported that Vitamin C levels have been lower than controls in patients with senile dementia of Alzheimer's type (Jeandel 1989). In a longitudinal and cross-sectional study it was found that higher vitamin C levels were associated with better memory performance (Perrig 1997).
Minerals		
Calcium	neuronal activity	Calcium ions regulate a number of physiological processes including neuronal gene expression and the neuronal secretion of neurotransmitters (Delage 2014; Dolphin 2012). Supplementation with calcium together with vitamin D was found to have no significant association with incident cogni-

		tive impairment (Rossom 2012). Ozawa 2012 concluded that, in the general Japanese population, higher self-reported dietary intakes of potassium, calcium, and magnesium reduced the risk of all-cause dementia, especially Vascular Dementia (VaD). The proposed mechanism was through the reduction of of vascular risk factors
Chromium	energy production; metabolism.	Chromium is needed for energy production and has been found to promote the effect of insulin involved in metabolism and storage of protein, carbohydrates and lipids within the CNS (Anderson 1997; Institute of Medicine 2011; Ozawa 2012). Chromium is involved in metabolism of nucleic acid, which is needed to build DNA, the genetic material in cells; and promotes synthesis of cholesterol and fatty acids needed for brain function. It may lower LDL cholesterol and triglyceride levels, raise HDL cholesterol levels and reduce high blood pressure (Preuss 1997), hence may affect vascular risk factors. Insulin resistance is implicated in the pathophysiological changes associated with Alzheimer's disease, and pharmaceutical treatments that overcome insulin resistance improve memory function in subjects with mild cognitive impairment (MCI) and early Alzheimer's disease. Chromium (Cr) supplementation improves glucose disposal in patients with insulin resistance and diabetes. A double-blind RCT suggested that supplementation with Chromium picolinate can enhance cognitive inhibitory control and cerebral function in older adults at risk for neurodegeneration (Krikorian 2010). An additional study reported a positive correlation between cognitive function and serum chromium levels (Smorgon 2004).
Copper	antioxidant	Copper is a component of an antioxidant enzyme called superoxide dismutase that protects cells from damage by harmful free radicals. Copper is necessary for a healthy nerve system and taste sensitivity (Institute of Medicine 2011).

(Continued)

		Copper may promote non-amyloidogenic processing of amyloid precursor protein (APP) and thereby lower the $A\beta$ production in cell culture systems, and it increases lifetime and decreases soluble amyloid production in APP transgenic mice (Borchardt 1999). In Alzheimer patients, a decline of $A\beta$ levels in CSF is reported in adults in the treatment group (Kaden 2011).
Iodine	neuronal development and structure.	Iodine is needed for the synthesis of thyroid hormones which, in turn, are needed for the myelination of the central nervous system. Iodine is necessary for the normal development of the brain. A deficiency of this mineral during critical periods of development in gestation can lead to intellectual disability and neurodevelopmental problems (Bath 2013a). Positive association was found between maternal iodine status and child IQ at age 8 years and reading ability at age 9 years (Bath 2013b).
Iron	neuronal activity	Iron is needed for development of oligodendrocytes and numerous enzymes that synthesise neurotransmitters such as noradrenaline, serotonin, and dopamine. It is important for production of the haemoglobin in red blood cells (Delage 2014; Institute of Medicine 2011). Regression analysis showed that non-anaemic iron-deficient adolescent girls who received iron performed better on a test of verbal learning and memory than girls in the control group (Bruner 1996)
Magnesium	energy, metabolism	Magnesium is involved in hundreds of enzyme reactions, including protein synthesis. It plays a role in energy production; can improve insulin sensitivity in diabetics; helps regulate blood sugar level; and regulates neuro-muscular transmission. Higher self-reported dietary intakes of potassium, calcium, and magnesium have been reported to reduce the risk of all-cause dementia, especially VaD, in the general Japanese population (Ozawa 2012).

Manganese	metabolism	Manganese is needed to synthesise fatty acids and cholesterol, and metabolise carbohydrates and proteins. It is important for energy production. It promotes utilisation of other key nutrients like vitamin B1 (thiamine), biotin, choline, ascorbic acid, and vitamin E (Delage 2014). Manganese is needed for glucose metabolism, which helps regulate blood glucose. It is needed to make manganese superoxide dismutase (MnSOD), one of the key antioxidants that protects cells from free radical damage, and so helps maintains healthy nerves. It works synergistically with the B-complex vitamins to generate an overall feeling of well-being (Institute of Medicine 2011).
Molybdenum	metabolism	Molybdenum promotes normal cell function; and functions as a cofactor for three essential enzymes that play a vital role in carbohydrate metabolism, utilisation of iron, sulphite detoxification, and uric acid formation (Delage 2014; Institute of Medicine 2011).
Phosphorus	metabolism, neuronal structure and function.	Phosphorus is needed for metabolism of carbohydrates and fats to produce energy and is involved in the production of ATP required for growth and repair of cells and tissues; it is needed to make cell membranes. It helps the body utilise the B-complex vitamins that support proper muscle and nerve function (Delage 2014; Institute of Medicine 2011).
Potassium	nerve transmission.	Potassium is involved in regulating nerve transmissions and muscle contractions. It helps the body handle sodium and so reduces the risk of high blood pressure (Berr 2012). It has been found to lower the risk of stroke and ischaemic heart disease. Potassium is needed for synthesis of protein from amino acids (Delage 2014; Institute of Medicine 2011). Higher self-reported dietary intakes of potassium, calcium, and magnesium reduce the risk of all-cause dementia, espe-

(Continued)

		cially VaD, in the general Japanese population (Ozawa 2012).
Selenium	antioxidant.	Selenium is an important antioxidant especially in combination with vitamin E, in the central nervous system and other body tissues (Delage 2014; Mehdi 2013; Rahman 2007). Low selenium levels were found to be associated with poorer cognitive function (Berr 2012; Smorgon 2004). Supplementation with selenium has been associated with improved overall health, reducing oxidative stress and reducing the risk of dementia (Mehdi 2013).
Sodium	neuronal activity	Sodium is essential for regulating muscle contractions, nerve transmissions essential for normal CNS physiological mechanisms and homeostasis (Delage 2014; Institute of Medicine 2011).
Zinc	antioxidant neuronal activity	Zinc is a constituent of the antioxidant enzyme superoxide dismutase that helps reduce the harm from free radicals. Zinc regulates cell division and synthesis of genetic cell DNA. It is essential for reproduction, repair, and normal growth within the CNS (Delage 2014). Zinc is found in high levels in the brain where it performs catalytic, structural and regulatory roles in cellular metabolism. Zinc is bound to proteins but free zinc is present in synaptic vesicles and performs a role in neurotransmission mediated by glutamate and gamma-aminobutyric acid (GABA). Short-term deficits of zinc have been shown to impair certain measures of mental and neurological function while long-term deficits of zinc, especially during gestation, result in malformation or deficits in attention, learning, memory and neuropsychological behaviour (Institute of Medicine 2011). Zinc was found to be capable of reducing post-ischaemic injury to a variety of tissues and organs through a mechanism that might involve the antagonism of copper reactivity. Although the evidence for the an-

tioxidant properties of zinc is compelling, the mechanisms are still unclear (Powell 2000).

* Only orally-administered supplements taken at any dose for at least 12 weeks are considered. Supplements that combine vitamins or minerals are eligible as well

Footnotes

 $^{A}\beta$: Amyloid β Protein

AD: Alzheimers' Disease

APP: amyloid precursor protein

ATP: Adenosine triphosphate

CNS: central nervous system

Cr: Chromium

CSF: CerebroSpinal Fluid

DNA: DeoxyriboNucleic Acid

GABA: Gamma-AminoButyric Acid

HDL: high-density lipoproteins

IQ: Intelligence quotient

LDL: Low-density lipoprotein

MCI: Mild Cognitive Impairment

MnSOD: manganese superoxide dismutase

PLP: Pyridoxal 5' phosphate

PMP: pyridoxamine 5' phosphate

PNP: Pyridoxine 5'-phosphate

VaD: Vascular Dementia

VDR: Vitamin D receptor

25(OH)D: serum hydroxyvitamin D

Appendix 2. Sources searched and search strategies

Source	Search strategy	Hits retrieved
ALOIS (www.medicine.ox.ac.uk/alois) (Date of most recent search: 25 January 2018)	Basic search: VIT [Studies within ALOIS are coded VIT if the intervention is a vitamin or mineral]	Dec 2014: 254 Jul 2015: 0 Mar 2016: 2 Aug 2016: 0 Mar 2017: 3 Jan 2018: 1
MEDLINE In-process and other non-indexed citations and MEDLINE 1950-present (Ovid SP) (Date of most recent search: 25 January 2018)	2. exp *Minerals/3. exp *Dietary Supplements/	Dec 2014: 1320 Jul 2015: 53 Mar 2016: 111 Aug 2016: 103 Mar 2017: 166 Jan 2018: 120

- 8. toxiferol.ti,ab.
- 9. retinol.ti,ab.
- 10. "retinoic acid".ti,ab.
- 11. Vitamin A/
- 12. Vitamin B 12/
- 13. Vitamin D/
- 14. Vitamin E/
- 15. "beta-carotene".ti,ab.
- 16. "alpha-carotene".ti,ab.
- 17. "gamma-carotene".ti,ab.
- 18. "beta-cryptoanthin".ti,ab.
- 19. thiamine.ti,ab.
- 20. riboflavin.ti,ab.
- 21. niacin.ti,ab.
- 22. nicotinamide.ti,ab.
- 23. pantothenic.ti,ab.
- 24. pyridoxine.ti,ab.
- 25. pyridoxal.ti,ab.
- 26. pyridoxamine.ti,ab.
- 27. biotin.ti,ab.
- 28. "folic acid".ti,ab.
- 29. Folic Acid/
- 30. cyanocobalamin.ti,ab.
- 31. methylcobalamin.ti,ab.
- 32. "l-ascorbic acid".ti,ab.
- 33. "ascorbic acid".ti,ab.
- 34. ascorbate.ti,ab.
- 35. Ascorbic Acid/36. phylloquinone.ti,ab.
- 37. phytomeadione.ti,ab.
- 38. phytonadine.ti,ab.
- 39. mineral*.ti,ab.
- 40. multivitamin*.ti,ab.
- 41. "diet* supplement*".ti,ab.
- 42. calcium.ti,ab.
- 43. Calcium/
- 44. iron.ti,ab.
- 45. zinc.ti,ab.
- 46. sodium.ti,ab.
- 47. potassium.ti,ab.
- 48. phosphorus.ti,ab.
- 49. magnesium.ti,ab.
- 50. chloride.ti,ab.
- 51. sulphur.ti,ab.
- 52. mangansese.ti,ab.
- 53. cobalt.ti,ab.
- 54. selenium.ti,ab.
- 55. copper.ti,ab.

- 56. iodine.ti,ab.
- 57. fluoride.ti,ab.
- 58. or/1-57
- 59. *Aging/
- 60. Aged/
- 61. "Aged, 80 and over"/
- 62. Middle Aged/
- 63. Age Factors/
- 64. "mild cognitive impairment".ti,ab.
- 65. Mild Cognitive Impairment/
- 66. MCI.ti,ab.
- 67. AAMI.ti,ab.
- 68. ACMI.ti,ab.
- 69. ARCD.ti,ab.
- 70. CIND.ti,ab.
- 71. (nMCI or aMCI or mMCI or MCIa). ti,ab.
- 72. "old* adults".ti,ab.
- 73. elderly.ti,ab.
- 74. "old* age*".ti,ab.
- 75. "middle age*".ti,ab.
- 76. seniors.ti,ab.
- 77. "senior citizens".ti,ab.
- 78. "community dwelling".ti,ab.
- 79. pensioners.ti,ab.
- 80. "aged sample".ti,ab.
- 81. "aged population".ti,ab.
- 82. or/59-81
- 83. 58 and 82
- 84. *Cognition/
- 85. *Cognition Disorders/
- 86. Memory/
- 87. Memory Disorders/
- 88. (cognit* adj3 (func* or declin* or reduc* or impair* or improve* or deficit* or progress* or perform* or abilit*)).ti,ab.
- 89. "mental perform*".ti,ab.
- 90. memory.ti,ab.
- 91. "executive function*".ti,ab.
- 92. Executive Function/
- 93. Attention/
- 94. (speed adj2 processing).ti,ab.
- 95. "episodic memory".ti,ab.
- 96. Memory, Episodic/
- 97. or/84-96
- 98. 83 and 97
- 99. randomized controlled trial.pt.
- 100. controlled clinical trial.pt.

Embase 1974-24 January 2018 (Ovid SP) (Date of most recent search: 25 January	101. randomized.ab. 102. placebo.ab. 103. drug therapy.fs. 104. randomly.ab. 105. trial.ab. 106. groups.ab. 107. or/99-106 108. exp Animals/ not humans.sh. 109. 107 not 108 110. 98 and 109 [all results] 111. *Vitamins/ 112. *Cognition/ 113. "Aged, 80 and over"/ or Aged/ or Middle Aged/ 114. Mild Cognitive Impairment/ 115. "mild cognitive impairment".ti,ab. 116. 113 or 114 or 115 117. 111 and 112 and 116 118. 99 or 100 119. 117 and 118 [results sent directly to author team] 120. 110 not 119 [results minus those sent directly to author team. These results will be screened by the 'crowd'] 1. exp *vitamin/ 2. exp *mineral/ 3. exp diet supplementation/	Dec 2014: 1275 Jul 2015: 114 Mar 2016: 184
2018)	4. calcium/ 5. vitamin*.ti,ab. 6. mineral*.ti,ab. 7. cholecalciferol.ti,ab. 8. ergocalciferol.ti,ab. 9. toxiferol.ti,ab. 10. retinol.ti,ab. 11. retinal.ti,ab. 12. "retinoic acid".ti,ab. 13. vitamin D/ 14. vitamin B complex/ or vitamin B group/ 15. vitamin D/ 16. vitamin K epoxide reductase/ or vitamin K group/ 17. colecalciferol/ or calcitriol/ or calcitriol derivative/ 18. ascorbic acid/ 19. vitamin supplementation/ 20. "beta-carotene".ti,ab. 21. beta carotene/	Aug 2016: 94 Mar 2017: 257 Jan 2018: 250

- 22. "alpha-carotene".ti,ab.
- 23. alpha carotene/
- 24. "gamma-carotene".ti,ab.
- 25. gamma carotene/
- 26. "beta-cryptoanthin".ti,ab.
- 27. thiamine.ti,ab.
- 28. thiamine/
- 29. riboflavin.ti,ab.
- 30. riboflavin/
- 31. niacin.ti,ab.
- 32. nicotinic acid/
- 33. nicotinamide.ti,ab.
- 34. pantothenic.ti,ab.
- 35. pyridoxamine.ti,ab.
- 36. pantothenic acid/
- 37. pyridoxamine/
- 38. biotin.ti,ab.
- 39. biotin/
- 40. "folic acid".ti,ab.
- 41. folic acid/
- 42. cyanocobalamin.ti,ab.
- 43. cyanocobalamin/
- 44. methylcobalamin.ti,ab.
- 45. "l-ascorbic acid".ti,ab.
- 46. "ascorbic acid".ti,ab.
- 47. phylloquinone.ti,ab.
- 48. phytonadine.ti,ab.
- 49. phytomeadione.ti,ab.
- 50. multivitamin*.ti,ab.
- 51. "vitamin* supple*".ti,ab.
- 52. "diet* supplement*".ti,ab.
- 53. calcium.ti,ab.
- 54. iron.ti,ab.
- 55. iron/
- 56. zinc.ti,ab.
- 57. zinc/
- 58. sodium.ti,ab.
- 59. sodium/
- 60. potassium.ti,ab.
- 61. citrate potassium/ or potassium/ or clavulanate potassium/ or diclofenac potassium/
- 62. phosphorus.ti,ab.
- 63. phosphorus/
- 64. magnesium.ti,ab.
- 65. magnesium/
- 66. chloride.ti,ab.
- 67. chloride/

68. sulphur.ti,ab. 69. mangansese.ti,ab. 70. cobalt.ti,ab. 71. cobalt/ 72. selenium.ti,ab. 73. selenium/ 74. copper.ti,ab. 75. copper/ 76. iodine.ti,ab. 77. fluoride.ti,ab. 78. fluoride/ 79. or/1-78 80. aging/ 81. aged/ 82. middle aged/ 83. mild cognitive impairment/ 84. "mild cognitive impairment".ti,ab. 85. MCI.ti,ab. 86. AAMI.ti,ab. 87. ACMI.ti,ab. 88. ARCD.ti,ab. 89. CIND.ti,ab. 90. (nMCI or aMCI or mMCI or MCIa). ti,ab. 91. "middle age*".ti,ab. 92. "old* age*".ti,ab. 93. "old* adults".ti,ab. 94. "community dwelling".ti,ab. 95. "senior citizens".ti,ab. 96. seniors.ti,ab. 97. pensioners.ti,ab. 98. "aged sample".ti,ab. 99. "aged population".ti,ab. 100. or/80-99 101. exp cognition/ 102. cognition disorders/ 103. episodic memory/ or memory/ 104. memory disorder/ 105. dementia/ 106. Alzheimer disease/ 107. dement*.ti,ab. 108. alzheimer*.ti,ab. 109. cognition.ti,ab. 110. cognitive.ti,ab. 111. or/101-110 112. 79 and 100 and 111 113. randomized controlled trial/ 114. controlled clinical trial/

	115. placebo.ab. 116. (random* adj2 divide*).ti,ab. 117. (random* adj2 allocate*).ti,ab. 118. trial.ab. 119. "double-blind*".ti,ab. 120. "single blind*".ti,ab. 121. or/113-120 122. 112 and 121 123. *Cognition/ 124. exp *vitamin/ or exp *vitamin supplementation/ 125. exp *mineral/ or exp *mineral supplementation/ 126. (vitamin* or mineral*).ti. 127. 124 or 125 or 126 128. exp *aging/ 129. (elderly or "middle age*" or "old* adults" or MCI or "mild cognitive impairment").ti 130. exp *middle aged/ 131. 128 or 129 or 130 132. 123 and 127 and 131 133. 113 or 114 134. 132 and 133 135. 122 not 134	
PsycINFO 1806-January week 2 2018 (Ovid SP) (Date of most recent search: 25 January 2018)	1. exp Aging/ 2. exp Cognitive Impairment/ 3. "cognit* impair*".ti,ab. 4. MCI.ti,ab. 5. AAMI.ti,ab. 6. ACMI.ti,ab. 7. ARCD.ti,ab. 8. CIND.ti,ab. 9. (nMCI or aMCI or mMCI or MCIa).ti, ab. 10. "old* age*".ti,ab. 11. elderly.ti,ab. 12. "middle age*".ti,ab. 13. "old* adults".ti,ab. 14. seniors.ti,ab. 15. "senior citizens".ti,ab. 16. "community dwelling".ti,ab. 17. pensioners.ti,ab. 18. or/1-17 19. exp Cognition/ 20. exp Dementia/ 21. 19 or 20 22. randomi?ed.ti.	Dec 2014: 202 Jul 2015: 15 Mar 2016: 10 Aug 2016: 0 Mar 2017: 14 Jan 2018: 9

- 23. (randomly adj2 allocat*).ab.
- 24. (randomly adj2 divide*).ab.
- 25. RCT.ti,ab.
- 26. "double-blind*".ti,ab.
- 27. "single blind*".ti,ab.
- 28. "randomi?ed trial".ab.
- 29. "randomi?ed control* trial".ab.
- 30. "random allocation".ab.
- 31. "controlled clinical trial".ti,ab.
- 32. or/22-31
- 33. exp Vitamins/
- 34. exp Dietary Supplements/
- 35. vitamin*.ti,ab.
- 36. mineral*.ti,ab.
- 37. calcium.ti,ab.
- 38. Calcium/
- 39. exp Ascorbic Acid/
- 40. exp Folic Acid/
- 41. "folic acid".ti,ab.
- 42. cholecalciferol.ti,ab.
- 43. ergocalciferol.ti,ab.
- 44. toxiferol.ti,ab.
- 45. retinol.ti,ab.
- 46. retinal.ti,ab.
- 47. "retinoic acid".ti,ab.
- 48. "beta-carotene".ti,ab.
- 49. "alpha-carotene".ti,ab.
- 50. "gamma-carotene".ti,ab.
- 51. "beta-cryptoanthin".ti,ab.
- 52. thiamine.ti,ab.
- 53. riboflavin.ti.ab.
- 54. niacin.ti,ab.
- 55. nicotinamide.ti,ab.
- 56. pantothenic.ti,ab.
- 57. pyridoxine.ti,ab.
- 58. pyridoxal.ti,ab.
- 59. pyridoxamine.ti,ab.
- 60. biotin.ti,ab.
- 61. "folic acid".ti,ab.
- 62. cyanocobalamin.ti,ab.
- 63. methylcobalamin.ti,ab.
- 64. "l-ascorbic acid".ti,ab.
- 65. "ascorbic acid".ti,ab.
- 66. ascorbate.ti,ab.
- 67. phylloquinone.ti,ab.
- 68. phytomeadione.ti,ab.
- 69. phytonadine.ti,ab.
- 70. multivitamin*.ti,ab.

	71. "diet* supplement*".ti,ab. 72. iron.ti,ab. 73. zinc.ti,ab. 74. sodium.ti,ab. 75. potassium.ti,ab. 76. phosphorus.ti,ab. 77. magnesium.ti,ab. 78. chloride.ti,ab. 79. sulphur.ti,ab. 80. mangansese.ti,ab. 81. cobalt.ti,ab. 82. selenium.ti,ab. 83. copper.ti,ab. 84. iodine.ti,ab. 85. fluoride.ti,ab. 86. 18 or 21 87. or/33-85 88. 86 and 87 89. 32 and 88 90. exp *Vitamins/ 91. (vitamin* or mineral*).ti. 92. 90 or 91 93. exp *Cognition/ 94. (cognition or cognitive).ti. 95. 93 or 94 96. (elderly or "middle age*" or "old* adults" or MCI or "mild cognitive impairment").ti	
	trial).ti. 99. 97 and 98	
	100. 89 not 99	
CINAHL (EBSCOhost) (Date of most recent search: 25 January 2018)	S1 (MM "Vitamins+") S2 (MM "Minerals+") S3 (MH "Dietary Supplements") OR (MH "Dietary Supplementation") OR (MH "Dietary Carbohydrates") OR (MH "Dietary Fiber") OR (MH "Sodium, Dietary") OR (MH "Dietary Fats") OR (MH "Dietary Proteins") OR (MH "Dietary Sucrose") S4 TX vitamin* S5 TX mineral* S6 TX "diet* supple*" S7 (MH "Fatty Acids") OR (MH "Fatty Acids, Omega-6") OR (MH "Fatty Acids, Unsaturated") OR (MH "Trans Fatty	Aug 2016: 15 Mar 2017: 24

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Acids") OR (MH "Fatty Acids, Monoun-
saturated") OR (MH "Fatty Acids, Satu-
rated") OR (MH "Fatty Acids, Essential")
S8 TX "fatty acid*"
S9 (MH "Vitamin A")
S10 (MH "Vitamin B12") OR (MH "Vita-
min B Complex") OR (MH "Thiamine")
OR (MH "Riboflavin") OR (MH "Pyri-
doxine") OR (MH "Carnitine")
S11 (MH "Folic Acid")
S12 (MH "Ascorbic Acid")
S13 (MH "Vitamin D") OR (MH "Chole-
calciferol") OR (MH "Ergocalciferols")
OR (MH "Calcitriol")
S14 (MH "Vitamin E") OR (MH "Pan-
tothenic Acid")
S15 (MH "Vitamin K") OR (MH "Osteo-
calcin")
S16 TX "beta-carotene"
S17 TX "alpha-carotene"
S18 TX thiamine
S19 TX riboflavin
S20 TX niacin
S21 TX pantothenic
S22 TX nicotinamide
S23 TX pyridoxine
S24 TX pyridoxal
S25 TX biotin
S26 (MH "Calcium")
S27 TX calcium
S28 TX iron
S29 (MH "iron")
S30 (MH "Zinc")
S31 TX zinc
S32 (MH "Sodium")
S33 TX sodium
S34 (MH "Potassium")
S35 TX potassium
S36 (MH "Phosphorus")
S37 TX phosphorus
S38 (MH "Magnesium")
S39 TX magnesium
S40 (MH "Sodium Chloride, Dietary")
S41 TX chloride
S42 TX sulphur
S43 TX cobalt
S44 TX selenium
S45 TX copper
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S46 TX iodine
S47 TX flouride
S48 S1 OR S2 OR S3 OR S4 OR S5 OR
S6 OR S7 OR S8 OR S9 OR S10 OR S11
OR S12 OR S13 OR S14 OR S15 OR S16
OR S17 OR S18 OR S19 OR S20 OR S21
OR S22 OR S23 OR S24 OR S25 OR S26
OR S27 OR S28 OR S29 OR S30 OR S31
OR S32 OR S33 OR S34 OR S35 OR S36
OR S37 OR S38 OR S39 OR S40 OR S41
OR S42 OR S43 OR S44 OR S45 OR S46
OR S47
S49 (MH "Aging")
S50 (MH "Aged") OR (MH "Aged, 80 and
Over")
S51 (MH "Middle Age")
S52 TX "Mild Cognitive Impairment"
S53 TX MCI OR AAMI OR ACMI OR
ARCD OR CIND
S54 TX nMCI OR aMCI OR mMCI OR
MCIa
S55 TX elderly
S56 TX "old* adults"
S57 TX "old* age*"
S58 TX pensioners
S59 TX "community dwelling"
S60 TX seniors
S61 TX "senior citizen*"
S62 TX "age* sample"
S63 TX "age* population"
S64 S49 OR S50 OR S51 OR S52 OR S53
OR S54 OR S55 OR S56 OR S57 OR S58
OR S59 OR S60 OR S61 OR S62 OR S63
S65 S48 AND S64
S66 (MH "Cognition") OR (MH "Cogni-
tion Disorders") OR (MH "Delirium, De-
mentia, Amnestic, Cognitive Disorders")
S67 TX cognition
S68 TX memory
S69 (MH "Memory") OR (MH "Mem-
ory Disorders") OR (MH "Memory, Short
Term")
S70 TX "executive function"
S71 TX "cognitive* declin*"
S72 TX "cognitive* improv*"
S73 TX "cognitive deficit*"
S74 TX "mental perform*"
S75 TX dementia
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	S76 TX alzheimer* S77 (MH "Dementia+") S78 S66 OR S67 OR S68 OR S69 OR S70 OR S71 OR S72 OR S73 OR S74 OR S75 OR S76 OR S77 S79 S65 AND S78 S80 (MH "Randomized Controlled Trials") S81 AB randomly S82 AB placebo S83 AB groups S84 AB RCT S85 TX "double blind*" S86 TX "single blind*" S87 TX "controlled clinical trial" S88 TI randomised S89 TI randomized S90 S80 OR S81 OR S82 OR S83 OR S84 OR S85 OR S86 OR S87 OR S88 S91 S79 AND S90	
ISI Web of Science [includes: Web of Science (1945-present); BIOSIS Previews (1926-present); MEDLINE (1950-present); Journal Citation Reports]; BIOSIS Previews (Date of most recent search: 25 January 2018)	("mild cognitive impairment" OR elderly OR "age* subjects" OR "old* adult*" OR "middle age*" OR MCI) AND TOPIC: ("randomly allocated" OR "random allocation" OR randomised OR randomized OR RCT OR "controlled trial" OR "double blind" OR "single blind") AND TOPIC: (vitamin* OR mineral* OR "diet* suppl*" OR "ascorbic acid" OR "folic acid" OR iron OR calcium OR sodium OR zinc OR potassium OR magnesium OR cobalt OR copper OR iodine) AND TOPIC: (cognition OR dementia OR memory OR "executive function" OR alzheimer*) Timespan: All years. Search language=Auto	Jul 2015: 34 Mar 2016: 100 Aug 2016: 43 Mar 2017: 67
LILACS (BIREME) (Date of most recent search: 25 January 2018)	cognition OR "mild cognitive impairment" OR elderly OR "aged subjects" OR "older adults" OR "middle aged" [Words] and randomly OR randomised OR randomized OR RCT OR "controlled trial" [Words] and vitamin OR vitamins OR mineral OR minerals OR "fatty acid" OR "folic acid" [Words]	Jul 2015: 0 Mar 2016: 0 Aug 2016: 1 Mar 2017: 0
CENTRAL (via CRSO) (Date of most recent search: 25 January 2018)	#1 MeSH descriptor: [Aged, 80 and over] explode all trees	Dec 2014: 395 Jul 2015: 10

#2 MeSH descriptor: [Aged] explode all Mar 2016: 50 Aug 2016: 48 trees #3 MeSH descriptor: [Middle Aged] ex- Mar 2017: 90 plode all trees Jan 2018: 54 #4 MeSH descriptor: [Mild Cognitive Impairment] explode all trees #5 "cognit* impair*" or MCI #6 elderly #7 "old* adults" #8 "old* age*" #9 "old* sample" #10 senior citizens #11 pensioners #12 seniors #13 #1 or #2 or #3 or #4 or #5 or #6 or # 7 or #8 or #9 or #10 or #11 or #12 #14 MeSH descriptor: [Cognition] explode all trees #15 MeSH descriptor: [Dementia] explode all trees #16 cognit* #17 memory #18 "executive function*" #19 processing #20 "mental perform*" #21 dement* #22 alzheimer* #23 #14 or #15 or #16 or #17 or #18 or # 19 or #20 or #21 or #22 #24 MeSH descriptor: [Vitamins] explode all trees #25 MeSH descriptor: [Minerals] explode all trees #26 vitamin* #27 mineral* #28 "ascorbic acid" #29 "folic acid" #30 MeSH descriptor: [Fatty Acids] explode all trees #31 zinc or iron or calcium or sodium or potassium or magnesium or cobalt or copper or selenium or iodine or flouride or chloride #32 "beta-carotene" #33 "alpha-carotene" #34 thiamine #35 riboflavin #36 niacin

	#37 biotin #38 pantothenic #39 nicotinamide #40 pyridoxal #41 "dier* suppl*" #42 #26 or #27 or #28 or #29 or #30 or # 31 or #32 or #33 or #34 or #35 or #36 or #37 or #38 or #39 or #40 or #41 #43 #13 and #23 and #42	
Clinicaltrials.gov (www.clinicaltrials.gov) (Date of most recent search: 25 January 2018)	In Intervention studies: [intervention] vitamin* OR mineral* OR "diet* suppl*" OR "ascorbic acid" OR "folic acid" OR iron OR calcium OR sodium OR zinc OR potassium OR magnesium OR cobalt OR copper OR iodine AND [condition]: cognition OR "mild cognitive impairment" OR elderly OR "aged subjects" OR "older adults" OR "middle aged" Trial Status: all	Jul 2015: 0 Mar 2016: 2 Aug 2016: 0
ICTRP Search Portal (http://apps.who.int/trialsearch) (includes: Australian New Zealand Clinical Trials Registry; ClinicalTrilas.gov; ISRCTN; Chinese Clinical Trial Registry; Clinical Trials Registry - India; Clinical Research Information Service - Republic of Korea; German Clinical Trials Register; Iranian Registry of Clinical Trials; Japan Primary Registries Network; Pan African Clinical Trial Registry; Sri Lanka Clinical Trials Registry; The Netherlands National Trial Register) (Date of most recent search: 25 January 2018)	tamin* OR mineral* OR "diet* suppl*" OR "ascorbic acid" OR "folic acid" OR iron OR calcium OR sodium OR zinc OR potassium OR magnesium OR cobalt OR	Jul 2015: 0 Mar 2016: 2 Aug 2016: 0
TOTAL before de-duplication		Dec 2014: 5071 Jul 2015: 233 Mar 2016: 496 Aug 2016: 304 Mar 2017: 631 Jan 2018: 522 TOTAL: 7257
TOTAL after de-duplication		Dec 2014: 3451 Jul 2015: 200 Mar 2016: 399 Aug 2016: 208 Mar 2017: 522 Jan 2018: 431

	TOTAL: 5211
TOTAL after first assessment by the Crowd and CDCIG information specialists	Dec 2014: 448 Jul 2015: 81 Mar 2016: 70 Aug 2016: 45 Mar 2017: 43 Jan 2018: 38 TOTAL: 725

Appendix 3. Definitions of design, patient and intervention characteristics as applied in the stratified analyses exploring between trial variations in intervention effect

Item	Definition
Bias-related characteristics*	
Concealment of allocation (avoiding selection bias)	The guidance from the <i>Cochrane Handbook for Systematic Reviews of Interventions</i> (Higgins 2011) will be used to judge bias related to sequence generation and concealment of allocation using the two Cochrane 'Risk of bias' items. From these, the statistician will derive a single variable to be used in the stratified analysis: allocation concealment will be judged at low risk of bias if the investigators responsible for patient selection were unable to suspect before allocation which treatment was next. Concealment will downgraded to high risk of bias if there is evidence of inadequate sequence generation
Blinding of patients and personnel (avoiding performance bias)	Low risk of bias will be judged if: - a credible sham procedure was used; or if a placebo supplement or pill was used that was reported to be identical in appearance to the experimental intervention and the specific outcome or group of outcomes is/are likely to be influenced by lack of blinding - blinding is absent or suboptimal and the specific outcome, such as mortality, is not likely to be influenced by lack of blinding
Blinding of outcome assessment (avoiding detection bias)	For self-reported/partner-reported outcomes: Low risk of bias will be judged if: - self-reported outcomes were assessed AND blinding of patients was considered adequate AND there was no information to suggest that there was an investigator involved during the process of outcome assessment; OR if blinding of investigators performing the outcome assessment was reported AND an attempt to blind patients was reported For other outcomes: Outcome assessment was considered to be blinded if the outcome

	assessment was reported to be blinded
Statistical analyses (avoiding attrition bias)	For continuous outcomes Low risk of bias will be judged if: - at least 90% of the patients randomised were analysed AND the difference in percentage of participants not analysed was 5% or lower across trial arms - for trials using imputations to handle missing data: the percentage of participants with missing data did not exceed 20% AND the difference in percentage of participants with imputed data was 10% or lower across trial arms AND applied imputation methods were judged to be appropriate For binary outcomes of rare events Low risk of bias will be judged if: - the event rate is low (e.g. incidence of dementia) AND at least 95% of the patients randomised were analysed AND there is no evidence of differential reasons for missing data that may alter the estimate AND the rate of missing data does not exceed the expected event rates For binary outcomes of non-rare events Low risk of bias will be judged if: - at least 90% of the patients randomised were analysed AND the difference in percentage of participants not analysed was 5% or lower across trial arms AND there is no evidence of differential reasons for missing data that may alter the estimate AND the rate of missing data does not exceed the expected event rates
Trial Size	A large trial will be defined by a sample size calculation for the primary outcome
Follow-up duration	For the cognitive outcomes, we will group studies according to these follow-up cut-offs to describe immediate results (up to 12 weeks), short-term (up to 1 year), medium-term (1 to 2 years) and longer-term results (more than 2 years). For the secondary outcome all-cause dementia, only outcome data at 1 year of follow-up or longer will be considered and therefore the grouping will include short-term (1 year), medium-term (up to 2 years) and longer-term results (more than 2 years)
Treatment-related characteristics	
Treatment duration	The minimum treatment duration of 3 months is considered short-term, 3 to 12 months as medium-term, and 12 months for long-term
Dose of treatment	Treatment will be analysed as high dose versus low dose according to previously reported cut-offs

Mechanisms of action of the supplements** Participant-related characteristics	Supplements postulated to share a main mechanism of action in preventing development of dementia, including: • Antioxidant properties - affecting superoxide dismutase (vitamin A, C, D, E, selenium) • Regulation/lowering levels of homocysteine: vitamins B12, folate and B6
Cognition-related criteria	No risk of deficiency versus at risk of deficiency for the type of vitamin and minerals investigated (e.g. presence of malabsorptive diseases, malnutrition, comorbidities or concomitant medications, and ethnicity (Vitamin D))

^{*} The descriptions depicted in this Table are in addition to the guidance provided by Cochrane (Higgins 2011).

Appendix 4. Composition of Centrum Silver ® used in Grodstein 2013

Vitamin A (IU)	5000*
Vitamin C (mg)	60
Vitamin D (IU)	400
Vitamin E (IU)	45
Vitamin K (μg)	10
Thiamin (mg)	1.5
Riboflavin (mg)	1.7
Niacin (mg)	20
Vitamin B6 (mg)	3
Folic Acid (µg)	400
Vitamin B12 (μg)	25
Biotin (μg)	30
Pantothenic Acid (mg)	10

^{**} Knowledge of possible mechanisms of actions is evolving, and we will consider other possible subgroups for data analysis as new information arises during the development of the review.

(Continued)

Calcium (mg)	200
Iron (mg)	4
Phosphorus (mg)	48
Iodine (μg)	150
Magnesium (mg)	100
Zinc (mg)	15
Selenium (µg)	20
Copper (mg)	2
Manganese (mg)	3.5
Chromium (µg)	130
Molybdenum (μg)	160
Chloride (mg)	72.6
Potassium (mg)	80
Boron (μg)	150
Nickel (μg)	5
Vanadium (µg)	10
Silicon (mg)	2

CONTRIBUTIONS OF AUTHORS

 $Completion \ of the \ protocol: \ Rutjes \ AWS, \ Al-Assaf \ AS, \ Denton \ DA, \ Abraham \ RP, \ Chong \ LY, \ Anderson \ J, \ Malik \ MA, \ Tabet \ N$

Completion of the searches: Noel-Storr A

Screening of references: Di Nisio M, Al-Assaf AS, Denton DA, Abraham RP, Chong LY, Anderson J, Malik MA, Rutjes AWS, Tabet N, Martinez G, McCleery J

Acquisition of data: Di Nisio M, Al-Assaf AS, Vernooij RWM, Rutjes AWS, Martinez G, McCleery J

Risk of bias and GRADE assessments: Di Nisio M, Vernooij RWM, Rutjes AWS, Martinez G, McCleery J

Statistical analysis: McCleery J, Martinez G

Overall interpretation of data: McCleery J

Manuscript preparation: McCleery J, Rutjes AWS

DECLARATIONS OF INTEREST

Rutjes AWS: Dr. Rutjes declares partial funding by a grant for the project 'OPERAM: OPtimising therapy to prevent Avoidable hospital admissions in the Multi-morbid elderly' supported by the European Union's Horizon 2020 research and innovation programme under the grant agreement No 6342388, and by the Swiss State Secretariat for Education, Research and Innovation (SERI) under contract number 15.0137.

Denton DA: none known

Di Nisio M: Di Nisio declares consultancy fees from Daiichi Sankyo, Bayer, and Aspen and participation to advisory boards for Daiichi Sankyo, Bayer, Pfizer outside the submitted work. M Di Nisio declares partial funding by a grant for the project 'OPERAM: OPtimising therapy to prevent Avoidable hospital admissions in the Multi-morbid elderly' supported by the European Union's Horizon 2020 research and innovation programme under the grant agreement No 6342388, and by the Swiss State Secretariat for Education, Research and Innovation (SERI) under contract number 15.0137

Chong LY: none known

Abraham RP: none known

Al-Assaf AS: none known

Anderson JL: none known

Malik MA: none known

Vernooij RWM: none known

Martinez G: none known

Tabet N: N Tabet reports I receive honorarium and consultation fees from the pharmaceutical industry

McCleery J: none known

SOURCES OF SUPPORT

Internal sources

• No sources of support supplied

External sources

• NIHR, UK.

This review was supported by the National Institute for Health Research, via a Cochrane Programme Grant to the Cochrane Dementia and Cognitive Improvement group. The views and opinions expressed therein are those of the authors and do not necessarily reflect those of the Systematic Reviews Programme, NIHR, NHS or the Department of Health

• SERI & Horizon 2020, Other.

The authors AR and MdN are partially funded by a grant for the project 'OPERAM: OPtimising therapy to prevent Avoidable hospital admissions in the Multi-morbid elderly' supported by the European Union's Horizon 2020 research and innovation programme under the grant agreement No 6342388, and by the Swiss State Secretariat for Education, Research and Innovation (SERI) under contract number 15.0137. The opinions expressed and arguments employed herein are those of the authors and do not necessarily reflect the official views of the EC and the Swiss government.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

There are a number of significant differences between this review and its protocol.

Published protocols anticipated the production of two reviews with distinct populations: cognitively healthy people in midlife (40 years to 65 years; Denton 2015) and in late life (> 65 years). However, it became apparent that many studies straddled this arbitrary age boundary so that it was not possible logically to assign them to one review rather than the other. Therefore, it was decided to produce a single review covering participants in mid- and late-life.

The protocols specified that participants should have been cognitively assessed at baseline. Again, this criterion would have led to exclusion of a large amount of potentially informative data and was dropped in favour of a more pragmatic approach. If a study stated that participants were cognitively healthy or if they were a large population sample with no cognitive inclusion criteria, then they were assumed to be eligible.

The protocol specified only three time points for grouping outcomes: 12 weeks to 1 year, 1 to 2 years and > 2 years. However, we considered that the very wide variation in durations of follow-up justified further subdivisions at 5 and 10 years. Particularly in the middle-aged and 'young' elderly, trial durations as short as 2 years are unlikely to be able to detect effects on the trajectory of age-related cognitive decline.

Appendix 3 described a stringent method for assessing blinding. In practice, we adopted a more flexible approach and for multicentre studies with generally rigorous methods, we accepted trial authors' statements about blinding of participants and investigators in the absence of further detail.