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Interventions to improve hearing aid use in adult auditory rehabilitation (Review)

Barker F, Mackenzie E, Elliott L, Jones S, de Lusignan S

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Interventions to improve hearing aid use in adult auditory rehabilitation

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ABSTRACT

Background

Acquired adult-onset hearing loss is a common long-term condition for which the most common intervention is hearing aid fitting. However, up to 40% of people fitted with a hearing aid either fail to use it or may not gain optimal benefit from it. This is an update of a review first published in *The Cochrane Library* in 2014.

Objectives

To assess the long-term effectiveness of interventions to promote the use of hearing aids in adults with acquired hearing loss fitted with at least one hearing aid.

Search methods

The Cochrane ENT Information Specialist searched the Cochrane ENT Trials Register; Central Register of Controlled Trials (CENTRAL 2016, Issue 5); PubMed; EMBASE; CINAHL; Web of Science; ClinicalTrials.gov; ICTRP and additional sources for published and unpublished trials. The date of the search was 13 June 2016.

Selection criteria

We included randomised controlled trials (RCTs) of interventions designed to improve or promote hearing aid use in adults with acquired hearing loss compared with usual care or another intervention. We excluded interventions that compared hearing aid technology. We classified interventions according to the 'chronic care model' (CCM). The primary outcomes were hearing aid use (measured as adherence or daily hours of use) and adverse effects (inappropriate advice or clinical practice, or patient complaints). Secondary patient-reported outcomes included quality of life, hearing handicap, hearing aid benefit and communication. Outcomes were measured over the short (≤ 12 weeks), medium (> 12 to < 52 weeks) and long term (one year plus).

Data collection and analysis

We used the standard methodological procedures expected by Cochrane.

Main results

We included 37 studies involving a total of 4129 participants. Risk of bias across the included studies was variable. We judged the GRADE quality of evidence to be very low or low for the primary outcomes where data were available.

The majority of participants were over 65 years of age with mild to moderate adult-onset hearing loss. There was a mix of new and experienced hearing aid users. Six of the studies (287 participants) assessed long-term outcomes.

All 37 studies tested interventions that could be classified using the CCM as self-management support (ways to help someone to manage their hearing loss and hearing aid(s) better by giving information, practice and experience at listening/communicating or by asking people to practise tasks at home) and/or delivery system design interventions (just changing how the service was delivered).

Self-management support interventions

We found no studies that investigated the effect of these interventions on adherence, adverse effects or hearing aid benefit. Two studies reported daily hours of hearing aid use but we were unable to combine these in a meta-analysis. There was no evidence of a statistically significant effect on quality of life over the medium term. Self-management support reduced short- to medium-term hearing handicap (two studies, 87 participants; mean difference (MD) -12.80, 95% confidence interval (CI) -23.11 to -2.48 (0 to 100 scale)) and increased the use of verbal communication strategies in the short to medium term (one study, 52 participants; MD 0.72, 95% CI 0.21 to 1.23 (0 to 5 scale)). The clinical significance of these statistical findings is uncertain. It is likely that the outcomes were clinically significant for some, but not all, participants. Our confidence in the quality of this evidence was very low. No self-management support studies reported long-term outcomes.

Delivery system design interventions

These interventions did not significantly affect adherence or daily hours of hearing aid use in the short to medium term, or adverse effects in the long term. We found no studies that investigated the effect of these interventions on quality of life. There was no evidence of a statistically or clinically significant effect on hearing handicap, hearing aid benefit or the use of verbal communication strategies in the short to medium term. Our confidence in the quality of this evidence was low or very low. Long-term outcome measurement was rare.

Combined self-management support/delivery system design interventions

One combined intervention showed evidence of a statistically significant effect on adherence in the short term (one study, 167 participants, risk ratio (RR) 1.06, 95% CI 1.00 to 1.12). However, there was no evidence of a statistically or clinically significant effect on daily hours of hearing aid use over the long term, or the short to medium term. No studies of this type investigated adverse effects. There was no evidence of an effect on quality of life over the long term, or short to medium term. These combined interventions reduced hearing handicap in the short to medium term (15 studies, 728 participants; standardised mean difference (SMD) -0.26, 95% CI -0.48 to -0.04). This represents a small-moderate effect size but there is no evidence of a statistically significant effect over the long term. There was evidence of a statistically, but not clinically, significant effect on long-term hearing aid benefit (two studies, 69 participants, MD 0.30, 95% CI 0.02 to 0.58 (1 to 5 scale)), but no evidence of an effect over the short to medium term. There was evidence of a statistically, but not clinically, significant effect on the use of verbal communication strategies in the short term (four studies, 223 participants, MD 0.45, 95% CI 0.15 to 0.74 (0 to 5 scale)), but not the long term. Our confidence in the quality of this evidence was low or very low.

We found no studies that assessed the effect of other CCM interventions (decision support, the clinical information system, community resources or health system changes).

Authors' conclusions

There is some low to very low quality evidence to support the use of self-management support and complex interventions combining self-management support and delivery system design in adult auditory rehabilitation. However, effect sizes are small. The range of interventions that have been tested is relatively limited. Future research should prioritise: long-term outcome assessment; development of a core outcome set for adult auditory rehabilitation; and study designs and outcome measures that are powered to detect incremental effects of rehabilitative healthcare system changes.

PLAIN LANGUAGE SUMMARY

Interventions to improve hearing aid use in adult auditory rehabilitation

Review question

We wanted to know if any interventions help people to wear their hearing aids more. We measured effects over the short term (less than 12 weeks), medium term (from 12 to 52 weeks) and long term (one year plus). This is an update of a review first published in *The Cochrane Library* in 2014.

Background

Hearing loss is very common. People who get hearing loss as adults are often offered a hearing aid(s). However, up to 40% of people fitted with a hearing aid choose not to use it.

Study characteristics

The evidence is up to date as of June 2016. We found 37 studies involving a total of 4129 people. Most of the people in the studies were aged over 65. There was a mix of new and experienced hearing aid users. Seven studies funded by the United States Veterans Association dominate the evidence. The 1297 people in these studies were serving in the military or military veterans. All but two of the other studies included fewer than 100 people in each study.

Results

Thirty-three of the 37 studies looked at ways to help someone to manage their hearing loss and hearing aid(s) better by giving information, practice and experience at listening/communicating or by asking people to practise tasks at home. These are forms of self-management support. Most of these studies also changed how the self-management support was provided, for example by changing the number of appointment sessions or using telephone or email follow-up.

Six studies looked at the effect of just changing how the service was delivered. No studies looked at the effect of using guidelines or standards, computerised medical record systems, community resources or changing the health system.

We found no evidence that the interventions helped people to wear their hearing aids for more hours per day over the short, medium or long term. One study that used interactive videos to give information after hearing aid fitting encouraged more people to wear their hearing aids.

We found no evidence of adverse effects of any of the interventions, but it was rare for studies to look for adverse effects.

Giving self-management support meant that people reported less hearing handicap and improved verbal communication over the short term. When this was combined with changing how the support was delivered people also reported slightly more hearing aid benefit over the long term.

Only six studies (287 people) looked at how people were doing after a year or more.

Conclusions

Complex interventions that deliver self-management support in different ways improve some outcomes for some people with hearing loss who use hearing aids. We found no interventions that increased self-reported daily hours of hearing aid use. Few studies measured how many people use hearing aids compared to how many are fitted (adherence). Many things that might increase daily hours of hearing aid use or encourage more people to wear the hearing aids they have been fitted with have not been tested. It was difficult to combine data across different studies because many outcome measures were used and results were not always fully reported. In future it would be helpful if researchers:

- used existing guidelines for presenting their results;
- agreed a set of outcome measures for use in this type of study; and
- focused on long-term outcomes where people are followed up for at least a year.

Quality of the evidence

We judged the evidence to be of very low or low quality. There was risk of bias in the way many of the studies were carried out or reported. The largest studies included only military veterans. We do not know whether studies would find the same results in more mixed populations. Most of the other studies had small sample sizes. Very few studies measured long-term outcomes.

SUMMARY OF FINDINGS FOR THE MAIN COMPARISON *[Explanation]*

Self-management support interventions for adults with hearing loss who use hearing aids						
Patient or population: adults with hearing loss who use hearing aids Settings: outpatient clinic Intervention: self-management support interventions						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	Control	Self-management support interventions				
Adherence	No studies identified					
Daily hours of hearing aid use	Two studies reported daily hours of hearing aid use but we were unable to combine these in a meta-analysis					
Adverse effects	No studies identified					
Quality of life Validated self-report measures. WHODAS 2.0 scale from: 0 to 100 Follow-up: 0 to 12 months	The mean quality of life in the intervention group was 9.1 lower (21.33 lower to 3.13 higher) than in the control group (on this generic health-related quality of life scale (WHODAS 2.0) a lower score indicates better quality of life)		-	35 (1 study)	⊕○○○ very low ¹	The minimal important difference on this scale has not been established for hearing health care
Self-reported hearing handicap Validated self-report measure: HHIE (Ventry 1982) scale from 0 to 100 Follow-up: 0 to 12 months	The mean self-reported hearing handicap in the intervention groups was 12.80 lower (23.11 lower to 2.48 lower) than in the control groups (lower score indicates less handicap)		-	87 (2 studies)	⊕○○○ very low ²	The minimal important difference on this scale is reported to be 18.7 for face-to face administration and 36 for pencil and paper (Weinstein 1986).

Hearing aid benefit	No studies identified			
Communication Validated self-report measure: verbal sub-scale of the CPHI (Demorest 1987) scale from 0 to 5 Follow-up: 0 to 12 months	The mean reported use of verbal communication strategy in the intervention group was 0.72 higher (0.21 higher to 1.23 higher) than in the control group (higher score indicates increased use of verbal communication strategy)	- 52 (1 study)	⊕○○○ very low ³	The minimal important difference for this sub-scale of the CPHI is 0.93 at the 0.05 level (Demorest 1988).

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (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; **CPHI:** Communication Profile for the Hearing Impaired; **RR:** risk ratio; **WHODAS 2.0:** World Health Organization Disability Assessment Schedule 2.0

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

¹Downgraded due to very serious concerns regarding limitations in study design (risk of bias), indirectness (participants were military veterans and only short- to medium-term outcomes were available) and serious concerns regarding imprecision (single study with small sample size).

²Downgraded due to very serious concerns regarding limitations in study design (risk of bias) and serious concerns due to indirectness (only short- to medium-term outcomes available) and imprecision (two small studies with a high risk of skewed data).

³Downgraded due to very serious concerns regarding limitations in study design (risk of bias) and serious concerns due to indirectness (only short- to medium-term outcomes available) and imprecision (single study with small sample size).

BACKGROUND

This is an update of a review first published in *The Cochrane Library* in 2014.

Description of the condition

Adult acquired hearing loss is a common long-term condition, which in the majority of cases is not remediable by surgical or medical intervention. It ranks 15th amongst the leading causes of global burden of disease and is the second leading cause of 'years lived with a disability' (WHO 2012). The prevalence of hearing loss increases with age, which has serious implications in a global population in which the proportion of elderly people is rising at unprecedented rates according to the World Health Organization (WHO 2011). The standard intervention for hearing loss, at least in the developed world, usually involves the provision of monaural or binaural hearing aids within an audiology clinic (Cox 2014). Despite the evidence of the negative consequences of hearing loss (Brooks 2001; Hallberg 1993; Lin 2011; Saito 2010), and the benefits of hearing aids (Bainbridge 2014; Chisolm 2007; Mulrow 1992; National Council on Aging 2000; Swan 2012), uptake of fitting is relatively low, even in countries where the provision of hearing aids is free at the point of use. In addition, results from studies on use and non-use of hearing aids support the finding that a proportion of those being prescribed a hearing aid do not use it. Estimates of non-use vary from 5% to 40% (Gimsing 2008; Lupsakko 2005; Smeeth 2002; Sorri 1984; Vuorialho 2006), and this is supported by commercial survey data from hearing aid dispensers (Hougaard 2011; Kochkin 2009). Some studies have highlighted poor sound quality or lack of perceived benefit as one of the reasons for non-use (Brooks 1985; Lupsakko 2005; Smeeth 2002), and it is likely that developments in sound processing technology have had an effect over time, such as the move from analogue to digital sound processing. The more recent studies tend to show higher levels of use but there is still room for improvement. Recent evidence suggests that increased cost does not necessarily improve outcome over and above that gained from more cost-effective options (Cox 2014). In addition, there is a reliable placebo effect when assessing different hearing aid technologies, which may have an impact on the results of unblinded studies (Dawes 2013). There are no data on rates of use in developing countries where access to hearing aid technology presents more of a challenge, although reasons for non-use are starting to be investigated in less well-resourced populations (Borg 2015).

Description of the intervention

This review considered any healthcare interventions aimed at improving or promoting the use of hearing aids in the context of acquired adult hearing loss. To provide a structure for this analysis we chose to classify interventions based on the chronic care model

(CCM) (Bodenheimer 2002). This is a framework used to develop and describe initiatives in the care of long-term conditions. Adult acquired hearing loss fits the World Health Organization definition of a long-term condition in that it is a health problem that requires ongoing management over a period of years or decades (WHO 2002). We chose the CCM because it is widely cited, has been used in a variety of healthcare settings and its implementation has been associated with improved outcomes (DH 2007; NHS 2006; Tsai 2005). It has also been used as a framework in previous reviews looking at the effects of interventions in the context of long-term conditions (Kreindler 2009; Tsai 2005).

We therefore hoped that this review would provide information on interventions and outcomes in the context of hearing loss as a long-term condition.

We classified potential interventions according to the six elements of the CCM as follows:

1. Self-management support interventions

Self-management support is at the heart of the CCM and other frameworks used in the context of long-term conditions. For chronic conditions such as hearing loss, patients themselves take on the primary responsibility for managing their condition. These are interventions that seek to empower and prepare patients to manage their own health and health care. They emphasise the patient's central role in managing their health. Self-management support involves collaborating with patients and their families to help them develop the skills and confidence they need to do this effectively. In their review of self-management approaches for people with long-term conditions, Barlow and colleagues state that "self-management refers to the individual's ability to manage the symptoms, treatment, physical and psychosocial consequences and lifestyle changes inherent in living with a chronic condition" (Barlow 2002). Self-management is a complex task.

The provision of self-management support might therefore include:

- self-management assessment - assessment of the impact of hearing loss, the difficulties it is causing and facilitators and barriers to potential solutions;
- patient education;
- patient activation - interventions that involve practice of the behaviour changes needed to develop practical, symptom and psychosocial management skills;
- self-management resources and tools - battery replacement services, provision of additional equipment to improve hearing aid benefit;
- collaborative decision-making (Tsai 2005).

These processes align to the Assess, Advise, Assist and Agree components of the 5As model of health behaviour change (Whitlock 2002). These have been applied previously in the context of long-term condition self-management (Glasgow 2003).

2. Delivery system design interventions

These interventions involve the introduction of systems to assure the delivery of efficient, effective care and self-management support. Kreindler 2009 and Tsai 2005 describe how this includes interventions that:

- reshape healthcare provider roles - for example, introducing the role of case manager or defining roles within a multi-disciplinary team;
- reorganise the scheduling or organisation of care - changes in care delivery, the provision of follow-up or planned visits, visit system change.

Delivery system design interventions involve changes in the mode (for example, group versus individual), format (face-to-face, on-line, booklet etc.), timing or follow-up pattern and location of delivery of self-management support rather than the content of the support itself. This category would include interventions such as group counselling and group rehabilitation, providing the same content is delivered in the same way to the intervention and control group. It will also include interventions where changes have been made to the post-fitting follow-up process in terms of timing, quantity, location, mode or format of delivery. The 'arrange follow-up' component of the 5As model would fit within this element.

In reality it is likely that many interventions will contain an element of self-management support and delivery system design because in order to provide self-management support some changes in delivery system design are likely to be needed (Tsai 2005).

3. Decision support interventions

Decision support interventions promote clinical care that is consistent with scientific evidence and patient preferences. They embed evidence-based guidelines into daily clinical practice and provide mechanisms to share evidence-based guidelines and information with patients to encourage their participation. They may involve the use of proven provider education methods or seek to integrate specialist expertise and primary care.

4. Clinical information system interventions

Interventions involving clinical information systems, generally computerised medical records systems, aim to organise patient and population data to facilitate care, provide timely reminders for providers and patients, identify relevant subpopulations for proactive care, facilitate individual care planning, share information with patients and providers to co-ordinate care, and monitor performance of the practice team and care system as a whole. For example, in audiology this might include the introduction of electronic patient records that facilitate the development of individual management plans or identify patients in need of routine review or follow-up.

5. Community interventions

Interventions falling into this category include those that mobilise community resources to meet the needs of patients, encourage patients to participate in community-based programmes or where partnerships have been formed with community organisations to support and develop interventions that fill gaps in services or advocate for policies to improve patient care. In audiology this might include partnerships with local deaf clubs or community volunteers who visit patients in their own homes.

6. Health system interventions

Health system interventions seek to create a culture, organisation and mechanisms to promote safe, high-quality care or visibly support improvement at all levels of the organisation, beginning with the senior leader. They may involve the introduction of policies that encourage open and systematic handling of quality problems or provide incentives based on quality of care. Health system interventions may also seek to develop agreements that facilitate care co-ordination within and across organisations. Examples from hearing health care would include the introduction of the Improving Quality in Physiological Diagnostic Services (IQIPS) programme. See Column 1, Table 1.

We recognise that within each of these elements there will be clinical diversity in the type of intervention delivered and we therefore planned to investigate this diversity with subgroup analyses where appropriate. However, we felt it would of interest to policy-makers to know the relative effects of different interventions grouped by element so that they can make an informed judgement about whether it is more cost-effective to make changes in intervention content (e.g. self-management support), how that content is delivered (delivery system design) or in how delivery is supported (decision support, clinical information system).

How the intervention might work

1. Self-management support interventions

These interventions act directly to promote behaviour change on the part of the patient. The behaviour change of primary interest in this review is increased hearing aid use. This might be achieved through:

- improving knowledge (advice);
- practising new skills - practical, symptom management and psychosocial management skills (assist through activation/engagement);
- providing self-management resources and tools (assist through resource provision);
- collaborating in decision-making (agree).

We recognised that these subtypes of self-management support may have an impact on behaviour to different extents and we ex-

plored this further with subgroup analyses (see [Subgroup analysis and investigation of heterogeneity](#)).

2. Delivery system design interventions

Delivery system design interventions work by making changes in the system to facilitate the delivery of self-management support through:

- reorganisation of staff roles;
- restructuring of care delivery.

In terms of clinical outcomes for patients, delivery system design interventions therefore have a less direct mode of action than self-management support interventions. They do not act directly to change patient behaviour but facilitate the delivery of interventions that do.

3. Decision support interventions

Decision support interventions work by promoting behaviour change on the part of the clinician. Again these have an indirect impact on patient behaviour. They work by providing the clinician with the knowledge and skills they need to provide self-management support as effectively as possible.

4. Clinical information system interventions

Again these have an indirect action on patient behaviour. These interventions work by using organising data to facilitate effective self-management support.

5. Community interventions

These interventions work by using resources within the wider community either by supporting the patient directly or by helping the health system function so that self-management support can be provided more effectively.

6. Health system interventions

How health system interventions work is complex, context-specific and less easy to quantify ([Kreindler 2009](#)). [Tsai 2005](#) also noted that health system interventions are difficult to manipulate empirically and that evidence is hard to find across the spectrum of long-term conditions.

Since the action and implementation of community and health system interventions cross the boundary between the direct healthcare patient-provider environment into the wider healthcare system and policy environment, we did not plan to carry out a detailed meta-analysis of effects for these two elements. Instead we documented whether any studies tested this type of intervention.

Why it is important to do this review

Researchers have argued that the negative consequences of hearing loss make a strong argument for early, effective hearing aid fitting ([Arlinger 2003](#)). Interventions that improve rates of hearing aid use should have an impact on such negative psychosocial consequences, both on an individual level and across the population with hearing loss who have been fitted with hearing aids.

In addition, if uptake of hearing aids is increased by the use of screening or education programmes ([Davis 2007](#); [Thodi 2013](#)), then it is important that subsequent hearing aid fitting is as effective as possible. There are also economic implications of non-use, both for national funding bodies and on an individual level for those purchasing their own hearing aids.

This review does not aim to compare the effects of context-specific interventions (e.g. auditory training, communication training) or modes of delivery (e.g. group versus individual interventions). However, adult hearing loss is an under-researched, under-theorised field. Hence we have employed a framework from the wider field of long-term conditions research and service development. We hope that this framework will provide information about high-level intervention types such as those that act directly to support patient behaviour change and those that seek to influence patient behaviour in less direct ways. However, we also hope to provide another level of detail using subgroup analyses for those stakeholders interested in, for example, subtypes of self-management support. We hope that by structuring the review in this way we will be able to encourage new research directions and highlight gaps in the evidence base.

OBJECTIVES

To assess the long-term effectiveness of interventions to promote the use of hearing aids in adults with acquired hearing loss fitted with at least one hearing aid.

METHODS

Criteria for considering studies for this review

Types of studies

We included randomised controlled trials (RCTs) that fulfilled the inclusion criteria. We included quasi-randomised trials such as those allocating by an arbitrary but not truly random process (e.g. day of the week) and cluster-randomised trials.

Types of participants

Adults with hearing loss greater than 25 dB hearing level (HL) in the better ear averaged across four frequencies (0.5 kHz, 1 kHz, 2 kHz and 4 kHz) who were fitted with a hearing aid for at least one ear. This is consistent with World Health Organization criteria for the definition of hearing loss (WHO 2000), and includes those with mild, moderate, severe and profound losses. Studies on the acceptability and benefit of hearing screening sometimes set different criteria for what constitutes a significant hearing loss (e.g. Davis 2007). These are generally more conservative and so would be included under the definition given above. Where trials did not give details of hearing levels for participants we assumed that those fitted with a hearing aid would have met these criteria. For the purposes of this review we considered adults to be aged 18 years and over. Trials that included participants under the age of 18 were included if the data for adults could be accessed separately by contacting the authors where it was not obvious from the trial data. We included adults with sensorineural, conductive and mixed hearing losses. We excluded trials that included participants using implantable devices such as bone-anchored hearing aids or cochlear implants.

Types of interventions

This review considered any healthcare interventions, classified according to the chronic care model (CCM), intended to increase the use of hearing aids. We excluded studies that tested or compared developments in hearing aid technology (see [Description of the intervention](#)).

Comparisons

- Self-management support interventions versus alternative interventions that control for other elements delivery method/pattern.
- Delivery system design interventions versus alternative interventions that control for content.
- Combined self-management support/delivery system design interventions versus standard care/control.
- Decision support interventions versus standard care.
- Clinical information system interventions versus standard care.

We planned to include subgroup analyses by self-management support content, delivery system design format and follow-up schedule (see [Subgroup analysis and investigation of heterogeneity](#)). Interventions were compared against each other, against no intervention or against 'standard care'. This review considered interventions supplementary to the hearing aid fitting process itself. We defined standard care as being a face-to-face individual hearing aid fitting typically lasting 45 to 60 minutes. We would expect a standard fitting to include a basic level of advice regarding use and management of the hearing aid with some practice at physical management of the device itself.

Types of outcome measures

The purpose of this review was to look at interventions that promote use of hearing aids once they have been fitted either by increasing the proportion of those fitted who become successful users or by increasing the amount of use per person. We recognise that for an individual use does not always equate to benefit but it is certainly a necessary starting point. At the most basic level it is not possible to benefit from a hearing aid if it is not in use for at least a proportion of the time.

As hearing loss is a long-term condition and hearing aids are usually intended as a long-term intervention, we were interested in hearing aid use after a follow-up period of at least a year. We also included short-term (≤ 12 weeks) and medium-term (> 12 to < 52 weeks) follow-up, but we considered this lower quality evidence than if long-term data were available for the same outcome.

Primary outcomes

1. Hearing aid use

The purpose of this review was to assess the degree to which any of the interventions described above resulted in the increased usage of hearing aids by the patient. This may be measured in many different ways (Perez 2012). This review uses the following measures:

1.1. Adherence (i.e. the proportion of participants who continued to use their hearing aids after fitting relative to the total number fitted). The World Health Organization defines adherence as "the extent to which a person's behaviour - taking medication, following a diet, and/or executing lifestyle changes, corresponds with agreed recommendations from a health care provider" (WHO 2003). This definition differs from purely behavioural definitions of use (is the patient wearing their hearing aid?) and compliance (is the patient wearing their hearing aid as recommended?). For the purposes of this review we assumed that those being fitted with a hearing aid had agreed to this management option. We have therefore defined number of aids in use/number fitted as adherence. Participants were classified as users or non-users. Users were defined as those who used their hearing aids on at least a weekly basis. Non-users were those who did not use their hearing aids at all or those who had not used their hearing aid for at least a week prior to follow-up data collection. Where it was unclear how often participants were using their hearing aids and how they had been classified as users or non-users, we attempted to contact the study authors for clarification. If we were unable to get clarification the study was excluded.

1.2. Daily hours of hearing aid use. This may be assessed using validated self-report measures that record the daily hours of hearing aid use or data-logging by the hearing aid itself. Modern hearing aids have the capacity to capture and record when the hearing aid is switched on. It does not represent a true objective measure of use because it is only able to measure whether the hearing aid is switched on and the acoustic environment it is in, not whether it

is switched on and in the patient's ear. However, we hoped to use it as a proxy measure of use. Both data collection methods yield continuous data either in terms of hours of use/time or proportion of the time the hearing aid(s) are worn. Since it is not the purpose of this review to compare methods of data collection, we combined data obtained using self-report and data-logging in our analyses of daily hours of hearing aid use.

2. Adverse effects

2.1 Inappropriate advice/clinical practice causing damage to patients' hearing.

2.2 Patient complaints:

- unresolved problems with physical management of the hearing aid;
- unresolved issues with symptom or psychosocial management;
- complaints relating to the nature of the intervention itself, such as having to make repeat visits to the clinic.

Secondary outcomes

For the purposes of this review we were interested in additional patient-reported outcomes that might be theoretically related to hearing aid use. Additional process-related outcomes such as utilisation, quality of care and resource use are outside the scope of this review.

We included validated measures of:

- quality of life - we included validated generic (e.g. SF-36, SF-12) and disease-specific measures of quality of life (e.g. IOI-HA item 7 [Cox 2002](#));
- hearing handicap - validated measures of residual handicap or activity limitations (e.g. Hearing Handicap Inventory for the Elderly (HHIE) ([Ventry 1982](#)), Hearing Coping Assessment (HCA) ([Andersson 1995a](#)), Hearing Measurement Scale (HMS) ([Noble 1970](#)), Hearing Performance Inventory (HPI) ([Giolas 1979](#)), QDS ([Alpiner 1978](#)), IOI-HA item 3);
- hearing aid benefit - validated measures of hearing aid benefit (e.g. Abbreviated Profile of Hearing Aid Benefit (APHAB) ([Cox 1995](#)), Glasgow Hearing Aid Benefit Profile (GHABP) ([Gatehouse 1999](#)), IOI-HA item 2);
- communication - any validated measure of communication ability or strategy (e.g. Communication Profile for the Hearing Impaired (CPHI) ([Demorest 1987](#))).

These measures might be completed by the patient, their communication partner(s) or both, with or without supervision from a clinician. We selected the outcomes we considered would be of most interest to patients, clinicians and policy-makers as recommended in the *Cochrane Handbook for Systematic Reviews of Interventions* ([Higgins 2011](#)). We reached the decision on which outcomes to include a priori following discussion between FB, EM and LE, who all have clinical experience in the context of hearing health care.

Search methods for identification of studies

The Cochrane ENT Information Specialist conducted systematic searches for randomised controlled trials and controlled clinical trials. There were no language, publication year or publication status restrictions. The date of the search was 13 June 2016.

Electronic searches

The Information Specialist searched:

- the Cochrane ENT Trials Register (searched 20 June 2016);
- the Cochrane Central Register of Controlled Trials (CENTRAL 2016, Issue 5);
- PubMed (1946 to 13 June 2016);
- Ovid EMBASE (1974 to 2016 June 10);
- Ovid CAB Abstracts (1910 to 2016 week 22);
- EBSCO CINAHL (1982 to 13 June 2016);
- Ovid AMED (1985 to 13 June 2016);
- LILACS, lilacs.bvsalud.org (searched 13 June 2016);
- KoreaMed (searched via Google Scholar 13 June 2016);
- IndMed, www.indmed.nic.in (searched 13 June 2016);
- PakMediNet, www.pakmedinet.com (searched 13 June 2016);
- Web of Knowledge, Web of Science (1945 to 13 June 2016);
- CNKI, www.cnki.com.cn (searched via Google Scholar 13 June 2016);
- ClinicalTrials.gov (searched via the Cochrane Register of Studies 13 June 2016);
- World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP), www.who.int/ictip (searched 13 June 2016);
- ISRCTN, www.isrctn.com (searched 13 June 2016);
- Google Scholar, scholar.google.co.uk (searched 17 June 2016);
- Google, www.google.com (searched 17 June 2016).

The Information Specialist modelled subject strategies for databases on the search strategy designed for CENTRAL. Where appropriate, they were combined with subject strategy adaptations of the highly sensitive search strategy designed by Cochrane for identifying randomised controlled trials and controlled clinical trials (as described in the *Cochrane Handbook for Systematic Reviews of Interventions* Version 5.1.0, Box 6.4.b. ([Higgins 2011](#))). Search strategies for major databases including CENTRAL are provided in [Appendix 1](#).

Searching other resources

We scanned the reference lists of identified publications for additional trials. We searched PubMed, *The Cochrane Library* and Google to retrieve existing systematic reviews relevant to this systematic review, so that we could scan their reference lists for ad-

ditional trials. We searched for conference abstracts using the Cochrane ENT Trials Register.

Data collection and analysis

Selection of studies

Material downloaded from electronic sources included details of author, institution or journal of publication and abstract. FB and EM inspected all reports independently in order to ensure reliable selection. We resolved any disagreement by discussion and, where there was still doubt, we acquired the full article for further inspection. Once the full articles were obtained, we decided whether the studies met the review criteria. If disagreement could not be resolved by discussion, we sought further information and added these trials to the list of those awaiting assessment.

Data extraction and management

Data extraction

Review authors FB and LE independently extracted data from all included studies. Again, we discussed any disagreement, documented decisions and, if necessary, contacted authors of studies for clarification. We extracted data presented only in graphs and figures whenever possible, but included them only if two review authors independently came to the same result. We attempted to contact authors through an open-ended request in order to obtain missing information or for clarification whenever necessary. If studies were multicentre, where possible, we extracted data relevant to each component centre separately.

Data management

Forms

We extracted data onto standard, simple forms, which are available on request from the corresponding author.

Scale-derived data

We included ordinal data from rating scales only if:

- the psychometric properties of the measuring instrument had been described in a peer-reviewed journal; and
- the measuring instrument had not been written or modified by one of the investigators for that particular trial.

We considered the ideal measuring instrument to be either i) self-report or ii) completed by an independent rater or relative (not the clinician).

Endpoint versus change data

There are advantages of both endpoint and change data. Change data can remove a component of between-person variability from the analysis. On the other hand, calculation of change needs two assessments (baseline and endpoint), which increases the likelihood of missing data points. We primarily used endpoint data and only used change data if the former were not available. Where appropriate we used standardised mean differences to combine endpoint and change data in the analyses (Higgins 2011).

Skewed data

Continuous data on clinical and social outcomes are often not normally distributed. To avoid the pitfall of applying parametric tests to non-parametric data, we applied the following standards to all data before inclusion:

- standard deviations and means were reported in the paper or obtainable from the authors;
- when a scale started from zero, the mean should be more than twice the standard deviation (as otherwise the mean was unlikely to be an appropriate measure of the centre of the distribution (Altman 1996);
- if a scale started from a positive value we modified the calculation described above to take the scale starting point into account. In these cases skew was present if $2SD > (S - S_{min})$, where S is the mean score and S min is the minimum score.

Endpoint scores on scales often had a finite start and endpoint and these rules could be applied. We entered potentially skewed endpoint data from studies into our analyses but noted the high risk of skew, downgrading our judgement of the quality of the evidence for a particular outcome where the majority of studies were at high risk of bias.

When continuous data were presented on a scale that included a possibility of negative values (such as change data) and it was difficult to tell whether data were skewed or not, we entered change data but again noted where we considered it to be of potential significance when interpreting the evidence.

Common measure

To facilitate comparison between trials, we converted variables that were reported in different metrics, such as hours of use (mean hours per day, per week or per month) to a common metric. We used mean hours per day. For conversion purposes we considered a full day to equal 12 hours since hearing aids are not normally worn at night.

Direction of graphs

For outcomes where a higher score was judged to be a positive outcome (such as daily hours of use or quality of life), we displayed the results so that the area to the left of the line of no effect indicated a favourable outcome for the control group. For outcomes where a

higher score was judged to be a negative outcome (such as hearing handicap), we displayed the results so that the area to the left of the line of no effect indicated a favourable outcome for the intervention group.

Assessment of risk of bias in included studies

Authors FB and EM independently undertook an assessment of the risk of bias of the included trials as guided by the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). This set of criteria is based on evidence of associations between overestimate of effect and high risk of bias of the article such as sequence generation, allocation concealment, blinding, incomplete outcome data and selective reporting.

We used the Cochrane 'Risk of bias' tool in RevMan 5.3 (RevMan 2014), which involves describing each of the domains as reported in the trial and then assigning a judgement about the adequacy of each entry: 'low', 'high' or 'unclear' risk of bias. We judged that any study that had a high risk of bias in three or more areas had an overall high risk of bias and we subjected those to a sensitivity analysis (see [Sensitivity analysis](#)).

Where the raters disagreed, we made the final rating by consensus, with the involvement of another member of the review group. Where inadequate details of randomisation and other characteristics of trials were provided, we attempted to contact authors of the studies in order to obtain further information. We recorded non-concurrence in quality assessment and, where there was disagreement as to which category a trial was to be allocated, again we resolved this by discussion.

We noted the level of risk of bias in both the text of the review and in the 'Summary of findings' tables.

Measures of treatment effect

Binary data

For binary outcomes we calculated a standard estimation of the risk ratio (RR) and its 95% confidence interval (CI). It has been shown that the RR is more intuitive than the odds ratio (Boissel 1999), and additionally that odds ratios tend to be interpreted as RR by clinicians (Deeks 2000). Where we identified heterogeneity we planned to use a random-effects model.

Continuous data

If continuous data, for example from hearing aid benefit questionnaires, were measured on the same scale, we used the mean difference to summarise the results between studies. For outcomes measured using different scales, we calculated the standardised mean difference (SMD) to combine the results.

Unit of analysis issues

Cluster trials

We anticipated that some studies might employ 'cluster-randomisation' (such as randomisation by clinician or practice) and we planned for how we would deal with this statistically to reduce the risk of 'unit of analysis' errors (Divine 1992). In the event no trials involving cluster-randomisation were included in this review.

2. Cross-over trials

A major concern of cross-over trials is the carry-over effect. It occurs if an effect (e.g. pharmacological, physiological or psychological) of the treatment in the first phase is carried over to the second phase. As a consequence, on entry to the second phase the participants can differ systematically from their initial state despite a wash-out phase. For the same reason cross-over trials are not appropriate if the condition of interest is unstable or progressive. As both these possibilities arise with hearing loss, we sought to use only data from the first phase of cross-over studies.

3. Studies with multiple treatment groups

Where a study involved more than two treatment arms, if relevant, we presented the additional treatment arms in comparisons. If data were binary we added and combined these within the two-by-two table. If data were continuous we combined data following the formula in section 7.7.3.8 ('Combining groups') of the *Cochrane Handbook for Systematic Reviews of Interventions* where appropriate (Higgins 2011). Where the additional treatment arms were not relevant, we did not use these data.

Dealing with missing data

Overall loss of credibility

At some degree of loss of follow-up data must lose credibility. We decided that, for any particular outcome, should more than 50% of data be unaccounted for, we would not present these data or use them within analyses. If, however, more than 50% of those in one arm of a study were lost, but the total loss was less than 50%, we marked such data with (*) to indicate that such a result may well be prone to bias.

Binary

In the case where attrition for a binary outcome was between 0% and 50% and where these data were not clearly described, we presented data on a 'once randomised always analyse' basis (an intention-to-treat analysis). We assumed those leaving the study early to have the same rates of negative outcome as those who completed. We planned to undertake a sensitivity analysis to test

how prone the primary outcomes are to change when data only from people who completed the study to that point were compared with the intention-to-treat analysis using the above assumption.

Continuous

Attrition

In the case where attrition for a continuous outcome was between 0% and 50% we reported data only from people who completed the study to that point.

Standard deviations

If standard deviations were not reported, we first tried to obtain the missing values from the authors. If not available, where there were missing measures of variance for continuous data, but an exact standard error and confidence intervals were available for group means, and either P value or t value were available for differences in mean, we calculated them according to the rules described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). When only the standard error (SE) was reported, we attempted to calculate standard deviations (SDs) by the formula $SD = SE * \text{square root } (n)$. Chapters 7.7.3 and 16.1.3 of the *Cochrane Handbook for Systematic Reviews of Interventions* present detailed formulae for estimating SDs from P values, t or F values, confidence intervals, ranges or other statistics (Higgins 2011).

Last observation carried forward

We anticipated that in some studies the method of last observation carried forward (LOCF) might be employed within the study report. As with all methods of imputation to deal with missing data, LOCF introduces uncertainty about the reliability of the results. Therefore, where LOCF data were used in a trial, if less than 50% of the data was assumed, we presented and used these data and indicated that they are the product of LOCF assumptions.

Assessment of heterogeneity

Clinical diversity

We considered all included studies initially, without seeing comparison data, to look for variations in participants, interventions and outcomes (clinical diversity). We inspected all studies for clearly outlying people or situations that we had not predicted would arise. We planned theory-led subgroup analyses based on CCM element definitions and long-term conditions research (see [Subgroup analysis and investigation of heterogeneity](#)).

Methodological diversity

We considered all included studies initially, without seeing comparison data, to look for variability in study design and risk of bias (methodological diversity). We inspected all studies for clearly outlying methods that we had not predicted would arise.

Statistical heterogeneity

Heterogeneity may arise as a result of clinical or methodological diversity, or both. We assessed it in two ways.

Visual inspection

We visually inspected graphs to investigate the possibility of statistical heterogeneity by looking at the degree of overlap between confidence intervals.

Employing the I^2 statistic

We investigated heterogeneity between studies by considering the I^2 statistic alongside the Chi^2 test P value. The I^2 statistic provides an estimate of the percentage of inconsistency thought to be due to chance. The importance of the observed value of I^2 depends on i) the magnitude and direction of effects and ii) the strength of evidence for heterogeneity (e.g. P value from Chi^2 test, or a confidence interval for I^2). We have interpreted an I^2 estimate greater than or equal to around 50% accompanied by a statistically significant Chi^2 value as evidence of substantial levels of heterogeneity (Section 9.5.2 - Higgins 2011).

Assessment of reporting biases

Reporting biases arise when the dissemination of research findings is influenced by the nature and direction of results (Egger 1997). These are described in section 10.1 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011).

Protocol versus full study

We attempted to locate protocols for included randomised trials. If the protocol was available, we compared outcomes in the protocol and in the published report. If the protocol was not available, we compared outcomes listed in the methods section of the trial report with the results actually reported.

Funnel plot

We are aware that funnel plots may be useful in investigating reporting biases but are of limited power to detect small study effects. We did not use funnel plots for outcomes where there were 10 or fewer studies, or where all studies were of similar sizes.

Data synthesis

Fixed-effect models hold that only within-study variation influences the uncertainty of an effect (as reflected in the confidence interval). Variation between the estimates of effect from each study (heterogeneity) does not influence the confidence interval in a fixed-effect model. Random-effects models incorporate an assumption that the different studies are estimating different (yet related) but not fixed intervention effects.

In a group of studies where there is low heterogeneity, fixed-effect and random-effects models will return the similar confidence intervals. However, where there is evidence of statistical heterogeneity this will be taken into account only by a random-effects model analysis and the confidence intervals will be wider than they would be when analysing the same data using a fixed-effect model. In terms of identifying evidence of significant effects a random-effects model is therefore more conservative. However, it does put more weight on the smaller studies, which are often the most biased. Depending on the direction of effect these studies can either inflate or deflate effect size.

Since we anticipated a degree of clinical and methodological heterogeneity in these data, given the wide range of interventions included, we used a random-effects model for all analyses. To investigate heterogeneity further, where appropriate, we carried out a series of theory-led subgroup analyses based on the CCM element definitions and previous research carried out in other long-term conditions and we assessed risk of bias (see [Subgroup analysis and investigation of heterogeneity](#)).

Subgroup analysis and investigation of heterogeneity

Subgroup analyses

In this review, we have grouped the results into comparisons within the CCM element. We anticipated that there would be diversity of intervention within a CCM element (see [Description of the intervention](#)) and so we planned to use the CCM element definitions and previous research analysing complex interventions in long-term conditions to perform subgroup analyses where appropriate.

Due to the wide range of skills needed to live well with a long-term condition, self-management support interventions can be varied and complex. Based on the work of [Lorig 2003](#), [Pearson 2007](#) and [Whitlock 2002](#) and previous reviews by [Barlow 2002](#), [Kreindler 2009](#) and [Tsai 2005](#), results for comparisons that include a component of self-management support should be subject to a subgroup analysis as follows:

- Advise: interventions aiming to inform and educate the patient about any aspect of self-management.
- Activate - practical skills: interventions that include practice of practical skills in terms of hearing aid management.
- Activate - symptoms management skills: interventions that include practice addressing the direct symptoms associated with

hearing loss, i.e. reduced sound/speech perception/discrimination.

- Activate - psychosocial management skills: interventions that include practice addressing the psychosocial and emotional consequences of hearing loss, i.e. communication difficulties, acceptance of hearing loss etc.
- Assist: interventions that include the provision of additional practical tools to support self-management.
- Agree: collaborative decision-making.

We have assumed that an assessment of need has been incorporated into all the self-management support interventions and so have chosen not to include this as a discrete subgroup. We have augmented the 5As model with behavioural activation subgroups based on the work of [Barlow 2002](#) and [Pearson 2007](#).

This subdivision of self-management provision was supported by the results of a Delphi review involving a panel of 26 hearing healthcare stakeholders including patients, clinicians, researchers and commissioners. It involved a three-round online Delphi process to investigate whether consensus could be reached on what it means to live well with a hearing loss, how this might be measured and the clinical processes that might support it ([Barker 2015](#)). The relative effect of these subgroups of self-management support would be of interest to patients, clinicians and policy-makers. The division into 'informing' and 'involving' processes has also recently been suggested as a way to operationalise patient-centred care within hearing healthcare ([Grenness 2014](#)).

Results for comparisons that include a component of delivery system design were subject to subgroup analyses as follows:

Delivery system design format:

- Face-to-face
- Telephone
- Booklet
- Remote (online/PC-based/DVD/video)
- Other

Delivery system design intensity:

- Low-intensity - single session interventions
- Medium-intensity - up to four session interventions
- High-intensity - five or more session interventions

The cut off between medium- and high-intensity interventions was chosen based on clinical experience.

This was based on the clinical experience of FB, LE and EM and on the results of the Delphi review described above where there was consensus that follow-up scheduling may be an important factor in supporting someone to live well with their hearing loss. The effect of delivery format may be of interest to patients and healthcare professionals and policy-makers interested in system redesign.

We recognise the possibility of interaction in effect between content, follow-up pattern and format but it was not the intention of this review to carry out a full multiple regression analysis to

investigate this. We review the relevance and usefulness of the use of these research-based subgroups in the [Discussion](#).

Investigation of heterogeneity

We anticipated that there might be a high degree of heterogeneity across eligible studies due to variations in patient populations, characteristics of interventions, outcome measurement, study design and risk of bias. Where this was found to be the case for a particular outcome we first checked that all data had been entered correctly, including checking for unit of analysis errors. We then sought to investigate remaining clinical heterogeneity using subgroup analyses. Where this did not adequately reduce heterogeneity we then went back to the original papers and study designs looking for studies that shared common characteristics in terms of population, intervention, comparison and outcome. We assessed the impact of risk of bias using sensitivity analysis (see [Sensitivity analysis](#)).

Sensitivity analysis

We carried out sensitivity analyses based on the quality criteria reported in this review.

Implication of randomisation

We included trials when they were described in some way as to imply randomisation even when details were not given of the allocation process. For the primary outcome we included these studies and if there was no substantive difference when the implied randomised studies were added to those with a better description of randomisation, then we employed all data from these studies.

Assumptions for lost binary data

Where assumptions had to be made regarding people lost to follow-up, we planned to compare the findings of the primary outcomes when using our assumptions compared with completer data only. If there was a substantial difference, we planned to report the results and discuss them but continue to employ our assumptions. We had also planned to follow a similar protocol where assumptions were made regarding missing SD data.

Risk of bias

We analysed the effects of excluding trials that we judged to be at overall high risk of bias (see [Assessment of risk of bias in included studies](#)). Where the exclusion of trials at high risk of bias did

not substantially alter the direction of effect or the precision of the effect estimates, then we included data from these trials in the analysis. If it did alter the direction or precision of effects we included the data but discussed the implications when presenting the results (see [Effects of interventions](#); [Summary of findings for the main comparison](#); [Summary of findings 2](#); [Summary of findings 3](#)).

Imputed values

We planned also to undertake a sensitivity analysis to assess the effects of including data from trials where we had to use imputed values for the intra-cluster correlation coefficient (ICC) in calculating the design effect in cluster-randomised trials.

RESULTS

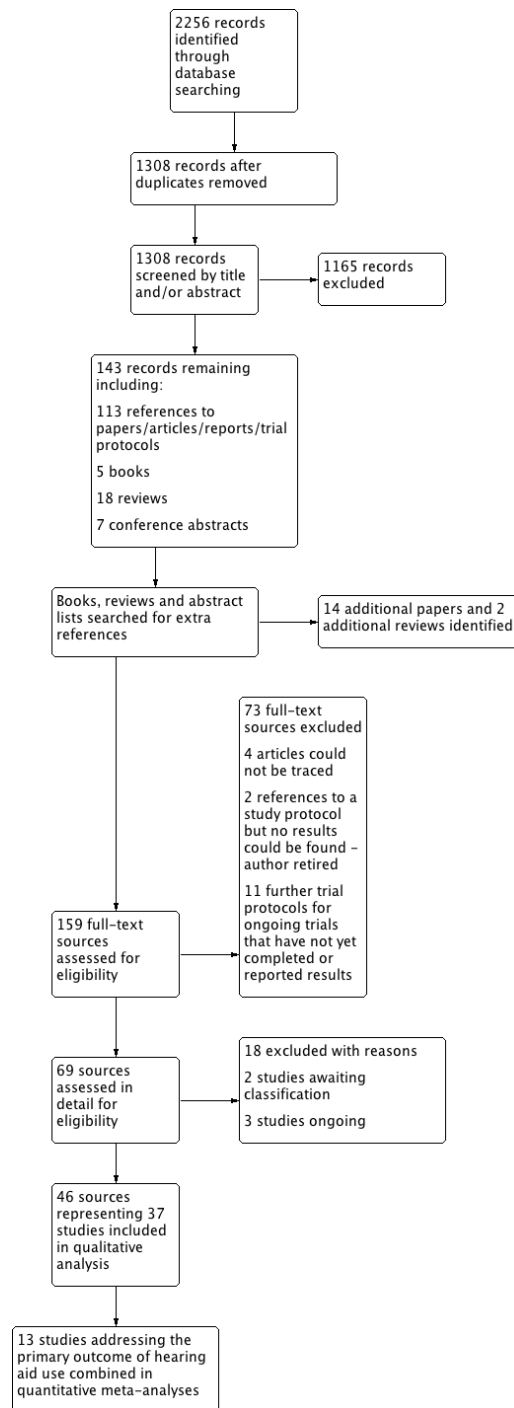
Description of studies

See: [Characteristics of included studies](#); [Characteristics of excluded studies](#); [Table 1](#).

Results of the search

The search identified 2256 papers, reviews, book chapters and conference abstracts, of which 1308 remained once duplicates were removed. We discarded 1165 papers on the basis of the title and/or abstract leaving 143 remaining sources for which we searched the full text. We also searched their reference lists and this identified a further 14 papers and two reviews, which we also attempted to access in full text. Of these 159 sources, we discarded 73 on the basis that they did not meet the inclusion criteria, four could not be traced, 11 referred to study protocols for which results were not available and two were abstracts for oral presentations but the authors could not be traced. This left 69 papers that we analysed in detail. We subsequently excluded 18 of these for the reasons given in [Excluded studies](#) and two completed studies await classification ([Henshaw 2013](#); [Malmberg 2015](#)). Three studies are ongoing ([ISRCTN77340339](#); [NCT02233361](#); [NCT02264314](#)). Forty-six papers giving results from 37 original studies were eligible for inclusion in the review ([Figure 1](#)). Quantitative data for the primary outcome of hearing aid use were reported in 22 studies, but we were able to combine only 13 of these studies in quantitative meta-analyses.

Figure 1. PRISMA diagram showing process for sifting search results and selecting studies for inclusion.



Included studies

Participants

In all the included studies participants were adults as defined in [Types of participants](#). Some studies included participants aged in their 20s and upwards (e.g. [Collins 2013](#); [Smaldino 1988](#); [Sweetow 2006](#); [Thoren 2011](#)), but the majority of the studies included participants aged 50 or above. Even in those studies that included younger participants the mean age was generally in the 60- to 70-year age range. Other frequently applied inclusion criteria were that participants should have no evidence of additional cognitive or physical impairment that might have an impact on hearing aid use and that their hearing loss was sensorineural in nature.

Where information was reported we also looked at the gender of participants. Seven of the studies were carried out in a US military veteran population and hence included an overwhelmingly male population ([Abrams 1992](#); [Chisolm 2004](#); [Collins 2013](#); [Preminger 2010a](#); [Saunders 2009](#); [Saunders 2016](#); [Turbin 2006](#)).

Interventions

1. Self-management support interventions

Five studies reported comparisons that changed the content of self-management support in isolation ([Fitzpatrick 2008](#); [Kricos 1996](#); [Preminger 2010a](#); [Saunders 2009](#); [Saunders 2016](#)). Alternative interventions were compared to control for changes in delivery system design (see [Table 1](#)).

There were no studies that sought to investigate the effect of providing resources to support self-management (assist) or the role of collaborative decision-making, goal-setting or action-planning (agree).

2. Delivery system design interventions

Six studies reported comparisons that changed the delivery of self-management support and included comparison interventions that controlled for changes in self-management support content ([Campos 2013](#); [Cherry 1994](#); [Collins 2013](#); [Cunningham 2001](#); [Lavie 2014](#); [Ward 1981](#)). Four studies changed the format of delivery ([Campos 2013](#); [Cherry 1994](#); [Lavie 2014](#); [Ward 1981](#)), two changed the intensity ([Cherry 1994](#); [Cunningham 2001](#)), and one changed the mode ([Collins 2013](#)).

There were no studies that sought to investigate staff roles and task distribution amongst team members on the usage of hearing aids. No studies specifically addressed participants' understanding

of the care they received or investigated whether it fitted in with their cultural background.

Twenty-nine studies reported on comparisons of combined self-management support/delivery system design interventions where the self-management support content and delivery were changed. Where interventions were compared against standard care or 'no intervention' we defined this as a standard individual hearing aid fitting comprising a single visit for the fitting itself and any routine follow-up session(s).

3. Decision support interventions

None found.

4. Clinical information system interventions

None found.

5. Community resource interventions

None found.

6. Health system interventions

None found.

Details of interventions are given in [Characteristics of included studies](#) and summarised in [Table 1](#). All interventions in the included studies could be classified according to the chronic care model (CCM). The majority involved both self-management support and delivery system design changes.

Outcomes

This review aimed to look at long-term outcomes as hearing loss is a long-term condition requiring self-management on the part of the patient over many years. Only six of the studies we identified looked at outcome over one year or longer ([Andersson 1994](#); [Andersson 1995](#); [Cherry 1994](#); [Chisolm 2004](#); [Oberg 2008](#); [Oberg 2009](#)), and only two of these addressed the primary outcome of hearing aid use ([Oberg 2008](#); [Oberg 2009](#)) (see [Summary of findings 3](#)).

Primary outcomes

1. Hearing aid use

1.1. [Collins 2013](#) and [Campos 2013](#) reported data that could be interpreted as adherence as defined in this review, although only the [Collins 2013](#) study actually specified adherence as an outcome.

1.2. Twenty-two studies measured daily hours of use or used a scale that could be converted to daily hours of use (Andersson 1995; Andersson 1997; Campos 2013; Cherry 1994; Collins 2013; Cunningham 2001; Eriksson-Mangold 1990; Ferguson 2016; Fitzpatrick 2008; Kemker 2004; Kramer 2005; Lavie 2014; Lundberg 2011; Oberg 2008; Oberg 2009; Olson 2013; Saunders 2009; Thoren 2011; Thoren 2014; Vreeken 2015; Ward 1978; Ward 1981). Campos 2013, Lavie 2014 and Ferguson 2016 used data-logging to measure hours of use per day in addition to or instead of self-reported hours of use.

2. Adverse effects

2.1 No studies reported on clinical adverse events.

2.2 Cherry 1994 looked at the number of outstanding complaints at one year after the provision of telephone follow-up.

Secondary outcomes

1. Quality of life

Twelve studies reported quality of life as an outcome measure (Ferguson 2016; Kramer 2005; Lundberg 2011; Oberg 2008; Oberg 2009; Olson 2013; Preminger 2008; Preminger 2010; Preminger 2010a; Thoren 2011; Thoren 2014; Vreeken 2015). Only Oberg 2008 and Oberg 2009 reported on long-term quality of life.

2. Hearing handicap

Twenty-seven studies reported hearing handicap as an outcome (Abrams 1992; Andersson 1994; Andersson 1995; Andersson 1997; Beynon 1997; Cherry 1994; Chisolm 2004; Collins 2013; Ferguson 2016; Kramer 2005; Kricos 1992; Kricos 1996; Lundberg 2011; Miranda 2008; Oberg 2008; Oberg 2009; Preminger 2008; Preminger 2010; Preminger 2010a; Saunders 2009; Saunders 2016; Smaldino 1988; Sweetow 2006; Thoren 2011; Thoren 2014; Ward 1978; Ward 1981). Only Andersson 1994; Andersson 1995; Oberg 2008 and Oberg 2009 reported long-term hearing handicap.

3. Hearing aid benefit

Fourteen studies reported hearing aid benefit as an outcome (Collins 2013; Cunningham 2001; Ferguson 2016; Gil 2010; Kemker 2004; Kramer 2005; Lundberg 2011; Oberg 2008; Oberg 2009; Olson 2013; Saunders 2009; Thoren 2011; Thoren 2014; Vreeken 2015), but only Oberg 2008 and Oberg 2009 did so over the long term.

4. Communication

Eight studies reported a measure of communication as an outcome (Andersson 1997; Chisolm 2004; Collins 2013; Kricos 1996; Oberg 2008; Preminger 2010; Sweetow 2006; Turbin 2006), but only Chisolm 2004 and Oberg 2008 did so over the long term. Only two of the studies reported an overall single score measure of communication (Preminger 2010; Sweetow 2006). The remaining studies used the Communication Profile for the Hearing Impaired (CPHI) to measure communication ability (Demorest 1987), with some choosing to use only the communication strategies subscale of this measure. This measures whether people use verbal, non-verbal and maladaptive strategies for communication. We took the verbal strategy subscale of the CPHI as an example measure of communication. This subscale was chosen post hoc on the basis that a primary aim of hearing aid fitting and subsequent rehabilitation is to improve verbal communication. However, we recognise that this scale does not represent the range of communicative ability or potential improvement. Where appropriate this has been taken into consideration when grading the quality of the evidence.

Excluded studies

Details of the 18 studies that we excluded after careful study of their methods are given in [Characteristics of excluded studies](#).

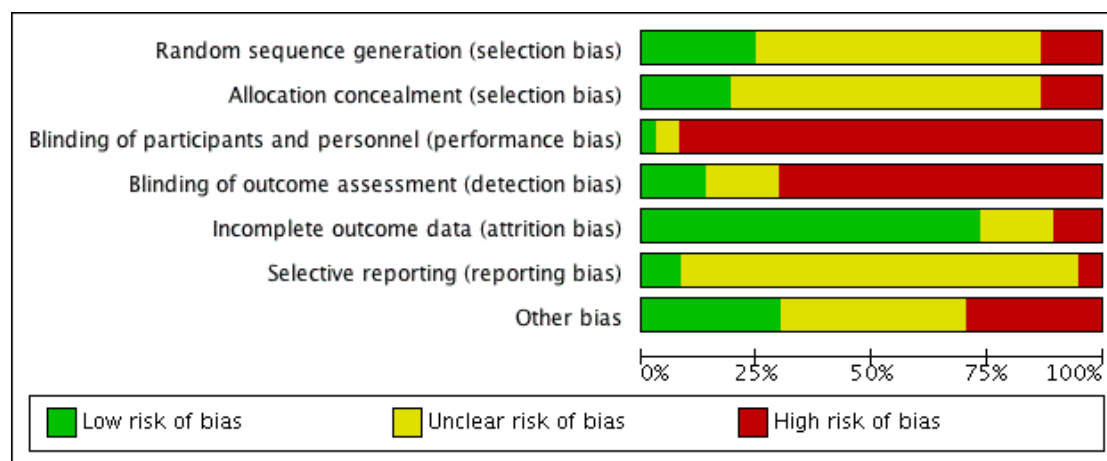
Risk of bias in included studies

In general the risk of bias was unclear or high in most studies. Please see [Figure 2](#) for the 'Risk of bias' analysis for the individual included studies and [Figure 3](#) showing the review authors' judgements about each 'Risk of bias' item presented as percentages across all included studies. Specific areas of concern are highlighted below.

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
Abrams 1992	?	?	?	?	?	?	?
Andersson 1994	?	?	?	?	?	?	?
Andersson 1995	?	?	?	?	?	?	?
Andersson 1997	?	?	?	?	?	?	?
Beynon 1997	?	?	?	?	?	?	?
Campos 2013	?	?	?	?	?	?	?
Cherry 1994	?	?	?	?	?	?	?
Chisolm 2004	?	?	?	?	?	?	?
Collins 2013	?	?	?	?	?	?	?
Cunningham 2001	?	?	?	?	?	?	?
Eriksson-Mangold 1990	?	?	?	?	?	?	?
Ferguson 2016	?	?	?	?	?	?	?
Fitzpatrick 2008	?	?	?	?	?	?	?
Gil 2010	?	?	?	?	?	?	?
Kemker 2004	?	?	?	?	?	?	?
Kramer 2005	?	?	?	?	?	?	?
Kricos 1992	?	?	?	?	?	?	?
Kricos 1996	?	?	?	?	?	?	?
Lavie 2014	?	?	?	?	?	?	?
Lundberg 2011	?	?	?	?	?	?	?
Miranda 2008	?	?	?	?	?	?	?
Oberg 2008	?	?	?	?	?	?	?
Oberg 2009	?	?	?	?	?	?	?
Olson 2013	?	?	?	?	?	?	?
Preminger 2008	?	?	?	?	?	?	?
Preminger 2010	?	?	?	?	?	?	?
Preminger 2010a	?	?	?	?	?	?	?
Saunders 2009	?	?	?	?	?	?	?
Saunders 2016	?	?	?	?	?	?	?
Smaldino 1988	?	?	?	?	?	?	?
Sweetow 2006	?	?	?	?	?	?	?
Thoren 2011	?	?	?	?	?	?	?
Thoren 2014	?	?	?	?	?	?	?
Turbin 2006	?	?	?	?	?	?	?
Vreeken 2015	?	?	?	?	?	?	?
Ward 1978	?	?	?	?	?	?	?
Ward 1981	?	?	?	?	?	?	?

Figure 3. 'Risk of bias' graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.



Allocation

It was rare for studies to give an adequate description of their randomisation process. Sequence generation and allocation concealment were frequently not mentioned at all so it was not possible to make a clear assessment of risk of selection bias. Only 16 of the 37 included studies gave any description of the allocation process. Of these, in nine studies the description was enough to allocate a low risk of selection bias. In the remaining seven studies the information given led us to judge that there was an unclear or high risk of selection bias.

Blinding

Due to the nature of the interventions in this context it is difficult to design studies that are blinded to participants and those delivering the intervention so performance bias is difficult to control for. See [Table 1](#) for a list of interventions and control conditions. Blinding in outcome assessment was mentioned more frequently than blinding for group allocation, although it was still rare.

Incomplete outcome data

A strength of the studies we identified is that they had low dropout rates even for long-term follow-up periods of over a year and there were only occasional instances of unexplained losses to follow-up.

Selective reporting

There was only one case of definite reporting bias ([Andersson 1994](#)), where an outcome had been recorded in the study but not reported in the paper. We discovered this because the data were later included in a paper ([Andersson 1998](#)), which combined data from three previous studies including [Andersson 1994](#). In most other cases it was not possible to make a clear judgement on reporting bias due to the lack of published protocols in this context. Where protocols were available, there was no evidence of selective reporting.

Other potential sources of bias

With a few exceptions, studies were small and lacked power calculations. Some studies were funded by hearing aid manufacturers, although this should not introduce undue bias as both control and interventions groups were provided with hearing aids in all cases. In the [Discussion](#) we consider the possible implications of studying participants from a tightly defined population such as military veterans, which was an issue in several of the included studies.

Effects of interventions

See: [Summary of findings for the main comparison Self-management support interventions for adults with hearing loss who use hearing aids](#); [Summary of findings 2 Delivery system design interventions for adults with hearing loss who use hearing](#)

aids; [Summary of findings 3](#) Combined self-management support/delivery system design interventions for adults with hearing loss who use hearing aids

[Summary of findings for the main comparison](#) summarises the evidence of effect for **self-management support** interventions on the primary and secondary outcomes.

[Summary of findings 2](#) summarises the evidence of effect for **delivery system design** interventions on the primary and secondary outcomes.

[Summary of findings 3](#) summarises the evidence of effect for **combined self-management support/delivery system design** interventions on the primary and secondary outcomes.

Where possible we have presented data on long-term outcomes. Short- and medium-term outcomes are included only where long-term outcome data were not available.

Self-management support interventions

Primary outcome

1. Hearing aid use

1.1 Adherence

We found no studies of self-management support interventions that reported adherence as an outcome.

1.2 Daily hours of hearing aid use

Two self-management support studies measured short- to medium-term daily hours of hearing aid use, but we were unable to combine them in a meta-analysis as they categorised daily use in a different way from our definition in this review ([Fitzpatrick 2008](#); [Saunders 2009](#)). [Fitzpatrick 2008](#) reported that for their auditory training intervention eight participants (57%) wore their hearing aids all of the time before, after and during therapy and six participants (43%) wore their hearing aids in more listening situations after therapy. In the control group (who received lectures on hearing loss, hearing aids and communication over the same time period) seven participants (70%) wore their hearing aids all the time and three participants (30%) wore their aids in limited situations before and after the lectures. [Saunders 2009](#) reported that when comparing a pre-fitting demonstration of listening situations with no demonstration, 4/20 participants in the intervention group and 1/20 participants in the control group wore their hearing aids for more than eight hours per day. The clinical significance of these results is unclear.

2. Adverse effects

No self-management support intervention studies reported on adverse effects.

Secondary outcomes

1. Quality of life

One self-management support intervention showed no statistically significant evidence of effect of adding psychosocial exercises to a communication training programme on short to medium-term quality of life (one study, 35 participants; mean difference (MD) -9.10, 95% confidence interval (CI) -21.33 to 3.13; [Analysis 1.1](#)) ([Preminger 2010a](#)). This represents a reduction of 9.1 points on the World Health Organization Disability Assessment Schedule 2.0 (WHODAS 2.0) 0- to 100-point scale. On this scale a lower score indicates improved quality of life. However, the minimal important difference on this scale for hearing loss has not been established. This means we cannot comment on the clinical significance of this result. We found no self-management support studies that reported long-term quality of life. Our confidence in the quality of the evidence for the effect of self-management support interventions on quality of life is very low.

2. Hearing handicap

We were able to combine the data from two self-management support interventions that assessed short- to medium-term hearing handicap ([Kricos 1996](#); [Preminger 2010a](#)). There was evidence of a short- to medium-term effect on hearing handicap (two studies, 87 participants; MD -12.80, 95% CI -23.11 to -2.48; [Analysis 1.2](#)). Although this represents a statistically significant change in the mean difference, it falls below the 18.7-point difference considered to represent a minimal important difference on this 100-point scale ([Ventry 1982](#); [Weinstein 1986](#)). The minimal important difference does fall within the confidence interval in this analysis, which suggests that there may have been a clinically significant effect on hearing handicap for some, but not all, participants. We found no self-management support interventions that reported long-term hearing handicap. Our confidence in the quality of evidence for the effect of self-management support interventions on self-reported hearing handicap is very low.

3. Hearing aid benefit

We found no studies of self-management support interventions that reported hearing aid benefit as an outcome.

4. Communication

One study that included a comparison of a self-management support intervention reported data on communication in the short to medium term (Kricos 1996). There was evidence of a short-term effect on the use of verbal communication strategies for this intervention, which compared an active listening programme with auditory training (one study, 52 participants; MD 0.72, 95% CI 0.21 to 1.23; Analysis 1.3). The minimal important difference on this subscale of the communication profile for the hearing impaired is 0.93 (Demorest 1988). The mean difference and confidence intervals suggest that for some, but not all, participants there was a clinically significant difference in the use of a communication strategy. We found no self-management support interventions that reported long-term communication. Our confidence in the quality of the evidence for the effect of self-management support interventions on communication is very low.

Delivery system design interventions

Primary outcome

1. Hearing aid use

1.1 Adherence

Two delivery system design studies yielded data that could be analysed as adherence (people fitted with aids/people using aids) (Campos 2013; Collins 2013). Collins 2013 asked participants whether they wore their hearing aids or not after six months. Campos 2013 used data-logging to record those with zero hours of use over the short term. These studies involved changes in mode (group fitting versus individual fitting; Collins 2013) and format (teleconsultation versus online fitting; Campos 2013). Combining these studies shows no evidence of short- to medium-term effects on adherence for these delivery system design interventions (two studies, 686 participants; risk ratio (RR) 1.02, 95% CI 0.99 to 1.05; Analysis 2.1). This equates, on average, to an additional 19 people out of 1000 wearing their hearing aid up to six months post-intervention. We found no studies that reported the effect of delivery system redesign on adherence in the long term. Our confidence in the quality of the evidence for the effect of delivery system design interventions on adherence is low.

1.2 Daily hours of hearing aid use

Six delivery system design studies reported daily hours of hearing aid use over the short- to medium-term (Campos 2013; Cherry 1994; Collins 2013; Cunningham 2001; Lavie 2014; Ward 1981). The data from the Ward 1981 and Lavie 2014 studies could not be

combined in a meta-analysis. Ward 1981 reported no significant difference in hours of use between a group given information about hearing tactics in written format versus face-to-face. Lavie 2014 reported that simultaneous bilateral hearing aid fitting resulted in significantly more hours of use per day than bilateral hearing aids fitted sequentially in two separate visits. In Campos 2013, they measured self-reported daily hours of use and data-logged hours of use. We could not use the self-reported hours of use in this analysis because no standard deviations or other measures of variance were reported in the study. However, they did report high levels of correlation ($r = 0.81$, P value = 0.00 for the intervention group and $r = 0.74$, P value = 0.00 for the control group) between the self-reported data and the data-logging. We have therefore combined the data-logging results in this analysis. There was no evidence of a short- to medium-term statistically significant effect on daily hours of hearing aid use for these delivery system design interventions (four studies, 700 participants; MD -0.06, 95% CI -1.06 to 0.95; Analysis 2.2). This MD equates to the participants in the intervention groups wearing their hearing aids for three to four minutes less in each day than those in the control groups. We found no delivery system design interventions that reported daily hours of hearing aid use in the long term. Our confidence in the quality of the evidence for the effect of delivery system design interventions on daily hours of hearing aid use is very low.

2. Adverse effects

2.1 No studies reported on clinical adverse events.

2.2 Only one study looked at the number of outstanding complaints after the provision of telephone follow-up and reported no statistically significant difference in the number of complaints at one-year follow-up (one study, 98 participants; RR 0.75, 95% CI 0.50 to 1.12; Analysis 2.3) (Cherry 1994). This difference equates to 142 fewer complaints per 1000 participants in the group who received scheduled telephone follow-up. Clinically this might represent a significant difference although this study was underpowered to detect it, hence the wide confidence intervals. Our confidence in the quality of the evidence for the effect of delivery system design interventions on the number of outstanding complaints in the long term is very low.

Secondary outcomes

1. Quality of life

No delivery system design intervention studies reported quality of life as an outcome.

2. Hearing handicap

Two studies measured the effect of delivery system design interventions on short- to medium-term hearing handicap and yielded

data in a form that we were able to combine in a quantitative analysis (Cherry 1994; Collins 2013). Data from these two studies showed no statistically or clinically significant short- to medium-term effect on hearing handicap, as measured using the Hearing Handicap Inventory for the Elderly (Ventry 1982), for delivery system design interventions as a whole (two studies, 628 participants; MD -0.70, 95% CI -5.22 to 3.81; Analysis 2.4). The Cherry 1994 study compared scheduled telephone follow-ups (delivery system design intervention - change in format) with face-to-face follow-up on request (control). The Collins 2013 study compared group fitting and follow-up (delivery system design intervention - change in mode) with individual fitting and follow-up. We found no delivery system design interventions that reported long-term hearing handicap. Our confidence in the quality of the evidence of the effect of delivery system design interventions on self-reported hearing handicap is very low.

3. Hearing aid benefit

A single delivery system design intervention showed no evidence of statistically or clinically significant effect on short- to medium-term hearing aid benefit (one study, 582 participants; MD 1.80, 95% CI -3.10 to 6.70; Analysis 2.5) (Collins 2013). We found no delivery system design studies that reported on long-term hearing aid benefit. Our confidence in the quality of the evidence of the effect of delivery system design interventions on hearing aid benefit is very low.

4. Communication

One delivery system design intervention reported data on communication in the short- to medium-term (Collins 2013). This showed no statistically or clinically significant effect on short- to medium-term use of verbal communication strategies for group versus individual hearing aid fittings (one study, 588 participants; MD -0.10, 95% CI -0.40 to 0.20; Analysis 2.6). We found no delivery system design studies that reported long-term communication outcome. Our confidence in the quality of the evidence of the effect of delivery system design interventions on communication is very low.

Combined self-management support/delivery system design interventions

Primary outcome

1. Hearing aid use

1.1 Adherence

One combined self-management support/delivery system design intervention study reported data on adherence as defined in this review (Ferguson 2016). They reported that at five to eight weeks post fitting no participants given access to remote learning objects post fitting were non-users compared to 5/88 in the control group (one study, 162 participants; risk ratio (RR) 1.06, 95% CI 1.00 to 1.12; Analysis 3.1). This equates, on average, to an additional 57 people out of 1000 wearing their hearing aid up to eight weeks post fitting. We found no studies that reported the effect of combined interventions on adherence in the long term. Our confidence in the quality of the evidence for the effect of combined self-management support/delivery system design interventions on adherence is low.

1.2 Daily hours of hearing aid use

Two combined studies measured daily hours of hearing aid use over the long term (Oberg 2008; Oberg 2009). There was no statistically or clinically significant evidence of overall long-term effect for these combined self-management support/delivery system design interventions (two studies, 69 participants; MD 0.04, 95% CI -0.64 to 0.73; Analysis 3.2). There was some heterogeneity in these data ($I^2 = 55\%$). The studies did not differ in self-management support content, delivery system design format or intensity as we have defined them, so our subgroup analyses failed to explain this heterogeneity. However, the participants in the Oberg 2009 study were able to gain some experience in their own home with an experimental hearing aid prior to fitting rather than only in a clinic setting as they did in the Oberg 2008 study.

Nine of the combined self-management support/delivery system design studies that measured short- to medium-term daily hours of hearing aid use yielded data in a form suitable for meta-analysis (Andersson 1995; Andersson 1997; Ferguson 2016; Kemker 2004; Lundberg 2011; Oberg 2008; Oberg 2009; Thoren 2011; Thoren 2014). There was no statistically or clinically significant evidence of overall short- to medium-term effect on daily hours of hearing aid use (see total in Analysis 3.3, nine studies, 534 participants; MD 0.19, 95% CI -0.01 to 0.40). There were no apparent subgroup differences for self-management support content (Analysis 3.3), delivery system design format (Analysis 3.4) or delivery system design intensity (Analysis 3.5).

The data from two combined self-management support/delivery system design studies could not be combined in the quantitative analysis because we could not obtain either means and/or standard deviations (Eriksson-Mangold 1990; Ward 1978). Data from two further studies could not be combined because they used variants of the same measurement instrument for the intervention and control groups to measure use (Kramer 2005; Olson 2013). This comparison may be invalid and should be interpreted with caution (Laplante-Levesque 2012).

Our confidence in the quality of the evidence of the effect of combined self-management support/delivery system design interventions on daily hours of hearing aid use is very low.

2. Adverse effects

No combined studies reported on clinical adverse events or the number of complaints.

Secondary outcomes

1. Quality of life

Two combined self-management support/delivery system design studies assessed long-term quality of life (Oberg 2008; Oberg 2009). There was no evidence of a statistically significant long-term effect on quality of life for these interventions over and above that provided by the hearing aid itself (two studies, 69 participants; MD 0.32, 95% CI -0.17 to 0.80; Analysis 3.6).

Eight combined self-management support/delivery system design interventions reported short- to medium-term quality of life (Ferguson 2016; Kramer 2005; Lundberg 2011; Oberg 2008; Oberg 2009; Preminger 2010; Thoren 2011; Thoren 2014). Overall there was no evidence of a statistically or clinically significant effect for these combined interventions on short- to medium-term quality of life (eight studies, 530 participants; standardised mean difference (SMD) 0.02, 95% CI -0.15 to 0.19). There were no significant subgroup differences by self-management support content (Analysis 3.7), delivery system design format (Analysis 3.8) or delivery system design intensity (Analysis 3.9).

Our confidence in the quality of the evidence of the effect of combined self-management support/delivery system design interventions on quality of life is low.

2. Hearing handicap

All of the studies reporting long-term hearing handicap were combined self-management support/delivery system design interventions. We were able to combine three of these studies in a meta-analysis, which showed no overall evidence of a statistically significant effect (three studies, 88 participants; SMD -0.31, 95% CI -1.06 to 0.44; Analysis 3.10) (Andersson 1994; Oberg 2008; Oberg 2009). However, there was evidence of heterogeneity in these data. A subgroup analysis by self-management support content suggests that the intervention containing components of psychosocial activation had a greater effect on hearing handicap than the two interventions that aimed to address symptom management skills. The three studies do not differ in delivery system design format or delivery system design intensity. However, we judged the Andersson 1994 study to have a high risk of bias. Based on this evidence, our confidence in the quality of the conclusion that psychosocial self-management support interventions might be more effective than symptom-focused self-management support interventions is very low.

We combined the data from 15 studies that assessed the effect of combined self-management support/delivery system de-

sign interventions on short- to medium-term hearing handicap in meta-analyses (Abrams 1992; Andersson 1995; Andersson 1997; Beynon 1997; Ferguson 2016; Kramer 2005; Kricos 1996; Lundberg 2011; Miranda 2008; Oberg 2008; Oberg 2009; Preminger 2010; Smaldino 1988; Thoren 2011; Thoren 2014) (Analysis 3.11; Analysis 3.12; Analysis 3.13). Overall there was evidence of a statistically significant effect on hearing handicap for these interventions (15 studies, 728 participants; SMD -0.26, 95% CI -0.48 to -0.04). A SMD of this magnitude reflects a small effect size (Cohen 1988). Subgroup analysis by self-management support content shows no significant subgroup differences (Analysis 3.11). Analysing the data by delivery system design format and delivery system design intensity suggests that an intervention involving telephone follow-up was more effective than interventions delivered face-to-face or remotely (Analysis 3.12) and that medium-intensity interventions are more effective than high-intensity (Analysis 3.13). However, a visual inspection suggests within-subgroup heterogeneity in these analyses. The interventions also varied by mode and location of care delivery and it is likely that interaction between these and the other variables is contributing to this heterogeneity. These subgroup analyses should therefore be viewed with caution.

3. Hearing aid benefit

We were able to combine two of the four combined self-management support/delivery system design interventions that assessed long-term hearing aid benefit in a quantitative analysis (Oberg 2008; Oberg 2009). This showed a statistically significant effect for these combined interventions on long-term hearing aid benefit (two studies, 69 participants; MD 0.30, 95% CI 0.02 to 0.58; Analysis 3.14). However, this does not represent a clinically significant difference on this scale (Cox 2002; Smith 2009). Both studies assessed the effect of changes in self-management support content (activate - symptoms versus no intervention) and delivery system design intensity (medium-intensity versus no intervention). We have therefore not performed a subgroup analysis of these data.

In the short to medium term there was no evidence of a statistically or clinically significant effect for combined self-management support/delivery system design interventions (see total in Analysis 3.15; seven studies, 361 participants; SMD 0.10, 95% CI -0.15 to 0.36). There were no apparent significant subgroup differences by self-management support content (Analysis 3.15), delivery system design format (Analysis 3.16) or delivery system design intensity (Analysis 3.17).

Our confidence in the quality of the evidence of the effect of combined self-management support/delivery system design interventions on hearing aid benefit is low.

4. Communication

Only two of the studies reported an overall single score measure of communication (Preminger 2010; Sweetow 2006), but we were

unable to combine these in meta-analyses. The [Sweetow 2006](#) study reported only combined data from both periods of their cross-over study and contact with the authors confirmed that it was not possible to extract the data for the first period of the study separately. The [Preminger 2010](#) study included data on two cochlear implant users and we were not able to separate the data for the hearing aid users only. The remaining studies used the Communication Profile for the Hearing Impaired to measure communication ability ([Demorest 1987](#)), with some choosing to use only the communication strategies subscale of this measure. This measures whether people use verbal, non-verbal and maladaptive strategies for communication. There was evidence of selective reporting in these data, with at least one of the studies reporting data only from scales where significant differences were seen ([Kricos 1996](#)). Only two studies reported effects on long-term communication for combined self-management support/delivery system design interventions ([Chisolm 2004](#); [Oberg 2008](#)). [Chisolm 2004](#) only provided mean scores with no measures of variance so data are only available from the [Oberg 2008](#) study. This showed no evidence of a statistically or clinically significant effect on the use of verbal communication strategies over the long term (one study, 34 participants; MD 0.30, 95% CI -0.20 to 0.80; [Analysis 3.18](#)). A meta-analysis of the four combined self-management support/delivery system design studies reporting short- to medium-term communication outcomes that we were able to combine showed evidence of a statistically significant short- to medium-term effect on the use of verbal communication for these combined self-management support/delivery system design interventions (see total

[Analysis 3.19](#); four studies, 223 participants; MD 0.45, 95% CI 0.15 to 0.74) ([Chisolm 2004](#); [Kricos 1996](#); [Oberg 2008](#); [Turbin 2006](#)). However, this mean difference does not represent a clinically significant difference based on a minimal important difference of 0.93 for this scale ([Demorest 1988](#)). All the studies involved face-to-face delivery and there were no significant subgroup differences by self-management support content ([Analysis 3.19](#)) and delivery system design intensity ([Analysis 3.20](#)). Our confidence in the quality of the evidence of the effect of combined interventions on communication is very low.

Decision support interventions

We found no studies that investigated the potential effects of decision support interventions.

Clinical information system interventions

We found no studies that investigated the potential effects of clinical information system interventions.

Community interventions

We found no studies that investigated the potential effects of community interventions.

Health system interventions

We found no studies that investigated the potential effects of health system interventions.

ADDITIONAL SUMMARY OF FINDINGS *[Explanation]*

Delivery system design interventions for adults with hearing loss who use hearing aids						
Patient or population: adults with hearing loss who use hearing aids Settings: outpatient clinic Intervention: delivery system design interventions						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	Control	Delivery system design interventions				
Adherence Number of people fitted with hearing aid/number of people who use the aids Follow-up: 0 to 12 months	948 per 1000	967 per 1000 (938 to 995)	RR 1.02 (0.99 to 1.05)	686 (2 studies)	⊕⊕○○ low ¹	-
Daily hours of hearing aid use Average self-reported or data-logged hours of use per day. Scale from: 0 to 12 hours Follow-up: 0 to 12 months	The mean daily hours of hearing aid use in the intervention groups was 0.06 lower (1.06 lower to 0.95 higher) than in the control groups. On average the intervention groups used their hearing aids for under a minute per day less than the control groups		-	700 (4 studies)	⊕○○○ very low ²	Participants in the intervention groups wore their hearing aids for 3 to 4 minutes less each day on average than those in the control group. This is not a clinically significant difference
Adverse effects Number of outstanding complaints Follow-up: 1+ years	571 per 1000	429 per 1000 (286 to 640)	RR 0.75 (0.5 to 1.12)	98 (1 study)	⊕○○○ very low ³	-

Quality of life	No studies identified				
Self-reported hearing handicap Validated self-report measure HHIE scale from: 0 to 100 (Ventry 1982) Follow-up: 0 to 12 months	The mean self-reported hearing handicap in the intervention groups was 0.7 lower (5.22 lower to 3.81 higher) than in the control groups (on this scale from 0 to 100, a lower score indicates less hearing handicap)	-	628 (2 studies)	⊕○○○ very low ⁴	The minimal important difference on this scale is reported to be 18.7 for face-to-face administration and 36 for pencil and paper (Weinstein 1986)
Hearing aid benefit Validated self-report measure. Outer EAR scale from: 0 to 100 Follow-up: mean 6 months	The mean hearing aid benefit in the intervention group was 1.8 higher (3.1 lower to 6.7 higher) than in the control group (on this scale from 0 to 100, a higher score indicates more hearing aid benefit)	-	582 (1 study)	⊕○○○ very low ⁴	While we were unable to reference a minimal important difference for this scale, a mean difference of 1.8 on a scale from 0 to 100 is unlikely to be a clinically significant change
Communication Validated self-report measure: verbal subscale of the CPHI scale from 0 to 5 (Demorest 1987) Follow-up: 0 to 12 months	The mean reported use of verbal communication strategy in the intervention group was 0.10 higher (0.40 lower to 0.20 higher) than in the control group (higher score indicates increased use of verbal communication strategy)	-	588 (1 study)	⊕○○○ very low ⁵	The minimal important difference for this subscale of the CPHI is 0.93 at the 0.05 level (Demorest 1988)

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (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; CPHI: Communication Profile for the Hearing Impaired; HHIE: Hearing Handicap Inventory for the Elderly; RR: risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

¹Downgraded due to very serious concerns regarding indirectness of the evidence (only short- to medium-term evidence and the majority of the participants were military veterans).

²Downgraded due to very serious concerns regarding indirectness (short- to medium-term data and military veteran participants) and serious concerns about limitations in study design (unclear risk of bias) and imprecision (standard deviations imputed in the largest study).

³Downgraded due to very serious concerns regarding indirectness (short- to medium-term data and military veteran participants) and serious concerns regarding limitations in study design (unclear risk of bias) and imprecision (small sample size, wide CIs).

⁴Downgraded due to very serious concerns regarding indirectness (short- to medium-term data and military veteran participants) and serious concerns about imprecision (standard deviations imputed).

⁵Downgraded due to very serious concerns regarding indirectness (short- to medium-term outcomes, military veteran participants and the lack of a global communication outcome measure) and serious concerns about imprecision (standard deviations imputed).

Combined self-management support/delivery system design interventions for adults with hearing loss who use hearing aids						
Patient or population: adults with hearing loss who use hearing aids Settings: outpatient clinic Intervention: combined self-management support/delivery system design interventions						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	Control	Combined SMS/DSD interventions				
Adherence Number of people fitted with hearing aid/number of people who use the aids Follow-up: 5 to 8 weeks	943 per 1000	1000 per 1000 (943 to 1000)	RR 1.06 (1 to 1.12)	162 (1 study)	⊕⊕○○ low ¹	-
Daily hours of hearing aid use Self-reported or data-logged average hours of use per day. Scale from: 0 to 12 hours Follow-up: 1+ years	The mean daily hours of hearing aid use in the intervention groups was 0.04 higher (0.64 lower to 0.73 higher) than in the control groups		-	69 (2 studies)	⊕○○○ very low ²	Participants in the intervention groups wore their hearing aids for 2 to 3 minutes more per day than those in the control group. This is not a clinically significant difference
Adverse effects	No studies identified					
Quality of life Validated self-report measures. IOI-HA item 7 scale from: 1 to 5 Follow-up: 1+ years	The mean quality of life in the intervention groups was 0.32 higher (0.17 lower to 0.8 higher) than in the control groups, measured on item 7 of the IOI-HA (Cox 2002)		-	69 (2 studies)	⊕⊕○○ low ³	The minimally important difference for this subscale of the IOI-HA is 0.32 for those with mild-moderate hearing

					loss and 0.28 for those with moderate-severe hearing loss (Smith 2009).
Self-reported hearing handicap Validated self-report measures Follow-up: 1+ years	The mean self-reported hearing handicap in the intervention groups was 0.31 standard deviations lower (1.06 lower to 0.44 higher) than in the control groups	-	88 (3 studies)	⊕○○○ very low ⁴	Using the classification suggested by Cohen 1988 a SMD of 0.31 represents a moderate effect size.
Hearing aid benefit Validated self-report measures (IOI-HA item 4). Scale from: 1 to 5 Follow-up: 1+ years	The mean hearing aid benefit in the intervention groups was 0.3 higher (0.02 to 0.58 higher) than in the control groups, measured on item 4 of the IOI-HA (Cox 2002)	-	69 (2 studies)	⊕⊕○○ low ³	This is a statistically significant difference. However, the minimally important difference for this subscale of the IOI-HA is 0.39 for those with mild-moderate hearing loss and 0.32 for those with moderate-severe hearing loss (Smith 2009), so this does not represent a clinically important difference
Use of verbal communication strategy Validated self-report measures (verbal subscale of the CPHI (Demorest 1987)). Scale from: 0 to 5 Follow-up: 1+ years	The mean use of verbal communication strategy in the intervention groups was 0.3 higher (0.2 lower to 0.8 higher) than in the control groups	-	34 (1 study)	⊕○○○ very low ⁵	The minimal important difference for this subscale of the CPHI is 0.93 at the 0.05 level (Demorest 1988).

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (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; **CPHI:** Communication Profile for the Hearing Impaired; **DSD:** delivery system design; **IOI-HA:** International Outcome Inventory for Hearing Aids; **RR:** risk ratio; **SMD:** standardised mean difference; **SMS:** self-management support

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

¹Downgraded due to serious concerns regarding risk of bias and serious concern regarding consistency (single study).

²Downgraded due to very serious concerns regarding imprecision (small sample size) and serious concerns regarding inconsistency (heterogeneity).

³Downgraded due to very serious concerns regarding imprecision (small sample size).

⁴Downgraded due to very serious concerns regarding imprecision (small sample size, risk of skewed data in two of the studies) and serious concerns regarding limitations in study design (high risk of bias in one study) and inconsistency (heterogeneity).

⁵Downgraded due to very serious concerns regarding imprecision (small sample size) and indirectness (lack of a global measure of communication, participants were all first-time hearing aid users, we do not know whether equivalent benefit could be gained in people already fitted with hearing aids).

DISCUSSION

Summary of main results

We reviewed the range, nature and long-term effects of any intervention supplementing, but not including, hearing aid fitting that had the aim of improving or encouraging hearing aid use in adult auditory rehabilitation.

All the studies we identified could be classified using the chronic care model (CCM) as self-management support and/or delivery system design interventions.

We found no self-management support studies that investigated the effect of self-management support on adherence, adverse effects or hearing aid benefit. Two studies reported daily hours of hearing aid use but we were unable to combine these in a meta-analysis. There was no evidence of a statistically significant effect on quality of life over the short to medium term (one study, 35 participants; mean difference (MD) -9.10, 95% confidence interval (CI) -21.33 to 3.13). Self-management support interventions reduce short- to medium-term hearing handicap (two studies, 87 participants; MD -12.80, 95% CI -23.11 to -2.48) and increase the use of verbal communication strategies in the short to medium term (one study, 52 participants; MD 0.72, 95% CI 0.21 to 1.23). The clinical significance of these statistical findings is open to question but, based on the minimal important differences on the scales used, it is likely that the outcomes were clinically significant for some, but not all, participants. Our confidence in the quality of this evidence was very low. No self-management support studies reported long-term outcomes (see [Summary of findings for the main comparison](#)).

Delivery system design interventions did not significantly effect adherence (two studies, 686 participants; risk ratio (RR) 1.02, 95% CI 0.99 to 1.05) or daily hours of hearing aid use (four studies, 700 participants; MD -0.06, 95% CI -1.06 to 0.95) in the short to medium term or adverse effects in the long term (one study, 98 participants; RR 0.75, 95% CI 0.50 to 1.12). We found no studies that investigated the effect of delivery system design changes on quality of life. There was no evidence of a statistically or clinically significant effect on hearing handicap (two studies, 628 participants; MD -0.70, 95% CI -5.22 to 3.81), hearing aid benefit (one study, 582 participants; MD 1.80, 95% CI -3.10 to 6.70) or the use of verbal communication strategies (one study, 588 participants, MD -0.10, 95% CI -0.40 to 0.20) in the short to medium term. Our confidence in the quality of this evidence was low or very low. Long-term outcome measurement was rare in delivery system design comparisons (see [Summary of findings 2](#)).

We found no studies that investigated the effect of complex interventions combining components of self-management support and delivery system redesign on adverse effects. A single study showed a probable effect on adherence in the short term (one study, 162 participants; risk ratio (RR) 1.06, 95% CI 1.00 to 1.12). There

was no evidence of a statistically or clinically significant effect on daily hours of hearing aid use over the long term (two studies, 69 participants; MD 0.04, 95% CI -0.64 to 0.73) or short to medium term (nine studies, 534 participants; MD 0.19, 95% CI -0.01 to 0.40). Similarly, there was no evidence of an effect on quality of life over the long term (two studies, 69 participants; MD 0.32, 95% CI -0.17 to 0.80) or short to medium term (eight studies, 530 participants; standardised mean difference (SMD) 0.02, 95% CI -0.15 to 0.19). Combined interventions reduced hearing handicap in the short to medium term (15 studies, 728 participants; SMD -0.26, 95% CI -0.48 to -0.04). This represents a small to moderate effect size but there is no evidence of a statistically significant effect over the long term (three studies, 88 participants; SMD -0.31, 95% CI -1.06 to 0.44). There was evidence of a statistically, but not clinically, significant effect on long-term hearing aid benefit (two studies, 69 participants; MD 0.30, 95% CI 0.02 to 0.58), but no evidence of an effect over the short to medium term (seven studies, 361 participants; SMD 0.10, 95% CI -0.15 to 0.36). There was evidence of a statistically, but not clinically, significant effect on the use of verbal communication strategies in the short term (four studies, 223 participants; MD 0.45, 95% CI 0.15 to 0.74), but not the long term (one study, 34 participants; MD 0.30, 95% CI -0.20 to 0.80). Our confidence in the quality of this evidence was low or very low (see [Summary of findings 3](#)). There were no studies investigating the effect of decision support, the use of clinical information systems, community resources or health system changes.

Overall completeness and applicability of evidence

Completeness

In terms of interventions, although we were able to identify 37 studies for inclusion in this review we classified all of them as delivery system design and/or self-management support interventions. There was a lack of randomised controlled trial (RCT) evidence looking at decision support, clinical information systems, community support or health system changes. Some components of delivery system design and self-management support have also not been fully explored. For example, there were no studies that specifically addressed the effect of supporting hearing aid use with the provision of additional services such as battery replacement services and the provision of, or referral for, additional equipment to improve hearing aid benefit. No studies involved the explicit use of collaborative goal-setting or action-planning and the patients' central role in managing their own health was often not explicitly acknowledged. These are central tenets of a self-management support approach. There has been relatively little focus on low-intensity interventions and no studies that consider the reorganisation of staff roles. Using a framework such as the CCM has

helped to highlight considerable gaps in the evidence base in terms of interventions that have been tested in RCTs in this context. The majority of interventions we found included components of both delivery system design and self-management support. This is consistent with the results of the review by Tsai 2005 for long-term conditions.

The CCM and other similar frameworks are general so that they can be applied in any healthcare context. This can mean that some of the detail about what works and what does not work can be lost. We hoped that the subgroup analyses might provide a useful model to explore what components of interventions may be most effective in changing particular outcomes. In the majority of cases our subgroup analyses did not help to answer this question. Partly this was due to a lack of data. Even for those comparisons and outcomes where we had more data (e.g. Analysis 3.13), there was a lack of data in some groups but not others, which makes a valid assessment of subgroup differences difficult. In addition, we did not analyse our data using other subgroups suggested by Barlow 2002, such as target population or delivery location. One possible avenue may be to explore not only the purpose of participant activation (addressing practical, symptom-management and psychosocial management skills) as we have done, but also the depth of activation. We would welcome a discussion on the potential viability of comparing interventions in which the minimum level of participant engagement is attendance, those that require some engagement in practical activities under the direct supervision of clinicians in a clinic-based environment and those which engage the participants in collaboratively agreed 'homework' under indirect professional guidance with appropriate follow-up. It would also be interesting to look more closely at the self-management support interventions and analyse the content in terms of behaviour change technique employed, for example using the behaviour change technique taxonomy (version 1) (Michie 2013). Combining this with a meta-regression could yield a more granular understanding of the relative contribution of different active ingredients of an intervention.

In terms of the primary outcomes, there was a relative lack of data on adherence and adverse effects. The problem of hearing aid non-use is always stated in terms of adherence (or lack of it) and so it is disappointing that few studies chose to report the outcome of their interventions in this way. It makes it difficult to relate the results of the studies back to the original problem. Adherence, using the definition we have adopted from the World Health Organization, implies a level of agreement with the chosen management option. For the purposes of this review we assumed that a hearing aid fitting was the agreed course of action. However, the level of collaboration between patient and clinician was not mentioned explicitly in any of the studies we included, something highlighted in our subgroup analyses by self-management support content. It is possible that the included studies were therefore measuring compliance rather than adherence as we have defined it. When studies did consider hearing aid use it was usually measured as self-reported hours of

use per day. It was rare for studies to make any mention of the potential for adverse effects, which is a limitation in study design and outcome measurement to date.

Patient-reported secondary outcomes were measured with a variety of metrics even for the same outcome. Hearing loss has complex consequences and the measurement of outcome is therefore complex (Granberg 2014). There is a lack of consensus over which outcomes are important in hearing health and a lack of agreement on which specific scales should be used to measure those outcomes (Hanratty 2000; Humes 2011). This diversity was reflected in this review and made meta-analysis for some of the outcomes difficult. Our results suggest that any positive outcomes due to changes in the way care is delivered are small and incremental compared to the benefits of the hearing aid itself. All of the scales used in our meta-analyses had minimal important differences of approximately 20% of the total scale score. This means that interventions would need to produce average mean differences of that magnitude to be considered clinically significant. Studies aiming to measure these potential incremental benefits should bear this in mind in their choice of patient-reported outcome measure.

This review also highlights the need for further studies that consider long-term outcomes over a year or more. It is of great importance to know whether a particular intervention has lasting effects over the long term, especially in the context of managing a long-term condition. It is not safe to assume that short-term positive outcomes translate into the long term. A patient may persevere with hearing aid use while they are still receiving relatively intense support from their clinician but then lapse when they are left to self-manage their condition over the longer term. However, the reverse may also be true for some outcomes. Positive outcomes may not been seen in the short term but may only be evident in the long term once participants have had the benefit of extended practice and experience.

The number of studies that provided data in a form that could be included in a meta-analysis was relatively low. This is not unusual in systematic reviews (Johnston 2013), but is not something to be applauded. Sometimes raw data were not available, with only the overall conclusion being reported in the paper, and sometimes particular figures such as standard deviations or other measures of data spread were missing. A significant amount of data could not be combined or had high standard deviations relative to the means and therefore carried a high risk of skew. This variability in the data highlights the need to include a priori estimates of effect size so that studies are appropriately statistically powered.

Applicability

All of the studies identified were carried out in countries with well-developed health systems; this limits the applicability of the findings beyond such systems.

Some of the studies involved the use of veterans as participants (Abrams 1992; Chisolm 2004; Collins 2013; Kemker 2004; Preminger 2010a; Saunders 2016; Turbin 2006). While in terms

of study numbers these were a minority, in terms of participants they represented almost a third of the total (1297/4129 participants). This weights the results towards a largely male, highly motivated population, which limits the generalisability of the findings to the non-military population, a limitation acknowledged in most of these studies.

The studies all had specific inclusion/exclusion criteria that often meant that people with comorbid conditions were excluded. Some had age restrictions either at the younger or older end, sometimes both. Again this limits the generalisability of the findings.

There were no large-scale effectiveness trials conducted in unselected populations.

Quality of the evidence

Twelve studies have publication dates from 2001 to 2009 but none referenced the CONSORT 2001 guidelines. A further 13 studies were published after 2010 and the updated CONSORT guidelines (CONSORT 2010), but only four referenced the updated guidance (Ferguson 2016; Saunders 2016; Thoren 2011; Thoren 2014).

The studies were of variable methodological rigour (see Figure 3) and many of them did not report raw data or reported data in such a way that they could not be included in a meta-analysis. There was a diversity of outcome metrics, which sometimes made comparisons between studies difficult.

We have assessed the results for the primary and secondary outcomes using GRADE protocols and the results are included in the 'Summary of findings' tables (Summary of findings for the main comparison; Summary of findings 2; Summary of findings 3). For self-management support interventions, delivery system design interventions and combined interventions we judged the limited evidence to be of very low to low quality against the GRADE criteria (Higgins 2011). Where evidence was downgraded this was due limitations in study design (high or unclear risk of bias across studies for a particular outcome), indirectness (in terms of population and outcome measurement) and imprecision (small sample sizes, large confidence intervals, high risk of skewed data).

Potential biases in the review process

We cannot exclude the possibility that other studies have been published showing positive or negative results, which have not been included here, but we are confident that the extensive electronic search and subsequent reference checking has captured most of the relevant literature. However, we invite readers to notify us of any trials or studies we may have missed so that they might be included in subsequent updates to this review. Similarly, although we did make efforts to contact study authors directly to clarify study methods and obtain raw data where possible, we were not always able to do so. We would very much welcome contact from

any of the authors cited in this review who feel that their data could be included in the meta-analysis in future updates.

Some of the studies were excluded on the basis that their outcomes did not meet the inclusion criteria for this review. As far as we are aware, based on an assessment of the methods sections of these studies, the relevant outcomes were not available because they were not measured. However, it is possible that other outcomes were measured and not reported. We invite the authors of these studies to contact us if additional outcomes, which could be included in future updates, were measured but not reported.

Two review authors independently selected trials, extracted data, assessed risk of bias and graded the quality of evidence in order to minimise bias in the review process.

None of the review authors had any involvement in any of the trials. This has not been the case in previous systematic reviews in the context of hearing healthcare (Chisolm 2011; Sweetow 2005).

Agreements and disagreements with other studies or reviews

Like Barlow 2002, Kreindler 2009 and Tsai 2005, we found in our review that many interventions were a complex combination of delivery system design and self-management support components. Developing the skills necessary to become a self-manager of a long-term condition requires information and support for behaviour change to deal with the symptom, physical and psychosocial consequences of the condition (Barlow 2002; Lorig 2003; Pearson 2007). We feel that the CCM has been a useful starting framework within which to separate out the possible effects of different aspects of complex interventions, e.g. components of self-management support and components of delivery system design. In their review of self-management approaches for people with long-term conditions, Barlow 2002 sought to identify approaches to self-management and to consider the effectiveness of these approaches. Of the 145 studies they identified, only one looked at a sensory problem: tinnitus (Jakes 1986). The results of our review suggest that many of the studies we identified could be included if the Barlow 2002 review were to be updated. Barlow 2002 found that self-management support interventions rarely target carers. In our review many of the studies included content addressing communication, which is necessarily a two-way process, but only one specifically addressed the effect of explicit involvement of significant others or communication partners (Preminger 2010). In the Barlow 2002 review, approximately half the studies were RCTs but with small sample sizes (20 to 30) and short follow-up periods (four to six months). They called, like us, for RCTs of sufficient power to enable change to be detected and for longer-term follow-up.

In the context of hearing health care, previous reviews have tended to concentrate on specific intervention types, such as auditory training or changes in delivery such as group versus individual delivery.

A previous systematic review conducted by [Sweetow 2005](#), subsequently updated to include a meta-analysis by [Chisolm 2011](#), addressed the evidence for individual auditory training. This type of intervention involves the patient participating in a programme of training designed to enhance speech perception. Training is typically provided on a repeated basis over a number of sessions and involves practice with listening and recognition of speech-based material. The speech-based training material may be broken down into its constituent parts with the aim of improving the discrimination and recognition of those parts (analytic training), or presented in sentence-length structures with the aim of improving listening skill and overall comprehension (synthetic training). Both the original [Sweetow 2005](#) review and the [Chisolm 2011](#) update included RCTs but also cohort and before/after study designs where participants may act as their own controls. [Sweetow 2005](#) identified six studies for inclusion with four studies being added in the [Chisolm 2011](#) update. Six of these 10 studies were RCTs and three of these were included in our review also ([Kricos 1992](#); [Kricos 1996](#); [Sweetow 2006](#)). The three other RCTs were excluded from this review because their only reported outcome was speech perception (see [Characteristics of excluded studies](#)). They concluded that there was evidence of improvement in speech perception in adults with hearing loss who undertake auditory training at least in the short term (i.e. immediately after training). A review by [Hawkins 2005](#) (and subsequently updated by [Chisolm 2011](#)) assessed the evidence for counselling-based group auditory rehabilitation programmes. They looked at effects on short- and long-term self-perceived benefits, satisfaction or both. Like the [Sweetow 2005](#) review they did not limit inclusion to RCTs. [Hawkins 2005](#) highlighted the need for further well-controlled studies, with adequate numbers of participants, given the variability evident in the reviewed studies. We would echo this call. [Chisolm 2011](#) updated this review, focusing particularly on RCTs but also including studies that included people who were not using hearing aids. They identified 10 studies, seven of which also met the inclusion criteria for this review (two were excluded as they included non-hearing aid users ([Hallberg 1994](#); [Hickson 2007](#)), and one was excluded as it was a second paper on the same set of participants as an already included study ([Chisolm 2004](#))). [Chisolm 2011](#) conducted a meta-analysis looking at hearing handicap as an outcome. They found a small but significant effect of group auditory rehabilitation on short-term hearing handicap. However, their analysis did include some double-counting, with the participants in [Chisolm 2004](#) counted twice and the control participants for [Preminger 2010](#) and [Smaldino 1988](#) counted three times. They highlighted the variability present in their data but did not investigate possible reasons for the apparent heterogeneity. A major weakness of both of these reviews is that they do not consider interactions between the content and delivery of interventions and comparisons. Auditory training is typically delivered over many sessions and would therefore constitute a high-intensity intervention as we have defined it, but it is often compared

with standard care which is low- or medium-intensity. It is rare for auditory training studies to control for this, although [Kricos 1996](#), [Fitzpatrick 2008](#) and [Saunders 2016](#) did do this and have hence been defined in our review as self-management support interventions. They therefore provide more robust evidence on the effect of changing the content of an intervention. We found a similar issue when comparing group interventions versus individual interventions. Studies often do not control for variations in what is delivered between intervention and control groups. The one study we found that did control for content showed no significant difference in hearing handicap between group and individual delivery mode when the same content was delivered to both ([Collins 2013](#)).

A second weakness in both of these reviews is a lack of acknowledgement or assessment of risk of bias and other factors impacting on confidence in the quality of the evidence as recommended in GRADE protocols.

Using the CCM and work by [Barlow 2002](#) and [Pearson 2007](#) as a framework for this review has demonstrated clearly that most interventions in hearing health care are a complex mix of self-management support and delivery system design changes. Using this framework we have attempted to identify some of the potential active components of these complex interventions. While we have been only partially successful we have at least highlighted that this issue exists. Careful delineation of the different factors that may have an impact on outcome for these complex interventions is essential in drawing conclusions when reviews are undertaken or updated in future.

AUTHORS' CONCLUSIONS

Implications for practice

There is some low to very low quality evidence to support the use of self-management support and complex interventions combining components of self-management support and delivery system design in hearing health care. However, the range of interventions that have been tested is relatively narrow. Data on long-term outcomes are sparse.

Implications for research

There are many opportunities for further research in this context. The design and funding of future research should include a focus on investigating long-term outcomes. This has also been highlighted in other systematic reviews ([Barlow 2002](#); [Chisolm 2011](#); [Hawkins 2005](#); [Sweetow 2005](#)), as has the need for larger, appropriately powered studies in this context.

Using the chronic care model (CCM) and the literature on self-management support and its delivery as a theoretical backbone for this review has highlighted gaps in the evidence base, particularly

in the elements of decision support, clinical information systems, health system and community-based interventions, where there is a total lack of high-level evidence. Some specific intervention types have received more attention, such as educative, counselling-based self-management support and auditory training. However, the implementation of these interventions frequently also necessitates changes in delivery system design. The interaction between these two elements is rarely explicitly explored in hearing health care research. In future it would be helpful if researchers clearly delineate and describe the potentially active components of their interventions and use mixed methods to investigate the relative contribution of different components of any intervention. Even within the CCM elements where data are available relatively little research has looked at explicitly engaging the patient as an active participant in their own rehabilitation. Collaborative goal-setting and problem-solving is an area that would benefit from further investigation.

In relation to the primary outcome in this review it would be helpful to see more studies consider behavioural outcomes such as hearing aid use in terms of adherence rather than hours of use per day. Careful consideration needs to be given to the definition of adherence used. As defined in this review it acts both as a behavioural outcome but also brings in a need to acknowledge explicitly collaborative goal-setting in intervention study design. Otherwise studies may choose to measure a purely behavioural outcome (is the patient wearing their hearing aid?) or compliance (is the patient wearing their hearing aid as recommended?). It would be useful to supplement self-report data on hearing aid use (either defined as adherence or use in hours per day) with data-logging. More recent studies are starting to do this. Although data-logging is not a perfect measure of actual behaviour it can act to triangulate purely self-reported results. Previous authors have called for more standardisation in the way that hearing aid use is assessed and categorised (Perez 2012). Researchers should also be alert to the possibility of adverse effects of interventions.

A wide variety of patient-reported outcomes measures were reported in this review. It would be beneficial, in terms of combining study results and comparing interventions, to agree a set of core outcomes for future research into auditory rehabilitation, both in terms of outcome type (e.g. benefit, hearing handicap,

quality of life etc.) and in the measure used to record that outcome. Agreed measures of outcome would also allow mean differences rather than standardised mean differences to be used, which will make it easier to convert results back into meaningful changes on the relevant scales. This will make results easier to interpret and relate back to clinical practice using minimal important differences where available. Measures used for patient-reported outcomes should be sensitive enough to detect incremental changes in outcome over and above those provided by a hearing aid.

This review has highlighted the variable risk of bias in studies to date (Figure 3). Although performance bias is difficult to remove or control for in studies of this type, it is possible to do a better job with detection bias (blinding of outcome assessment) and this would significantly reduce the risk of bias in many of these studies (see, for example, Hickson 2007). Studies should include a better description of the randomisation procedure to allow an accurate assessment of the risk of selection bias to be made. Wider publication of study protocols would allow a clearer assessment of publication bias.

It should be a priority for future randomised controlled trials to cite and adhere to the CONSORT guidelines (CONSORT 2010), something that has been largely lacking in the evidence thus far.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Abrams 1992

Methods	Randomised trial (also had a control group with no intervention but control group inclusion was determined by eligibility for VA-funded HA so not randomised)
Participants	N = 22 in randomised groups Age: 55 and over, PTA 4 frequency average > 30 dB HL in better ear, no previous HA use, women not excluded but none in study Excluded known neurological deficiencies
Interventions	HA + AR group programme versus HA alone AR programme was 90-minute group session once a week for 3 weeks post-fitting
Outcomes	Short-term: baseline and 2 months HANDICAP Hearing Handicap Inventory for the Elderly (HHIE) total score, emotional subscale and social subscale
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "those who received hearing aids were randomly assigned to one of the two treatment groups" Comment: no details given about how sequence was generated Control group not randomised
Allocation concealment (selection bias)	Unclear risk	Comment: no information about how sequence generated or whether it was concealed
Blinding of participants and personnel (performance bias) All outcomes	High risk	Comment: participants not blinded (due to the nature of the intervention) and experimenters almost certainly not, but it was not explicitly stated in the text
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: no missing data

Abrams 1992 (Continued)

Selective reporting (reporting bias)	Unclear risk	No study protocol available
Other bias	Low risk	Study appears to be free of other sources of bias

Andersson 1994

Methods	Randomised
Participants	N = 20 Age: range 64 to 72 11 male, 9 female “Recently” retired, existing HA users (mean duration 2.8 years)
Interventions	HA alone versus HA + AR AR: 60-minute behavioural counselling session over 3 consecutive weeks with homework tasks - could be group, individual or combined depending on functional analysis and practical considerations
Outcomes	Short-term: baseline and 4 weeks later (post AR) - Life Orientation Test (PSYCHOLOGICAL/OPTIMISM) Long-term: 15 months post-intervention - Hearing Coping Assessment (HANDICAP/DISABILITY)
Notes	Could not include Hearing Questionnaire developed by the authors for this study and post-counselling questions also developed for this study Have included Hearing Coping Assessment as it was separately validated (although by the same authors)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: “subjects were randomly assigned to one of two groups” Comment: no details of randomisation process given
Allocation concealment (selection bias)	Unclear risk	Comment: no details of randomisation process
Blinding of participants and personnel (performance bias) All outcomes	High risk	Comment: not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	Comment: authors comment in discussion on the potential effect on non-blinding

Andersson 1994 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: 1 patient in the intervention group not reached at long-term FU - not clear whether results for long-term FU analysed on an intention-to-treat basis but only 1 patient lost
Selective reporting (reporting bias)	High risk	Comment: no protocol available Also, in a later 1998 paper they describe how HA use was measured in this study but not reported
Other bias	Unclear risk	2 of the outcome measures in the study could not be used as they were developed specifically for this study by the authors

Andersson 1995

Methods	Randomised after initial interview and video session
Participants	N = 24 Age: range 64 to 72 (mean 69.8) 14 male, 10 female Recently retired HA users
Interventions	HA alone versus HA + group AR AR consisted of 4 x 2-hour sessions including video feedback, applied relaxation, information and homework
Outcomes	Short-term: baseline and post-intervention (5 weeks) - USE of aid (hours/day), VAS scores for daily hearing problems, Hearing Coping Assessment (HANDICAP/DISABILITY) Long-term: 2-year follow-up - Hearing Coping Assessment (HANDICAP/DISABILITY), Communication Profile Hearing Impaired-Communication Strategy Subscale (COMMUNICATION)
Notes	4 patients lost to long-term follow-up - 2 in each group

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "after which a code was broken and subjects were assigned to respective groups" Comment: probably done
Allocation concealment (selection bias)	Unclear risk	Comment: probably done
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants were not blinded due to the nature of the intervention

Andersson 1995 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Interviewers appear to have been blinded to group allocation but these data were not included in our outcomes
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing data in the original phase of the study and number/reasons for dropout given in follow-up paper
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	The HCA was developed and validated by the author In a follow-up study, after dropouts, the 2 groups differed at baseline on HCA score

Andersson 1997

Methods	Randomised
Participants	N = 19 Age: range 67 to 75, mean 71.5 11 male, 8 female Inclusion criteria: HA users, 65 to 80 years old, able to use telephone Exclusion: previous attendance at a rehabilitation course at the centre, severe tinnitus or vestibular symptoms
Interventions	HA alone versus HA + self-help manual supplied with 1-hour face-to-face training session including relaxation training followed by telephone contacts during 4 consecutive weeks
Outcomes	Short-term: post-intervention: USE hours/day, Hearing Coping Assessment (HANDI-CAP), VAS scores, Communication Profile Hearing Impaired-Communication Strategy Subscale (COMMUNICATION)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "following the structured interview a code was broken and they were assigned to the groups" Comment: probably done
Allocation concealment (selection bias)	Unclear risk	Comment: probably done
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention

Andersson 1997 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Single-blinded interviewer at FU - blind to allocation BUT cannot use these outcomes
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Low risk	Study appears to be free of other sources of bias

Beynon 1997

Methods	Quasi-randomised using last digit of hospital number
Participants	N = 53 randomised but data analysed from 47 after dropouts Age: range 47 to 80 20 male, 27 female Inclusion criteria: first-time HA users, patients had to attend 3 out of 4 intervention sessions Exclusion criteria: over 80 years old, severe or profound hearing loss
Interventions	HA alone versus HA + AR group course AR course: 4 weekly sessions, 5 to 7 people, not clear how long the sessions were
Outcomes	Medium-term: 13 weeks - QDS (HANDICAP)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Comment: allocation by odd or even hospital record number so quasi-randomised
Allocation concealment (selection bias)	Unclear risk	Allocation by hospital number, which presumably investigators knew in advance
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding of outcome measurement

Beynon 1997 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Some missing data but reasons given. Post hoc analysis with imputed data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Low risk	Study appears to be free of other sources of bias

Campos 2013

Methods	Stratified randomised controlled trial. Stratified by age, hearing loss and HA features prior to randomisation	
Participants	N = Tx: 25, Cx: 25 Age range: 39 to 88 years Gender: 30 men, 20 women Inclusion criteria: bilateral mild to severe SNHL Exclusion criteria: no associated disabilities and no previous HA use	
Interventions	Cx - face-to-face consultation for HA fitting Tx - synchronous teleconsultation with facilitator present	
Outcomes	Short-term: 1 month post-intervention - time taken for HA programming and instruction (RESOURCE USE), daily hours of HA use as measured by data-logging and self-report on IOI-HA (USE), HINT-Brazil (SPEECH PERCEPTION)	
Notes	No raw data quoted for IOI-HA. The figures are only available in graph form and standard deviations were not displayed	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote "an equal number of participants from each stratum were allocated to experimental or control groups by a simple raffle"
Allocation concealment (selection bias)	Unclear risk	Comment - no details of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Evaluator was blinded

Campos 2013 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	There were dropouts but the authors explained and accounted for these as far as possible
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	Quote: "It must be emphasized that three participants in the experimental group failed to perform the evaluation of speech perception in quiet and in noise, despite the various attempts made by the evaluator. Thus, the values of the SRT and the S/N ratio of these participants were not included in calculating the average, which may have contributed to the results of the experimental group being more favorable (lower values)"

Cherry 1994

Methods	Randomised
Participants	N = 60 Age: range not given but all over 50 years Gender: not specified Inclusion criteria: 50 years old or over, unaided speech recognition thresholds no greater than 70 dB HL in the aided ear, agreement to buy a HA and kept them at the end of the trial period, mix of new and previous HA users
Interventions	Standard HA fitting versus HA fitting plus scheduled telephone contact post-fitting on 3 occasions
Outcomes	Medium-term: 4 months - USE hours/month, HHIE (HANDICAP) Long-term: 12 months - number of complaints
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly assigned" Comment: no details of randomisation process
Allocation concealment (selection bias)	Unclear risk	Comment: no details of randomisation process

Cherry 1994 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants and personnel not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: there was a dropout rate for the interview and questionnaire that was not completely addressed. Results were not analysed on an intention-to-treat basis but there was a similar dropout in both groups, although reasons are not clear so not sure whether they would be relevant
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Low risk	The study appears to be free of other sources of bias

Chisolm 2004

Methods	Randomised VA funded May 1999 to December 2001
Participants	N = 106 Age: range not given - average approximately 75 years 68 male, 38 female Inclusion criteria: US veterans, new HA users Exclusion criteria: more than mild depression on Beck Depression Inventory
Interventions	HA alone versus HA + AR AR = 4-week group programme, 2 hours once a week
Outcomes	Short-term: 8 weeks - CPHI (HANDICAP and COMMUNICATION), SF-36V (QUALITY OF LIFE)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly assigned" Comment: no details of randomisation process given
Allocation concealment (selection bias)	Unclear risk	Comment: no details given

Chisolm 2004 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants and personnel not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding in measurement of outcome
Incomplete outcome data (attrition bias) All outcomes	Low risk	Missing data at long-term FU but this was accounted for statistically
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Low risk	Study appears to be free of other sources of bias

Collins 2013

Methods	Cluster-randomised after enrolment VA provided HAs free of charge and participants paid USD 50 if they completed all the questionnaires February 2006 to October 2007
Participants	N = 659 randomised but results based on 644 who completed the study Age: range 23 to 93 years, mean 65.5 98.5% male Inclusion criteria: no previous HA use Exclusion criteria: unable/unwilling to participate in a group, fill in questionnaires in English, give informed consent or return for a FU visit
Interventions	Individual or group fitting with follow-up in an individual or group format
Outcomes	Medium-term: 6 months - inner EAR (HEARING FUNCTION), USE hours/day, costs of planned and unplanned visits over the 6 months FU period (ECONOMIC), HHIE (HANDICAP), CPHI, SADL (SATISFACTION), IOI-HA (USE and HANDICAP), SF-12 (QoL)
Notes	NB data for group and individual arms added together. Patients randomised twice, first prior to fit and then again prior to FU

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomized" Comment: no details of randomisation protocol but probably done

Collins 2013 (Continued)

Allocation concealment (selection bias)	Low risk	Quote: “sealed opaque envelopes”
Blinding of participants and personnel (performance bias) All outcomes	High risk	Acknowledged limitations - shame as could have been single-blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding of outcome assessment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Some dropouts but overall quite low in this large cohort so unlikely to affect results
Selective reporting (reporting bias)	Low risk	Protocol published in 2009 so able to compare aims with outcomes
Other bias	Unclear risk	Participants paid for their participation

Cunningham 2001

Methods	Randomised Funding: participants were provided with ITEs free of charge, Mary and Mason Rudd Surgical Research Fund, Siemens provided the HAs
Participants	N = 18 Age: mean intervention 65.22, control 68.78 Inclusion criteria: 50 to 75 years, moderate symmetrical SNHL, no Hx of otologic/neurologic disease, good general health Exclusion criteria: other aural or vestibular signs or Sx, previous HA use
Interventions	Control 'usual care' versus as many post-fitting adjustments as patients requested
Outcomes	Medium-term: APHAB (BENEFIT), SIN test (SPEECH PERCEPTION), hours per day (USE), satisfaction scale
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: “randomly assigned” Comment: no detail given and indeed no detail given of number in each group
Allocation concealment (selection bias)	Unclear risk	No detail on allocation procedure

Cunningham 2001 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	High risk	No blinding
Blinding of outcome assessment (detection bias) All outcomes	High risk	No blinding
Incomplete outcome data (attrition bias) All outcomes	Low risk	No apparent missing data post-randomisation
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	Patients given access to previous test scores for APHAB administration No power calculation to determine if sufficient numbers to demonstrate an effect

Eriksson-Mangold 1990

Methods	Randomised Conducted in 1985
Participants	N = 56 “picked out from the waiting list of new hearing aid candidates” probably randomly 28 in AF group 28 in control group Age: range 50 to 74 years Inclusion criteria: hearing loss at least 35 dB across 3 frequencies, speech discrimination 50% or more
Interventions	HA plus standard FU appointments versus ‘active fitting’ programme (including task-orientated diary to complete at home)
Outcomes	Medium-term: 10 months post-fit structured telephone interview including a 5-point scale of daily use
Notes	-

Risk of bias

Bias	Authors’ judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: “randomised into 2 groups” Comment: procedure for randomisation not given
Allocation concealment (selection bias)	Unclear risk	Comment: no information given on allocation

Eriksson-Mangold 1990 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Comment: psychologists carrying out the FU interview were blind to group allocation
Incomplete outcome data (attrition bias) All outcomes	Low risk	Low numbers of dropouts and reasons given
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Low risk	Study appears to be free of other sources of bias

Ferguson 2016

Methods	Randomised
Participants	N = 203; 103 intervention, 100 control Age range: 42 to 94 years Gender: 41% female Inclusion criteria: aged < 18, first time hearing aid user, English as a first language or good understanding of English Exclusion criteria: unable to access PC, DVD or internet, unable to complete questionnaires due to age-related problems
Interventions	Educational material delivered via DVD, PC or internet (patient preference) post fitting, 7 modules covering acclimatisation, getting to know the hearing aid, insertion of hearing aid, troubleshooting, expectations, phones and assistive listening devices, communication
Outcomes	Short term: GHABP, PHAST, SADL, IOI-HA, HHIE, HACK, HADS, PAM, EQ-5D, IT literacy and data logged HA use
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "allocation was based on a computer-generated pseudo-random code" Comment: sequence generation was adequate
Allocation concealment (selection bias)	Low risk	Quote: "allocations were revealed to the research team on completion of the study" Comment: adequate allocation concealment

Ferguson 2016 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	High risk	The participants could not be blinded due to the nature of the intervention and control
Blinding of outcome assessment (detection bias) All outcomes	High risk	Attempts were made to blind researchers to group allocation at the assessment stage but this was not successful
Incomplete outcome data (attrition bias) All outcomes	Low risk	Incomplete outcome data were accounted for
Selective reporting (reporting bias)	Unclear risk	No protocol available All expected outcomes appear to have been reported
Other bias	High risk	Some post hoc secondary subgroup analyses

Fitzpatrick 2008

Methods	Randomised but control participants crossed over to intervention after the control sessions	
Participants	N = 24 (14 intervention and 10 control) Age: range intervention 45 to 86, mean 69.5; control 61 to 88, mean 70.1 intervention 9 female, 5 male; control 6 females, 4 male Inclusion criteria: 18 years plus, have high school diploma, native English speakers, SNHL, used binaural HAs for at least 6 weeks Exclusion criteria: SF-12 score < 50%, word recognition score < 60%, no known neurological or psychiatric problems	
Interventions	Auditory training versus lectures on HL and HAs and discussion of communication tactics Auditory training consisted for 16 sessions - 13 training and 3 test sessions of 1 hour each	
Outcomes	Medium-term: NU-6, CST, CCT (SPEECH PERCEPTION), HA use and satisfaction questionnaire (USE)	
Notes	-	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Comment: random numbers table used - even number experimental, odd numbers control

Fitzpatrick 2008 (Continued)

Allocation concealment (selection bias)	Unclear risk	Comment: although random number tables used it is unclear who undertook the allocation and whether this was concealed from the researchers
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blind due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Apparently no missing data - must have had very highly motivated patients
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	Intervention group had training with one of the tests used in the evaluation sessions Also there was a baseline difference between the groups with the control group having higher scores on 2 of the speech perception tests

Gil 2010

Methods	Randomised	
Participants	N = 14 (7 control, 7 intervention) Age: details of actual age range not given but all must have been under 60 Inclusion criteria: 16 to 60 years old, mild to moderate bilateral sloping SNHL, word recognition 72% or more, 3 months + HA use Exclusion criteria: other neurological, psychological, cognitive disorders or mental disturbances	
Interventions	Auditory training - 8 x 1-hour sessions held twice a week for 4 weeks	
Outcomes	Short-term: electrophysiological (long-latency auditory evoked potentials), behavioural auditory processing (sound localisation, memory for verbal sounds in sequence, memory for non-verbal sounds in sequence, word recognition score, speech in noise test, synthetic sentence identification, dichotic digits), APHAB (BENEFIT)	
Notes	-	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Random sequence generation (selection bias)	Unclear risk	Quote: “randomly divided” Comment: no details given on procedure
Allocation concealment (selection bias)	Unclear risk	No details of allocation procedure
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Evaluation after intervention was carried out by a researcher who was blind to the participant’s group and was blind to participant’s baseline results
Incomplete outcome data (attrition bias) All outcomes	Low risk	No apparent missing data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	Only change scores presented and there was a reported difference between groups at baseline, which may have affected the outcome and was not fully addressed

Kemker 2004

Methods	Randomised but with balanced group allocation
Participants	N = 45 (1 participant excluded and his data not included so 44 - he was in the post-fit group) Age: range 60 to 80 Inclusion criteria: new HA users, US veterans, 23 or higher on mini mental state exam Exclusion criteria: patients being followed by VA visual impairment team
Interventions	HA alone versus pre-fitting hearing aid orientation + HA versus HA + pre and post-fit hearing aid orientation Orientation was 2 x 1-hour sessions 1 week apart
Outcomes	Short-term: GHABP (which includes measure of USE)
Notes	-

Risk of bias

Bias	Authors’ judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: “systematic random sampling scheme” Comment: to give 15 in each group - process not described beyond that except that

Kemker 2004 (Continued)

		word recognition scores were monitored to ensure balancing. Insufficient detail
Allocation concealment (selection bias)	High risk	Clearly not as the groups were balanced on the basis of word recognition scores
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: "double-blind" Comment: participant would know which group they were in
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not stated whether researchers administering the questionnaires/analysing results knew which group patients were in
Incomplete outcome data (attrition bias) All outcomes	Low risk	1 patient dropped out - reasons given and not study related; their data were excluded
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	No power calculation to determine if sufficient numbers to demonstrate an effect

Kramer 2005

Methods	Randomised	
Participants	N = 24 intervention and 24 control (plus their significant others) completed all and data analysed but 58 were initially recruited and randomised. 2 dropped out of training group (ill health and problems operating the video) and 8 further HI participants failed to return questionnaires (not clear which group they were in) Inclusion criteria: mix of new and existing HA users	
Interventions	HA alone versus HA plus home education programme for patients and significant others Home education programme - 5 video tapes and an instruction booklet Tapes sent out one at a time - only sent next one when returned previous	
Outcomes	Medium-term: 6 months post-intervention IOI-HA/IOI-AI	
Notes	Cannot include these outcomes in a meta-analysis as the 2 versions of the questionnaire are measuring different things - not a valid comparison The other outcomes were amended from other questionnaires for this study and so could not be used	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Kramer 2005 (Continued)

Random sequence generation (selection bias)	Unclear risk	Quote: “randomly allocated” Comment: no details given on procedure
Allocation concealment (selection bias)	Unclear risk	No details given on randomisation procedure
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants could not be blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding in collection of outcomes
Incomplete outcome data (attrition bias) All outcomes	High risk	Details are not given about numbers in each randomised group - only N post dropout/non-returned questionnaires. No details about which group the non-returned were in. 1 patient in intervention group dropped out due to problems with using the video - their results were not included
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	The intervention group and control group were evaluated using different versions of the same questionnaire. Subsequent research suggests this is not valid Also no power calculation

Kricos 1992

Methods	Randomised	
Participants	N = 26 (control 13, intervention 13) Age: range 61 to 83 years, mean 70.8 Inclusion criteria: HA users, no previous AR, bilateral SNHL, corrected vision of 20/30	
Interventions	4-week communication training programme - individual, twice a week 1-hour sessions - 8 hours in total	
Outcomes	Short-term: HHIE (HANDICAP), speech recognition test - audiovisual and audio only Central Institute for the Deaf Everyday Sentence Test % syllables correctly identified	
Notes	-	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No details given of randomisation procedure

Kricos 1992 (Continued)

Allocation concealment (selection bias)	Unclear risk	No details given
Blinding of participants and personnel (performance bias) All outcomes	High risk	The participants were not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No apparent missing data but not explicitly stated
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	No power calculation

Kricos 1996

Methods	Assigned on a rotating basis to 1 of 3 groups
Participants	N = 78 Age: range 52 to 85 Inclusion criteria: significant handicap score on HHIE, native English speakers with adult onset HL, existing HA users, 20/40 corrected vision
Interventions	Analytic auditory training (N = 26) same/different judgements between syllable pairs Active listening (N = 26) communication training Control (N = 26)
Outcomes	Short-term: CST (SPEECH PERCEPTION), HHIE (HANDICAP), CPHI (PSYCHOSOCIAL FUNCTION, COMMUNICATION AND HANDICAP)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Assigned on a rotating basis to 1 of 3 groups - quasi-randomised
Allocation concealment (selection bias)	High risk	No allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention

Kricos 1996 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	No apparent missing data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	It is unclear whether there are further sources of bias

Lavie 2014

Methods	Randomised No details of data collection period
Participants	N = 36, 12 simultaneous binaural fitting, 24 sequentially (12 left ear first, 12 right ear first) Age range: 64 to 88 years Gender: 20 men, 16 women Inclusion criteria: mild to moderate hearing loss, symmetric speech discrimination scores for each ear, first time hearing aid users, willing to try 2 aids Exclusion criteria: mini-mental state exam ≤ 24
Interventions	Group 1 - fitted binaurally Group 2 - fitted with aid for right ear and then left ear a month later Group 3 - fitted with aid for left ear and then right ear a month later Otherwise all groups received same rehabilitation/counselling/instruction
Outcomes	Short-term: 'compliance' assessed as high, fair or poor at 1 month and 2 months, average hours per day (data-logged)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Participants were randomly divided in to three equal-size groups" Comment: no further details of how this was done
Allocation concealment (selection bias)	High risk	Comment: unable to conceal allocation from participants or investigators during the first month, unclear whether allocation concealment from investigators was

Lavie 2014 (Continued)

		achieved in the second month
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	Investigators not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Some data excluded but reasons given
Selective reporting (reporting bias)	Unclear risk	No protocol available but published data include all expected outcomes
Other bias	Low risk	The study appears to be free of other sources of bias

Lundberg 2011

Methods	Randomised Data collection 2007 to 2008
Participants	N = 69 final numbers but there is some discrepancy in the way this was reached depending on where you read in the text Inclusion criteria: 60 to 75 years of age, mild-moderate HL, to have had HA fitted at least 1 year prior to study, to have HHIE score of > 20 (indicative of residual handicap)
Interventions	Intervention group received a booklet with weekly topic-based reading instructions based on chapters of the book plus 5 x 10- to 15-minute telephone calls with an audiologist to discuss the content of the book Control group received the booklet but no instructions or telephone follow-up
Outcomes	Short-term: HHIE (HANDICAP), HADS (PSYCHOLOGICAL IMPACT), IOI-HA (inc USE)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomised" Comment: probably done but no details of exact randomisation procedure given

Lundberg 2011 (Continued)

Allocation concealment (selection bias)	Unclear risk	Randomisation was carried out by someone independent of the study data collection but not clear whether concealed
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	Comment: authors do comment that blinding the questionnaire administrators may have improved quality of the study. They recognised the potential bias
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	They did explain how many dropped out and gave reasons and those included under ITT where included on a LOCF basis 1 participant in the control group was deemed an outlier and was thereby excluded from analyses because the participant's data differed by more than 2 SD from the control group mean measured before and after the intervention. Unclear whether this is appropriate
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Low risk	Authors give a good discussion of other potential sources of bias

Miranda 2008

Methods	Randomised Data collection 2005 and first quarter 2006
Participants	N = 13 (control 7, intervention = 6) Age: range 60 to 74 years, mean 65.3 Inclusion criteria: mild to moderately severe bilateral sensorineural hearing, symmetrical hearing loss of flat or slightly descending curve shape in the high frequencies; speech recognition index equal to or above 72% bilaterally; received a HA donation in the last 3 months; use or have the indication to use intracanal HAs in binaural fitting; not having perceivable cognitive alteration or speech alteration; more than 60 years of age
Interventions	Auditory training: 7 sessions, held once a week, with duration of 50 minutes each
Outcomes	Short-term: HHIE (HANDICAP), speech perception
Notes	-

<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "the individuals themselves pick a number to be randomized to which group they would be sent to" Comment: not 100% clear how this worked but almost certainly randomised
Allocation concealment (selection bias)	Unclear risk	Exact randomisation procedure unclear but may have been concealed if out of a hat
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention Those in control group who were interested in training were offered the chance - not clear whether this offer was made before or after the study
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Single-blinded - evaluations carried out but researcher blind to treatment group
Incomplete outcome data (attrition bias) All outcomes	Low risk	No apparent missing data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	Significant difference between the groups at baseline For the treatment group they reported the results for the 2 ears separately to double the sample size - incorrect assumption

Oberg 2008

Methods	Randomised Data collection Autumn 2005
Participants	Diagnosis: symmetrical mild to moderate SNHL N = 38 (19 intervention, 19 control) Age: range not given but mean intervention 67.1 and control 65.5 Inclusion criteria: first-time users aged 20 to 80, good general health, fluent in Swedish Exclusion criteria: evidence of cognitive deficits during the interview or on a test of verbal fluency

Oberg 2008 (Continued)

Interventions	Individual pre-fitting sound awareness training 3 visits each with different listening exercises and also use of the experimental adjustable aid	
Outcomes	Short-term: post-fitting - HHIE, SADL, CSS, HADS, IOI-HA Long-term: 1 year - HHIE, SADL, CSS, HADS, IOI-HA, COSI, speech recognition	
Notes	-	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: “the randomisation procedure was performed by an independent researcher. The researcher allocated the participants according to a computer-generated randomisation list”
Allocation concealment (selection bias)	Low risk	The audiologists who saw the participants in the clinic were blind to this list
Blinding of participants and personnel (performance bias) All outcomes	High risk	The participants were not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	All telephone consultations were conducted by an “independent audiologist”
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: missing outcome data balanced across groups with similar reasons across groups but sometimes reasons for dropouts not clear
Selective reporting (reporting bias)	Low risk	No protocol published but was described in thesis (which we have)
Other bias	Unclear risk	Study was under-powered Non-responders who declined the telephone interview but completed the IOI-HA reported significantly less use of aids than responders - not clear which groups the non-responders came from

Oberg 2009

Methods	Randomised Data collection autumn 2004
Participants	Diagnosis: symmetrical mild to moderate SNHL N = 39 (19 intervention, 20 control) Age: range not given, mean 68.6 intervention and 69.8 control Inclusion criteria: first-time users aged 20 to 80, good general health, fluent in Swedish Exclusion criteria: evidence of cognitive deficits during the interview or on a test of verbal fluency
Interventions	3 visits to clinic - 1 per week First week fitted with a user-controlled adjustment experimental aid Subsequent visits they adjusted the aid to preferred settings Wore aids at home in between
Outcomes	Short-term: week 6 post-intervention (pre HA) - HHIE, ECHO, CSS, HADS, COSI Medium-term: week 18 post-fitting - HHIE, SADL, CSS, HADS, IOI-HA Long-term: 1 year FU - HHIE, SADL, CSS, HADS, IOI-HA, COSI
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "the randomisation procedure was performed by an independent researcher. The researcher allocated the participants according to a computer-generated randomisation list."
Allocation concealment (selection bias)	Low risk	The audiologists who saw the participants in the clinic were blind to this list
Blinding of participants and personnel (performance bias) All outcomes	High risk	The participants were not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	All telephone consultations were conducted by an "independent audiologist"
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: missing outcome data balanced across groups with similar reasons across groups
Selective reporting (reporting bias)	Low risk	No protocol published but was described in thesis (which we have)

Other bias	Low risk	The study appears to be free of other sources of bias
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Olson 2013

Methods	Randomised trial (also had a group of experienced HA users who received training but allocation to this group was not randomised)
Participants	N = 29 in total but we cannot include data for experienced users. N = 8 new users plus training, N = 7 new users control Age range: mean 66 years in both groups Inclusion criteria: 'new' HA users (4 week to 6 months experience), 50 to 81 years old, mild-moderate bilateral SNHL and bilateral HAs, native speakers of American English, adequate vision, daily access to TV and DVD player with remote control Exclusion criteria: neurological, psychiatric disorder, conductive or asymmetric hearing loss
Interventions	LACE DVD - 20 x 30-minute sessions at home over a 4-week period
Outcomes	Short-term: at end of 4-week home training period - QuickSIN, Compressed Speech Test (word recognition), Synthetic Sentence Identification (competing speaker task) ALL SPEECH PERCEPTION, IOI-HA/AI, Speech, Spatial and Qualities of Hearing Scale - only 2 subscales as spatial considered not relevant
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "New HA users were randomly assigned to the training or control (nontraining) group as determined by random tables"
Allocation concealment (selection bias)	Unclear risk	No details of allocation concealment given
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	There were 4 dropouts. The reasons were given and were unrelated to the study in 2 cases but it is not clear which groups they

Olson 2013 (Continued)

		came from. N in each group prior to the dropouts was not quoted. The data from the 4 participants that did not complete the study were excluded from the analysis
Selective reporting (reporting bias)	High risk	No protocol available but data for SSQ not reported although it was listed as an outcome measure
Other bias	High risk	The Tx group had an additional test session at 2 weeks, which the control group did not have, and so they had extra experience with the test situation and material The study was also under-powered after dropouts by their own calculation Subjects were paid for their participation

Preminger 2008

Methods	Randomised
Participants	N = 53 (3 dropped out during study and were not included in analysis) Age: control mean 66, training + psychosocial (T + P) 65.3, training only (TO) 64.9 Gender: control 75% male, T + P 37.5%, TO 66.7% apparently not statistically significant on Chi ² test BUT is a big difference Inclusion criteria: aged 55 to 75, at least 3 months HA experience and > 20 score on HHIE or HHIA, corrected binocular visual acuity 20/40, passed MMSE, passed a screen for APD
Interventions	Training group: hour-long classes of speech training once a week for 6 weeks Training plus psychosocial exercises: as above plus an extra 30 minutes psychosocial exercises At least 2 instructors per class
Outcomes	Short-term: 6 weeks - CUNY AB wordlists auditory and audio-visual (SPEECH PERCEPTION), CUNY topic-related sentences auditory and audio-visual (SPEECH PERCEPTION), HHIE (HANDICAP, HEARING RELATED QoL), WHO Disability Assessment Schedule II (GENERIC QoL) Medium-term: 6 months - AS ABOVE
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly allocated" Comment: no details of procedure given

Preminger 2008 (Continued)

Allocation concealment (selection bias)	Unclear risk	Comment: no details given
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	3 dropouts which were excluded from the study - only evident from reading carefully. All dropouts from treatment groups. Reasons given but only partially clear; sensible management of dropouts in analysis
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	Quality of life measures completed with researcher present The gender and hearing handicap differences present at baseline, while not statistically significant between groups, apparently may have had an effect

Preminger 2010

Methods	Randomised BUT was made on basis of preference regarding class time so 'quasi-randomised'	
Participants	N = 36 Age range: no range given but average Cx 72.2, Tx 63.5 - significant difference Inclusion criteria: all PHL had to score over 20 on HHIE, scores below 25 on QuickSIN so they would have no problems communicating in group class, SO had to have PTA over 30 dB HL (near normal hearing at least)	
Interventions	AR group programme just for people with hearing loss (spouses no treatment) versus AR group programme plus separate group programme for spouses 90-minute sessions 1 x weekly for 4 weeks (no specific homework)	
Outcomes	Short-term: 4 weeks - HHIE (HEARING HANDICAP), perceived stress scale and affect rating scale (PSYCHOLOGICAL), primary communication inventory (COMMUNICATION) Medium-term: 6 months - as above	
Notes	-	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Preminger 2010 (Continued)

Random sequence generation (selection bias)	High risk	Quote: "Couples were assigned to either the control or the experimental AR sessions based on the couples' preferred class meeting times. Participants were only given class meeting times and no information about the class content (control versus experimental); in this way they could not choose to be in either the control group or the experimental group." Comment: quasi-randomised
Allocation concealment (selection bias)	High risk	No information re sequence generation but researchers presumably knew which class was which and therefore which participants were choosing
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants were not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding of outcome assessment
Incomplete outcome data (attrition bias) All outcomes	Low risk	No apparent dropouts or missing data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	Participants were a mix of CI and HA patients Also age difference in groups and in mood scores pre-intervention Scales all completed in presence of a researcher "to answer questions and make sure they were filled out correctly"

Preminger 2010a

Methods	Randomised on the basis of their choice of class time
Participants	N = 52 (18 group 1, 17 group 2, 17 Group 3) but there were 4 on top of this who dropped out but were not included Age: no range given - no significant differences in means between groups. Overwhelmingly male, VA population Inclusion criteria: 55 to 75, experience HA users (3 months plus), score at least 20 on HHIE, corrected binocular vision 20/40, passed MMSE, passed screen for APD described in 2008 study

	Exclusion criteria: fluctuating hearing loss during study	
Interventions	Group 1: communication strategies group Group 2: communication plus psychosocial group Group 3: informational lecture plus psychosocial group 1-hour lecture per week for 6 weeks - all participants completed 5 of the 6 classes	
Outcomes	Short-term: post-intervention - HHIE/A (HEARING RELATED QoL, HANDICAP) , WHODAS 2.0 (GENERIC QoL) Medium-term: 6 months post-class - as above	
Notes	-	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Quote: "Participants were randomly assigned to each treatment group based on their preferred class meeting times. Participants were given class meeting times and no information about the class content" Comment: quasi-randomised
Allocation concealment (selection bias)	High risk	Patients given the choice based on the above - researchers knew which group was at which time
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants were not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No blinding of outcome measurement
Incomplete outcome data (attrition bias) All outcomes	Low risk	There were dropouts but reasons were given
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	Questionnaires completed in the presence of a researcher (who was not blind to the group allocation)

Saunders 2009

Methods	Randomised
Participants	Diagnosis: symmetrical SNHL (< 15 dB HL difference between ears on 4 frequencies average) N = 60 (18 female, 42 male), 2 dropouts (1 from group 1 and 1 dropout from group 2, reasons given); analysed data from 58 people Age: range 55 to 81 years Inclusion criteria: first-time users
Interventions	Group 1: pre-fitting counselling including demonstration of listening situations, post-fit fine tune if wanted Group 2: pre-fitting counselling including demonstration of listening situations but no fine tuning Group 3: pre-fitting counselling but no demonstration and no fine tune post-fit Pre-fitting counselling based on COSI
Outcomes	Short-term: 8 to 10 weeks - HHIE/A aided, APHAB-A, PIADS-A, SADL, categorical assessment of USE PIADS = Psychosocial Impact of Assistive Devices Scale
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly assigned" Comment: no details given
Allocation concealment (selection bias)	Unclear risk	No details given
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	2 missing data - reasons given and not both from same group so unlikely to affect analysis
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	Questionnaires completed in clinic - not clear whether researcher present No power calculation No control group who were only aided without the pre-fitting counselling

Methods	Randomised
Participants	<p>N = 279: 136 were new HA users and 143 experienced users</p> <p>Age range: age range not specified but average age was approx 68 across the 4 groups of new users</p> <p>Gender: not specified but given that these were veterans it is very likely that they were nearly all or all male</p> <p>Inclusion criteria: English as first language, MMSE scores age appropriate, ability to read at 5th grade level or higher, good corrected vision, symmetrical hearing loss, 3 frequency average hearing loss < 50 dB HL, unaided speech recognition scores of > 40%, had to be willing to have HAs set to NAL prescription</p> <p>Exclusion criteria: no external ear disease, conductive or retrocochlear pathology, comorbidities that would interfere with participation</p>
Interventions	<p>4 groups:</p> <p>LACE DVD: auditory training 10 x 30-minute sessions over a 2-week period at home on DVD</p> <p>LACE C: auditory training 20 x 30-minute sessions over a 4-week period at home on computer</p> <p>Placebo: listening to an audio book 20 x 30-minute sessions over a 4-week period</p> <p>Control: standard HA fitting</p>
Outcomes	<p>Short-term: 2 to 6 weeks - HHIE/A, 5 behavioural measures of speech reception</p> <p>Medium-term: 6 months - as above</p>
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Each site was provided with sequentially numbered randomization envelopes that specified the intervention group to which the participant was assigned. A block randomization scheme with a block size of eight was utilized with stratification of hearing aid experience (new or experienced listener) across the four intervention groups."
Allocation concealment (selection bias)	Unclear risk	No information about allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Quote: "...participants in the LACEDVD, LACE-C, and placebo groups were masked regarding whether they were receiving experimental or placebo training"
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "The research audiologists were not blinded to the participant's intervention-arm assignment"
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition was clearly explained and reasons for dropouts given

Selective reporting (reporting bias)	Unclear risk	No protocol available but paper includes all expected outcomes
Other bias	Low risk	The study appears to be free of other sources of bias

Smaldino 1988

Methods	Randomised
Participants	N = 40 (19 females and 21 males) 10 in each group Age: range 30 to 90 years, mean 69 Inclusion criteria: new HA users
Interventions	Control group: HA with simple orientation Cognitive style: as control but also given information about their learning style Cognitive + AR - 4-week AR programme plus information on cognitive style AR was individual computer-based
Outcomes	Short-term: HPI (HANDICAP)
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Protocol for randomisation not given
Allocation concealment (selection bias)	Unclear risk	No details provided
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing data
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	Only change scores available - not post-score Not clear whether there was a difference at baseline in the outcome measure as handicap was assessed pre-intervention with a different measure

Sweetow 2006

Methods	Randomised cross-over trial	
Participants	N = 65 across 5 sites Age: range - trained 28 to 85 (average 63.15); control 32 to 82 (average 64.2 years)	
Interventions	Home-based interactive PC-based program (Listening and Communication Enhancement LACE) 30 minutes 5 times a week for 4 weeks	
Outcomes	Short-term: 4 weeks - QuickSIN, Hearing in Noise Test (HINT) (SPEECH PERCEPTION), HHIE/A (HANDICAP), Communication Scale for Older Adults (COMMUNICATION)	
Notes	Group 2 cross-over arm - as there were no significant differences between group 1 and cross-over arm of group 2 they pooled the data for these groups meaning we cannot use the data. Also data from HA users mixed with non-users. This means that the data from this study have not been included in any meta-analysis Outcomes for a 'subset' of participants - not clear how this set was decided on	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No details of sequence generation
Allocation concealment (selection bias)	Unclear risk	No details of randomisation protocol
Blinding of participants and personnel (performance bias) All outcomes	High risk	This study was not blinded due to the nature of the intervention
Blinding of outcome assessment (detection bias) All outcomes	High risk	No apparent blinding of outcome assessment
Incomplete outcome data (attrition bias) All outcomes	High risk	Some confusion over numbers for how many started and completed in each group. N is stated as 65 but this is not the number randomised - it is the number who completed. Number randomised is unclear
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	Participants at one of the sites were paid Data not available separately for users versus non-users, for different sites or for the 2 arms of the trial The authors have a financial interest in the company that produces the intervention LACE

Thoren 2011

Methods	Randomised
Participants	N = 59 (intervention group 29, control 30) recruited through adverts in newspapers and referred to a website Age: range 24 to 84, mean 63.5 29 women and 30 men; majority (67%) had education equivalent to university level Inclusion criteria: experienced HA users, hearing impairment with subjective significant communication difficulties (defined as over 20 on HHIE), using HA for at least 1 year, 18 years plus, Swedish as first language, access to PC and internet Exclusion criteria: not able to have a conversation by telephone, severe tinnitus, Ménière's
Interventions	Intervention: online education programme including professional guidance (5-week programme including information, tasks, assignments and professional contact) Control: online discussion forum with weekly topics but no professional guidance
Outcomes	Short-term: immediately post-intervention - questionnaires administered online HHIE, IOI-HA, SADL, HADS Medium-term: 6 months - as above
Notes	This was a well-designed study with a good attempt made to include a placebo intervention References CONSORT guidelines

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "...randomly assigned by an independent researcher" Comment: almost certainly done
Allocation concealment (selection bias)	Low risk	Allocation undertaken by researcher independent of the study
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Comment: probably was blinded to the participants - they were all given the same instructions pre-study and both intervention and control group involved the internet However, blinding not explicitly stated but implied
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Outcome assessment was online
Incomplete outcome data (attrition bias) All outcomes	Low risk	Results analysed on ITT LOCF - very clearly explained. Reasons for dropouts given
Selective reporting (reporting bias)	Unclear risk	As far as one can tell but no protocol available
Other bias	Low risk	This study appears to be free of other sources of bias. Limitations discussed No 'no treatment' control but the placebo control group was

Thoren 2011 (Continued)

	well thought out
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Thoren 2014

Methods	Randomised
Participants	N = 76 (38 in each group) Age range: 26 to 81 years Gender: 32 women, 44 men Inclusion criteria: 1 year + HA use, over 18 years of age, significant hearing impairment and significant communication difficulties
Interventions	Online rehabilitation for hearing aid users including self-study, training and professional coaching in hearing physiology, hearing aids and communication strategies, as well as online contact with peers across 5 weekly modules. The intervention group was compared to a waiting list control group
Outcomes	Short-term: immediately following intervention, IOI-HA, HHEI and HADS Medium-term: at 3 months measures repeated
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "The participants were randomized by an independent person (not involved in the study or recruitment) to either participate in the intervention group or in the control group." Comment: insufficient information about the sequence generation process in study
Allocation concealment (selection bias)	Low risk	Comment: the use of an independent person performing the randomisation is suggestive of allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Missing outcome data balanced in numbers across intervention groups; it was explained and due to attrition. Missing data were imputed using appropriate methods
Selective reporting (reporting bias)	Unclear risk	No protocol available

Other bias	Unclear risk	Quote: "Of the participants, 75% had completed education at university level." Comment: study appears to have a risk of recruitment bias
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Turbin 2006

Methods	Randomised
Participants	N = 135 (only 1 female), 90% non-Hispanic whites Age: range 46 to 85 Inclusion criteria: new and experienced HA users Exclusion criteria: participated in AR in last 5 years, no neurological, neuromuscular, psychiatric diagnosis to interfere with use of HA or participation in age-normal social activities
Interventions	HA alone (N = 66) versus HA plus single AR workshop (N = 69)
Outcomes	Short-term: 8 weeks post-fit - CPHI (communication strategies subscale, personal adjustment subscale, other scales), COSI, NEO-FFI - 5-factor personality inventory, WOCQ - ways of coping questionnaire Medium-term: 6 months - as above
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Author reports group allocation was randomised but process not clear
Allocation concealment (selection bias)	Unclear risk	No details available
Blinding of participants and personnel (performance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Not clear whether results analysed only for those who remained or on an ITT basis Dropout rate was higher from the intervention group at 8 weeks and at 6 months, which could have affected the results Reasons for dropout not given High dropout rate overall

Turbin 2006 (Continued)

Selective reporting (reporting bias)	Unclear risk	Results not published - data taken from presentation obtained from the author
Other bias	Unclear risk	Not enough information to make a judgement on other sources of bias

Vreken 2015

Methods	Randomised
Participants	N = 131: 64 intervention group, 67 in control group Age range: intervention group - mean age in years 81.2 (SD 10.0), control group - mean age in years 81.8 (SD 10.1) Gender: intervention group - 41.3% female, control group - 60% female Inclusion criteria: reported hearing disability, possessed a HA and had the cognitive ability (reported by a spouse, other relative or care personnel) and sufficient knowledge of the Dutch language to comprehend or respond to questions. Attendance at a low vision clinic, reporting hearing disability and owned a hearing aid
Interventions	Dual sensory loss protocol consisting of a handbook with background information and a checklist accompanied with exercises. The intervention was delivered in 3 to 5 weekly home visits. The protocol covered: hearing aid use, maintenance and handling; living environment; and hearing assistive devices and communication strategies and coping with DSL. The intervention group was compared to a waiting list control group
Outcomes	Medium-term: at 3 months IOI-HA
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "randomization was stratified per OTs' area of practice (eight strata). After completion of baseline measurements, an independent researcher not involved in the trial used randomization software to assign participants in each stratum. Participants were randomly allocated to either the IG or CG in blocks of two." Comment: randomisation was appropriate
Allocation concealment (selection bias)	Low risk	The use of an independent person performing the randomisation is suggestive of appropriate allocation concealment
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants were not blinded

Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The investigators and research assistants performing the measurement were not aware of the treatment allocation."
Incomplete outcome data (attrition bias) All outcomes	Low risk	Detailed information was provided on the numbers of participants not receiving the intervention and those lost to follow-up and how the data were included/excluded from the analysis
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	The study may have been under-powered

Ward 1978

Methods	6 consecutive patients in each group
Participants	Diagnosis: better ear 35 to 62 dB across 3 frequencies N = 36 (1 dropout who was discovered to have already had a HA), 15% of those fitted over that period so admit group was more selective than they intended Age: range 60 to 80 Inclusion criteria: new HA users, over retirement age Exclusion criteria: over 80 years old, predominantly conductive losses
Interventions	Group 1: fitting plus 2 group sessions of 2 hours each at 2 and 4 weeks post-fit Group 2: fitting plus 4 group sessions of 2 hours each at 2-week intervals Group 3: fitting only Up to 6 patients in each group
Outcomes	Medium-term: 6 months - hours of USE (patient report and battery use), change in hearing handicap (Hearing Measurement Scale), AB word lists score
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Comment: in fact a cluster quasi-randomised trial
Allocation concealment (selection bias)	Unclear risk	No details given, but as it was possibly allocated on a rotating basis every 6 patients allocation concealment is unlikely
Blinding of participants and personnel (performance bias) All outcomes	High risk	Participants not blinded due to the nature of the intervention

Ward 1978 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Comment: data for group 2 not analysed at all due to high dropout rate. Reasons for dropout given
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	Unclear risk	Not enough information to make a judgement about other sources of bias

Ward 1981

Methods	Randomised
Participants	N = 31 Age: range not given Inclusion criteria: sequential patients seen 3 months post-fitting, over 65, scored 2 or more on a questionnaire on hearing tactics (poor performance), only measured for those who had a HMS score of 15 or more and who wore aids for less than 8 hours per day (so were capable of improvement) at 3 months Exclusion criteria: frail, poor sight
Interventions	Control (N = 13) versus self-instruction package on hearing tactics (N = 9) versus hearing tactics instruction (individual) (N = 9)
Outcomes	Medium-term: 6 months after fitting - change in HMS score 3 to 6 months
Notes	-

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "each person was randomly allocated" Comment: no details of sequence generation
Allocation concealment (selection bias)	Unclear risk	No details given
Blinding of participants and personnel (performance bias) All outcomes	High risk	No blinding
Blinding of outcome assessment (detection bias) All outcomes	High risk	No blinding

Ward 1981 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Reasons for dropouts given and numbers balanced across groups
Selective reporting (reporting bias)	Unclear risk	No protocol available
Other bias	High risk	The study was small and the high number of exclusions prior to randomisation meant that groups were not balanced for age or gender

Abbreviations used:

AF: active fitting
 APD: auditory processing disorder
 APHAB: Abbreviated Profile of Hearing Aid Benefit (Cox 1995)
 AR: auditory rehabilitation
 ARS: Affect Rating Scale (Lawton 1992)
 CCT: California Consonant Test (Owens 1977)
 CI: cochlear implant
 COSI: Client Orientated Scale of Improvement (Dillon 1997)
 CPHI: Communication Profile for the Hearing Impaired (Demorest 1987)
 CSOA: Communication Scale for Older Adults (Kaplan 1997)
 CSS: Communication Strategies Scale (Demorest 1987)
 CST: Connected Speech Test (Cox 1998)
 CUNY: City University of New York
 Cx: control group
 ECHO: Expected Consequences of Hearing Aid Ownership (Cox 2000)
 EQ-5D: EuroQoL 5D (EuroQol 1990)
 FU: follow-up
 GHABP: Glasgow Hearing Aid Benefit Profile (Gatehouse 1999)
 HA: hearing aid
 HACK: Hearing Aid Communication Knowledge (Ferguson 2015)
 HADS: Hospital Anxiety and Depression Scale (Zigmond 1983)
 HCA: Hearing Coping Assessment (Andersson 1995a)
 HF: high frequency
 HI: hearing impaired
 HHIE: Hearing Handicap Inventory for the Elderly (Ventry 1982)
 HL: hearing loss
 HMS: Hearing Measurement Scale (Noble 1970)
 HPI: Hearing Performance Inventory (Giolas 1979)
 HR QoL: Hearing-Related Quality of Life
 Hx: history
 inner EAR and outer EAR: Effectiveness of Auditory Rehabilitation scales (Yueh 2005)
 IOI-AI: International Outcome Inventory - Alternative Interventions
 IOI-HA: International Outcome Inventory for Hearing Aids (Cox 2002)
 IT: information technology
 ITE: in-the-ear
 ITT: intention-to-treat
 LACE: Listening And Communication Enhancement (Sweetow 2006)
 LOCF: last observation carried forward

MMSE: mini mental state exam
 NEO-FFI: Neuroticism Extroversion Openness Five Factor Inventory ([Costa 1992](#))
 NST: Nonsense Syllable Test ([Dubno 1981](#))
 NU-6: Northwestern University auditory test no. 6 ([Tillman 1966](#))
 PAM: Patient Activation Measure ([Hibbard 2005](#))
 PCI: Primary Communication Inventory ([Navran 1967](#))
 PHAST: Practical Hearing Aid Skills Test ([Desjardins 2009](#))
 PHL: person with hearing loss
 PIADS-A: Psychological Impact of Assistive Devices Scale ([Day 2002](#))
 PTA: pure tone audiogram, a standardised measure of hearing threshold
 QDS: Quantified Denver Scale of Communication ([Alpiner 1978](#); [Schow 1980](#))
 QoL: quality of life
 SADL: Satisfaction with Amplification in Daily Life ([Cox 1999](#))
 SD: standard deviation
 SF-12: Short form 12 ([Ware 1998](#))
 SF-36: Short form 36 ([Ware 1992](#))
 SIN: Speech In Noise
 SNHL: sensorineural hearing loss
 SO: significant other
 SSQ: Speech, Spatial and Qualities of hearing scale ([Gatehouse 2004](#))
 Sx: symptoms
 Tx: treatment/intervention group
 VA: (United States) Veterans Association
 VAS: visual analogue scale
 WHODAS 2.0: World Health Organization Disability Assessment Scale ([WHO 2001](#))
 WOCQ: Ways of Coping Questionnaire ([Folkman 1988](#))

Characteristics of excluded studies *[ordered by study ID]*

Study	Reason for exclusion
Aazh 2016	ALLOCATION: randomised PARTICIPANTS: participants were people who had been previously fitted with hearing aids but were not using them
Andersson 1998	Collated data on hearing aid use from 3 previous studies (see Andersson 1994 ; Andersson 1995 ; Andersson 1997). Where possible we have included the relevant data on hearing aid use as reported in the original studies
Bevilacqua 2013	ALLOCATION: randomised PARTICIPANTS: not all adults according to the definition given in this review and cannot extract the data for the adults separately
Boymans 2012	ALLOCATION: randomised PARTICIPANTS: new and experienced adult hearing aid users INTERVENTION: audiologist versus patient-driven fine tuning OUTCOME: speech perception measures
Cardemil 2014	ALLOCATION: unclear in published study but not randomised according to trial registration

(Continued)

Hallberg 1994	ALLOCATION: randomised PARTICIPANTS: not all hearing aid users and cannot extract the data for the hearing aid users independently
Hennig 2012	ALLOCATION: not randomised
Hickson 2007	ALLOCATION: randomised PARTICIPANTS: not all hearing aid users and cannot extract the data for the hearing aid users independently
Kuk 2014	ALLOCATION: not randomised
Lavie 2013	ALLOCATION: randomised PARTICIPANTS: adult new hearing aid users INTERVENTION: individual listening training OUTCOME: speech perception measures only
Montgomery 1984	ALLOCATION: randomised PARTICIPANTS: new adult hearing aid users INTERVENTION: individual versus group auditory training OUTCOME: speech perception measures
Norman 1994	ALLOCATION: not randomised
Preminger 2003	ALLOCATION: part-randomised and data not available for randomised participants only
Reber 2005	ALLOCATION: part-randomised and data not available for randomised participants only
Ruschel 2007	ALLOCATION: randomised PARTICIPANTS: new adult hearing aid users INTERVENTION: 5 sessions of auditory rehabilitation including guidance on communication strategy OUTCOME: non-validated questionnaire relating to ease of use and communication
Stecker 2006	ALLOCATION: randomised PARTICIPANTS: new adult hearing aid users INTERVENTION: 40 x 1-hour sessions of analytic auditory training OUTCOME: speech perception measures
Walden 1981	ALLOCATION: randomised PARTICIPANTS: new adult hearing aid users INTERVENTION: 7 hours of individual versus group auditory training OUTCOME: speech perception measures
Yueh 2010	ALLOCATION: randomised PARTICIPANTS: at the level of randomisation no participants were hearing aid users and only a proportion became hearing aid users

Characteristics of studies awaiting assessment *[ordered by study ID]*

Henshaw 2013

Methods	Randomised
Participants	Adult existing hearing aid users (50 to 74 years old) with mild to moderate hearing loss
Interventions	Working memory training
Outcomes	Self-reported hearing ability HHIE and GHABP at baseline, 2, 7 and 31 weeks
Notes	Study has been completed We made attempts to obtain study data for this review but data were unavailable in time for the update

Malmberg 2015

Methods	Randomised
Participants	Adults with hearing loss. Not clear if they were fitted with hearing aids for this study
Interventions	Auditory rehabilitation delivered over the internet
Outcomes	Self-reported hearing handicap HHIE, CSS, CPHI, HADs and IOI-HA at 5 weeks and 6 months
Notes	Study is reported as being completed but only implementation challenges so far reported

Characteristics of ongoing studies *[ordered by study ID]*

ISRCTN77340339

Trial name or title	A study on the effectiveness of a support programme (SUPR) for adult hearing aid users
Methods	Randomised controlled trial
Participants	Hearing-impaired hearing aid users, aged 50 and over, including both first-time hearing aid users and experienced hearing aid users, and their communication partners
Interventions	Hearing aid dispenser shops are randomly allocated to offer their customers either care as usual (hearing aid care) or hearing aid care including the SUPR support programme
Outcomes	Measurements (CPHI) are performed at the start of the study and after 6, 12 and 18 months
Starting date	February 2015
Contact information	Dr M Pronk Dept. of Otolaryngology - Head and Neck Surgery section Ear & Hearing

	Room pk2Y150 P.O. Box 7057 Amsterdam 1007 MB Netherlands
Notes	-

NCT02233361

Trial name or title	Use of hearing aids development and implementation of a counselling program for hearing aid users
Methods	Randomised controlled trial
Participants	New hearing aid users, aged 65 years and over
Interventions	The intervention group will be informed in advance of a follow-up appointment 6 months after they have received their hearing aid. They will know that support will be given and time-use of the hearing aid will be checked. Counselling on hearing aid use will be given The control group will not receive any information about a follow-up appointment. However, they will receive a notice on this after 6 months
Outcomes	Hours per day of hearing aid use (measured using data-logging) at 6 months post fitting
Starting date	January 2014
Contact information	jorunn.solheim@lds.no
Notes	-

NCT02264314

Trial name or title	Tele-educative program to improve adherence to the use of hearing aids in patients with hearing loss
Methods	Randomised controlled trial
Participants	New hearing aid users, aged 65 years and older
Interventions	The active arm consists of the implementation of an educational programme called Active Communication Education, consisting of 4 sessions with a trained rehabilitator. Participants will be also monitored by telephone headset use by personnel trained for 3 months The control arm will consist of the usual care received by these patients
Outcomes	Adherence to hearing aid use and hearing-related quality of life both assessed at 3, 6, 9 and 12 months post randomisation
Starting date	July 2012

NCT02264314 (Continued)

Contact information	Felipe Cardemil, MD, University of Chile
Notes	-

DATA AND ANALYSES

Comparison 1. Self-management support interventions versus control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Quality of life - short/medium-term	1	35	Mean Difference (IV, Random, 95% CI)	-9.10 [-21.33, 3.13]
2 Self-reported hearing handicap - short/medium-term	2	87	Mean Difference (IV, Random, 95% CI)	-12.80 [-23.11, -2.48]
3 Use of verbal communication strategy - short-term	1	52	Mean Difference (IV, Random, 95% CI)	0.72 [0.21, 1.23]

Comparison 2. Delivery system design interventions versus control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Adherence - short/medium-term	2	686	Risk Ratio (M-H, Random, 95% CI)	1.02 [0.99, 1.05]
2 Daily hours of hearing aid use - short/medium-term	4	700	Mean Difference (IV, Random, 95% CI)	-0.06 [-1.06, 0.95]
3 Adverse effects - long-term	1	98	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.50, 1.12]
4 Self-reported hearing handicap - short/medium-term	2	628	Mean Difference (IV, Random, 95% CI)	-0.70 [-5.22, 3.81]
5 Hearing aid benefit - short/medium-term	1	582	Mean Difference (IV, Random, 95% CI)	1.80 [-3.10, 6.70]
6 Use of verbal communication strategy	1	588	Mean Difference (IV, Random, 95% CI)	-0.10 [-0.40, 0.20]

Comparison 3. Combined SMS/DSD interventions versus control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Adherence - short/medium-term	1	167	Risk Ratio (M-H, Random, 95% CI)	1.06 [1.00, 1.12]
2 Daily hours of hearing aid use - long-term	2	69	Mean Difference (IV, Random, 95% CI)	0.04 [-0.64, 0.73]
3 Daily hours of hearing aid use - short/medium-term - SMS content	9	534	Mean Difference (IV, Random, 95% CI)	0.19 [-0.01, 0.40]
3.1 Advise	1	44	Mean Difference (IV, Random, 95% CI)	0.08 [-1.18, 1.34]
3.2 Activate - practical	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.3 Activate - symptoms	2	76	Mean Difference (IV, Random, 95% CI)	0.28 [-0.04, 0.59]

3.4 Activate - psychosocial	6	414	Mean Difference (IV, Random, 95% CI)	0.10 [-0.24, 0.45]
3.5 Assist	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.6 Agree	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Daily hours of hearing aid use - short/medium-term - DSD format	9	534	Mean Difference (IV, Random, 95% CI)	0.19 [-0.01, 0.40]
4.1 Face-to-face	5	163	Mean Difference (IV, Random, 95% CI)	0.24 [-0.06, 0.54]
4.2 Telephone	1	69	Mean Difference (IV, Random, 95% CI)	0.20 [-0.30, 0.70]
4.3 Booklet	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.4 Remote (online, PC, video/DVD)	3	302	Mean Difference (IV, Random, 95% CI)	0.08 [-0.55, 0.71]
5 Daily hours of hearing aid use - short/medium-term - DSD intensity	9	534	Mean Difference (IV, Random, 95% CI)	0.19 [-0.01, 0.40]
5.1 Low-intensity	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Medium-intensity	4	189	Mean Difference (IV, Random, 95% CI)	0.25 [-0.01, 0.51]
5.3 High-intensity	5	345	Mean Difference (IV, Random, 95% CI)	0.03 [-0.49, 0.55]
6 Quality of life - long-term	2	69	Mean Difference (IV, Random, 95% CI)	0.32 [-0.17, 0.80]
7 Quality of life - short/medium-term - SMS content	8	530	Std. Mean Difference (IV, Random, 95% CI)	0.02 [-0.15, 0.19]
7.1 Advise	1	48	Std. Mean Difference (IV, Random, 95% CI)	0.11 [-0.46, 0.67]
7.2 Activate - practical	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.3 Activate - symptoms	2	76	Std. Mean Difference (IV, Random, 95% CI)	-0.07 [-0.52, 0.38]
7.4 Activate - psychosocial	5	406	Std. Mean Difference (IV, Random, 95% CI)	0.04 [-0.18, 0.25]
7.5 Assist	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.6 Agree	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Quality of life - short/medium-term - DSD format	8	530	Std. Mean Difference (IV, Random, 95% CI)	0.02 [-0.15, 0.19]
8.1 Face-to-face	3	111	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.28, 0.47]
8.2 Telephone	1	69	Std. Mean Difference (IV, Random, 95% CI)	0.30 [-0.18, 0.77]
8.3 Booklet	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.4 Remote	4	350	Std. Mean Difference (IV, Random, 95% CI)	-0.05 [-0.26, 0.16]
9 Quality of life - short/medium-term - DSD intensity	8	530	Std. Mean Difference (IV, Random, 95% CI)	0.02 [-0.15, 0.19]
9.1 Low-intensity	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Medium-intensity	3	111	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.28, 0.47]
9.3 High-intensity	5	419	Std. Mean Difference (IV, Random, 95% CI)	0.00 [-0.19, 0.20]
10 Self-reported hearing handicap - long-term	3	88	Std. Mean Difference (IV, Random, 95% CI)	-0.31 [-1.06, 0.44]
10.1 Advise	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Activate - practical	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.3 Activate - symptoms	2	69	Std. Mean Difference (IV, Random, 95% CI)	0.04 [-0.43, 0.51]
10.4 Activate - psychosocial	1	19	Std. Mean Difference (IV, Random, 95% CI)	-1.27 [-2.28, -0.26]
10.5 Assist	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.6 Agree	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Self-reported hearing handicap - short/medium-term - SMS content	15	728	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.48, -0.04]
11.1 Advise	4	153	Std. Mean Difference (IV, Random, 95% CI)	-0.27 [-0.59, 0.05]

11.2 Activate - practical	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.3 Activate - symptoms	3	89	Std. Mean Difference (IV, Random, 95% CI)	-0.34 [-0.76, 0.08]
11.4 Activate - psychosocial	8	486	Std. Mean Difference (IV, Random, 95% CI)	-0.24 [-0.61, 0.13]
11.5 Assist	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.6 Agree	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Self-reported hearing handicap - short/medium-term - DSD format	15	728	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.48, -0.04]
12.1 Face-to-face	9	289	Std. Mean Difference (IV, Random, 95% CI)	-0.16 [-0.39, 0.07]
12.2 Telephone	1	69	Std. Mean Difference (IV, Random, 95% CI)	-0.83 [-1.33, -0.34]
12.3 Booklet	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.4 Remote	5	370	Std. Mean Difference (IV, Random, 95% CI)	-0.28 [-0.72, 0.16]
13 Self-reported hearing handicap - short/medium-term - DSD intensity	15	728	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.48, -0.04]
13.1 Low-intensity	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Medium-intensity	7	249	Std. Mean Difference (IV, Random, 95% CI)	-0.35 [-0.60, -0.10]
13.3 High-intensity	8	479	Std. Mean Difference (IV, Random, 95% CI)	-0.17 [-0.52, 0.17]
14 Hearing aid benefit - long-term	2	69	Mean Difference (IV, Random, 95% CI)	0.30 [0.02, 0.58]
15 Hearing aid benefit - short/medium-term - SMS content	7	361	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.15, 0.36]
15.1 Advise	2	92	Std. Mean Difference (IV, Random, 95% CI)	-0.14 [-1.10, 0.83]
15.2 Activate - practical	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.3 Activate - symptoms	2	76	Std. Mean Difference (IV, Random, 95% CI)	0.17 [-0.28, 0.62]
15.4 Activate - psychosocial	3	193	Std. Mean Difference (IV, Random, 95% CI)	0.22 [-0.07, 0.50]
15.5 Assist	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.6 Agree	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Hearing aid benefit - short/medium-term - DSD format	7	361	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.15, 0.36]
16.1 Face-to-face	3	120	Std. Mean Difference (IV, Random, 95% CI)	0.24 [-0.13, 0.60]
16.2 Telephone	1	69	Std. Mean Difference (IV, Random, 95% CI)	0.38 [-0.09, 0.86]
16.3 Booklet	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.4 Remote	3	172	Std. Mean Difference (IV, Random, 95% CI)	-0.12 [-0.63, 0.39]
17 Hearing aid benefit - short/medium-term - DSD intensity	7	361	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.15, 0.36]
17.1 Low-intensity	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Medium-intensity	3	120	Std. Mean Difference (IV, Random, 95% CI)	0.24 [-0.13, 0.60]
17.3 High-intensity	4	241	Std. Mean Difference (IV, Random, 95% CI)	0.01 [-0.41, 0.43]
18 Use of verbal communication strategy - long-term	1	34	Mean Difference (IV, Random, 95% CI)	0.30 [-0.20, 0.80]
19 Use of verbal communication strategy - short/medium-term - SMS content	4	223	Mean Difference (IV, Random, 95% CI)	0.45 [0.15, 0.74]
19.1 Advise	1	115	Mean Difference (IV, Random, 95% CI)	0.25 [-0.07, 0.57]
19.2 Activate - practical	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.3 Activate - symptoms	1	37	Mean Difference (IV, Random, 95% CI)	0.40 [-0.06, 0.86]
19.4 Activate - psychosocial	2	71	Mean Difference (IV, Random, 95% CI)	0.70 [0.01, 1.39]
19.5 Assist	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.6 Agree	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

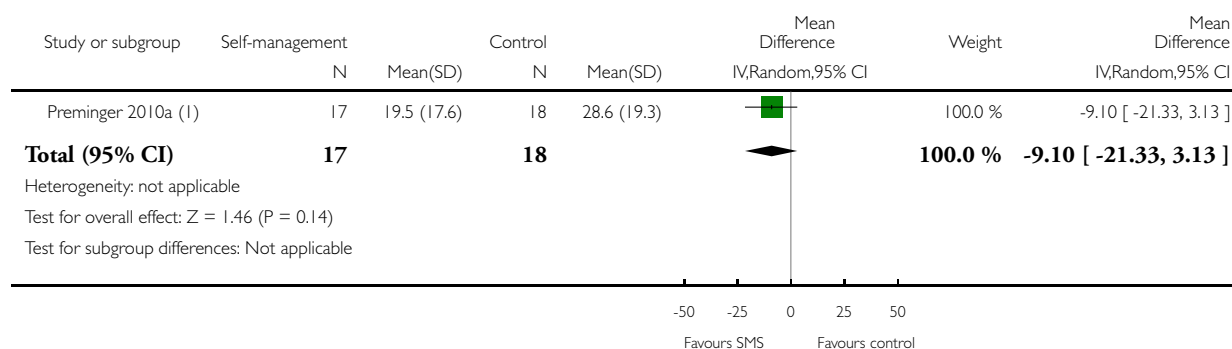
20 Use of verbal communication strategy - short/medium-term - DSD intensity	4	223	Mean Difference (IV, Random, 95% CI)	0.45 [0.15, 0.74]
20.1 Low-intensity	1	115	Mean Difference (IV, Random, 95% CI)	0.25 [-0.07, 0.57]
20.2 Medium-intensity	2	89	Mean Difference (IV, Random, 95% CI)	0.40 [0.07, 0.72]
20.3 High-intensity	1	19	Mean Difference (IV, Random, 95% CI)	1.1 [0.43, 1.77]

Analysis 1.1. Comparison 1 Self-management support interventions versus control, Outcome 1 Quality of life - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 1 Self-management support interventions versus control

Outcome: 1 Quality of life - short/medium-term



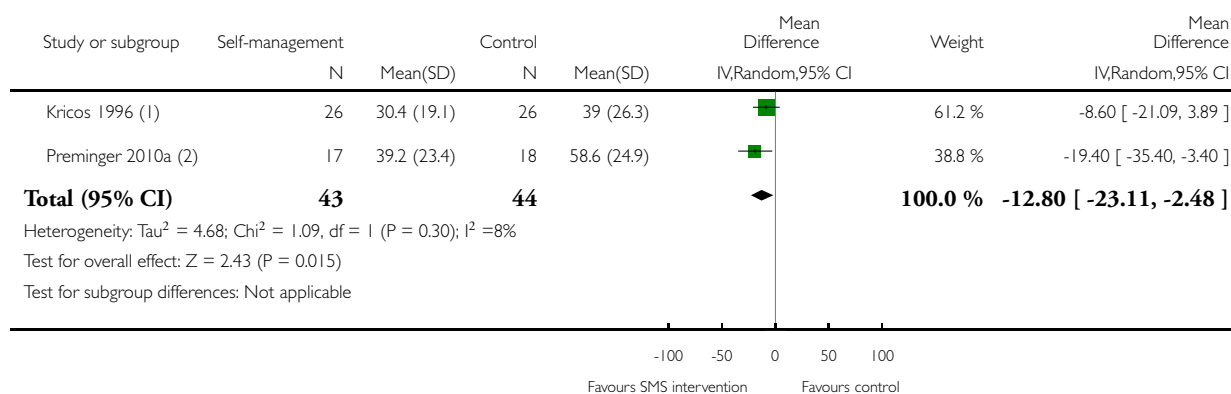
(1) Medium term data, WHO-DAS II - lower score = better QoL

Analysis 1.2. Comparison 1 Self-management support interventions versus control, Outcome 2 Self-reported hearing handicap - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 1 Self-management support interventions versus control

Outcome: 2 Self-reported hearing handicap - short/medium-term



(1) High risk of bias

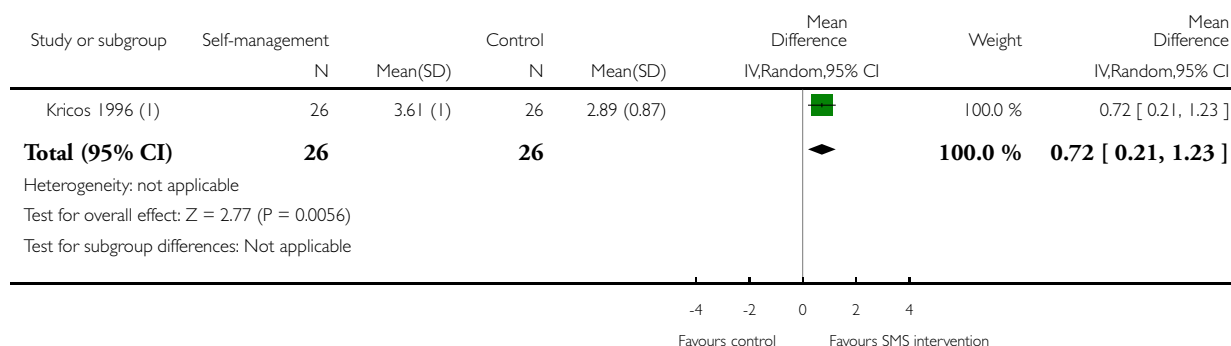
(2) Medium term data, high risk of bias

Analysis 1.3. Comparison 1 Self-management support interventions versus control, Outcome 3 Use of verbal communication strategy - short-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 1 Self-management support interventions versus control

Outcome: 3 Use of verbal communication strategy - short-term



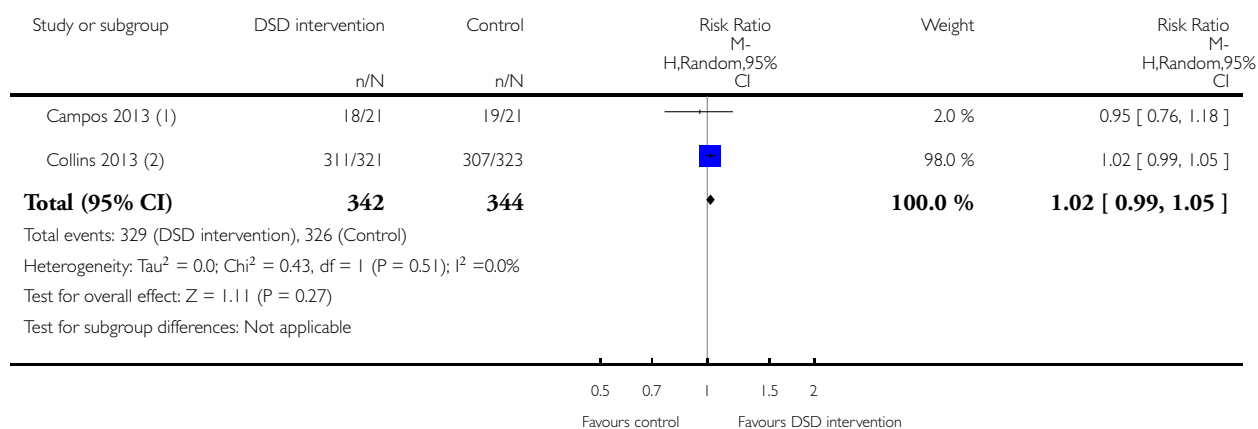
(1) High risk of bias

Analysis 2.1. Comparison 2 Delivery system design interventions versus control, Outcome 1 Adherence - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 2 Delivery system design interventions versus control

Outcome: 1 Adherence - short/medium-term



(1) Remote online fitting vs face-to-face fitting (short term)

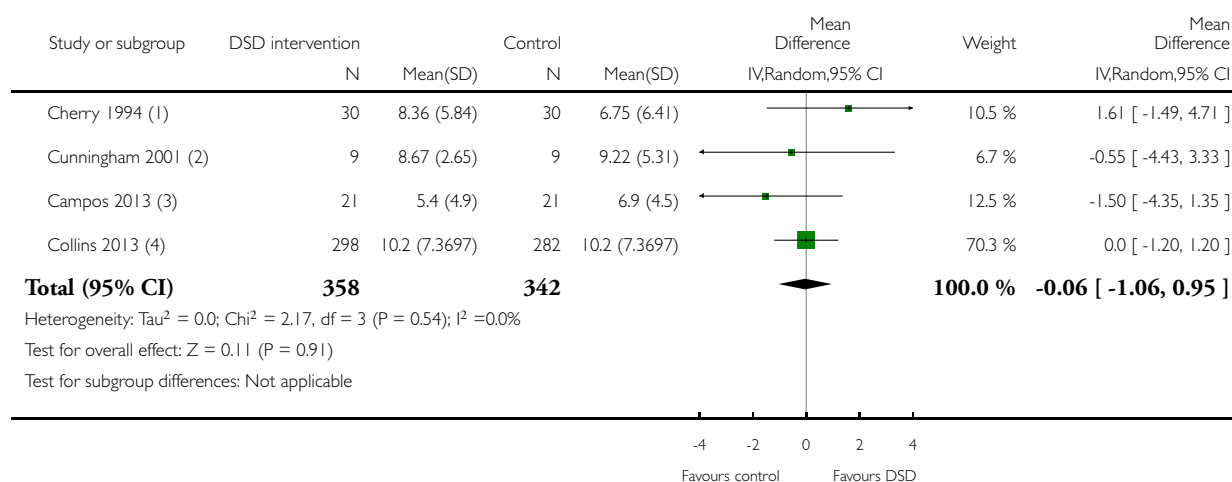
(2) Group vs individual fitting (medium term)

Analysis 2.2. Comparison 2 Delivery system design interventions versus control, Outcome 2 Daily hours of hearing aid use - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 2 Delivery system design interventions versus control

Outcome: 2 Daily hours of hearing aid use - short/medium-term



(1) Medium term data

(2) Medium term data

(3) Short term data - measured with data-logging

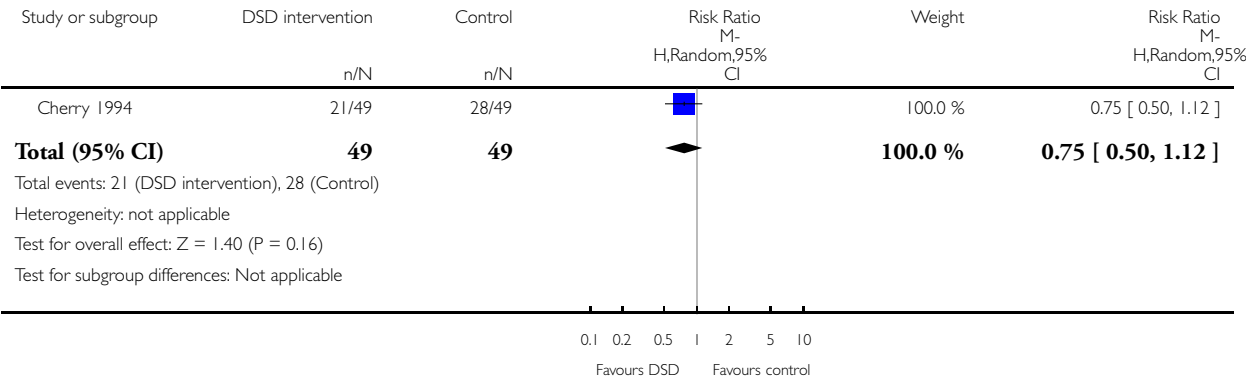
(4) Medium term data - Standard deviations calculated from mean difference and CIs reported in study

Analysis 2.3. Comparison 2 Delivery system design interventions versus control, Outcome 3 Adverse effects - long-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 2 Delivery system design interventions versus control

Outcome: 3 Adverse effects - long-term

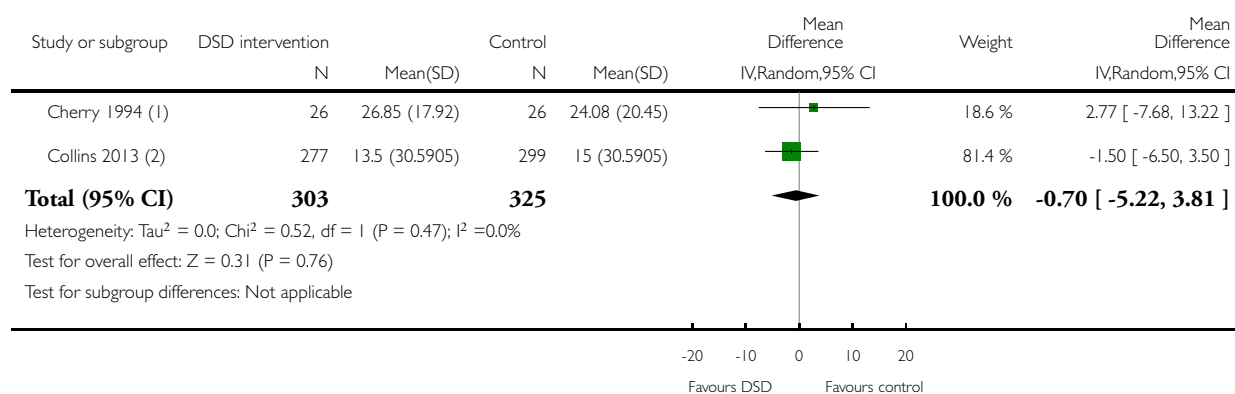


Analysis 2.4. Comparison 2 Delivery system design interventions versus control, Outcome 4 Self-reported hearing handicap - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 2 Delivery system design interventions versus control

Outcome: 4 Self-reported hearing handicap - short/medium-term



(1) Medium term data

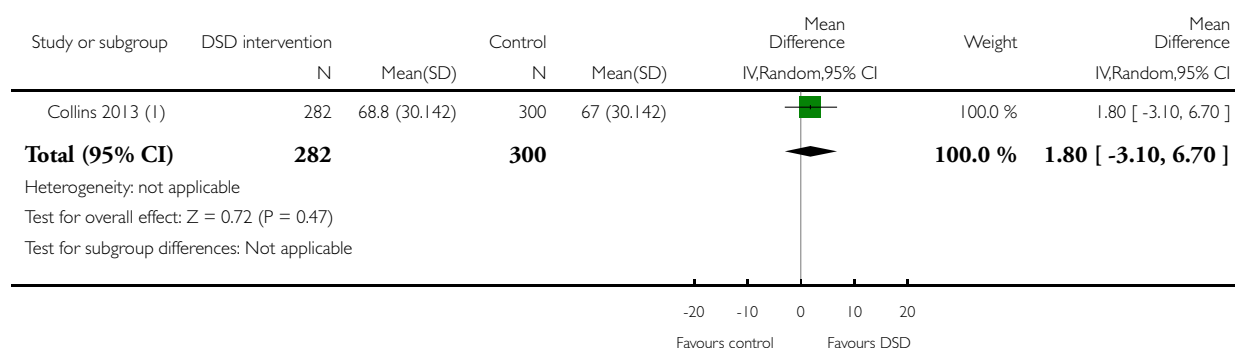
(2) Medium term data - SDs calculated from reported CIs and p-value

Analysis 2.5. Comparison 2 Delivery system design interventions versus control, Outcome 5 Hearing aid benefit - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 2 Delivery system design interventions versus control

Outcome: 5 Hearing aid benefit - short/medium-term



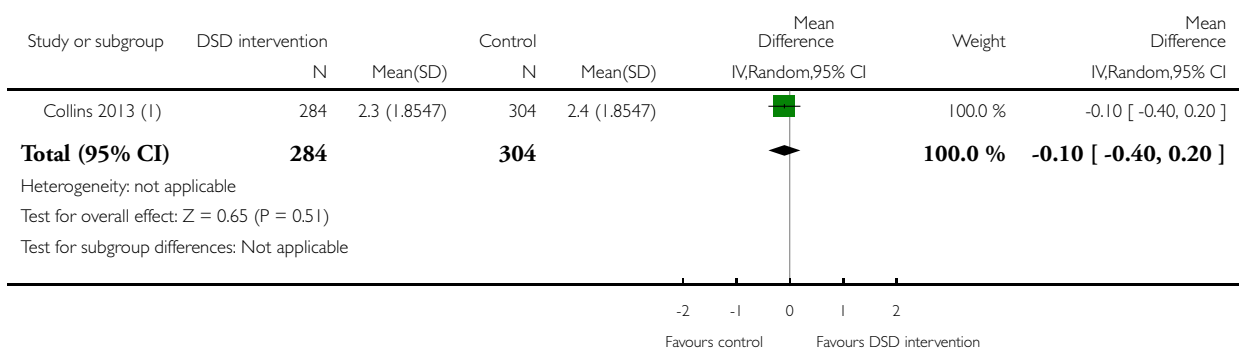
(I) Measured using Outer EAR, SDs calculated from p-value and confidence intervals

Analysis 2.6. Comparison 2 Delivery system design interventions versus control, Outcome 6 Use of verbal communication strategy.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 2 Delivery system design interventions versus control

Outcome: 6 Use of verbal communication strategy



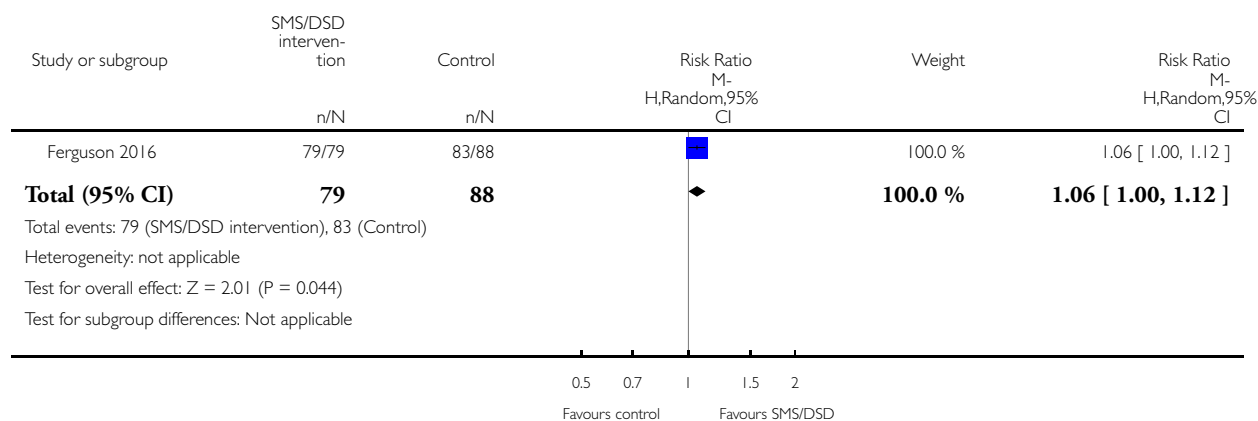
(I) SDs calculated based on p-value and CIs

Analysis 3.1. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 1 Adherence - short/medium-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 1 Adherence - short/medium-term

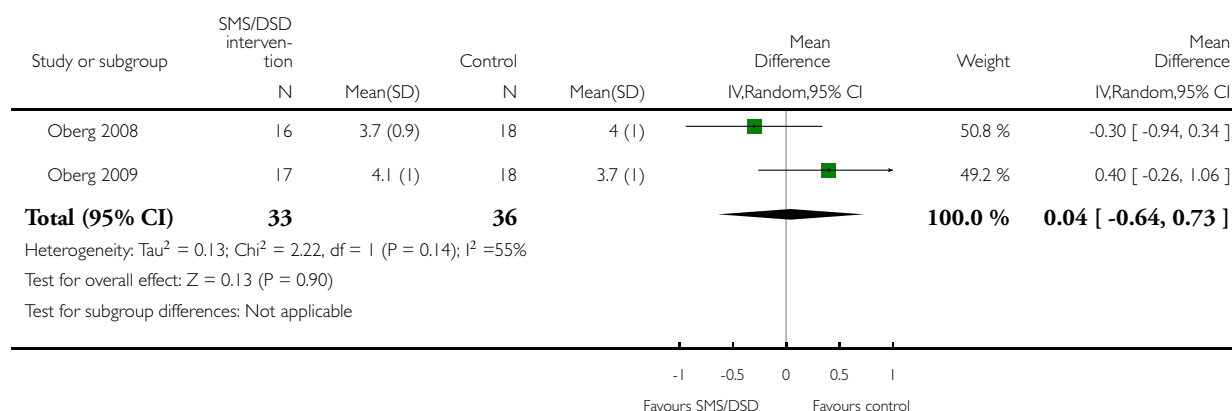


Analysis 3.2. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 2 Daily hours of hearing aid use - long-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 2 Daily hours of hearing aid use - long-term

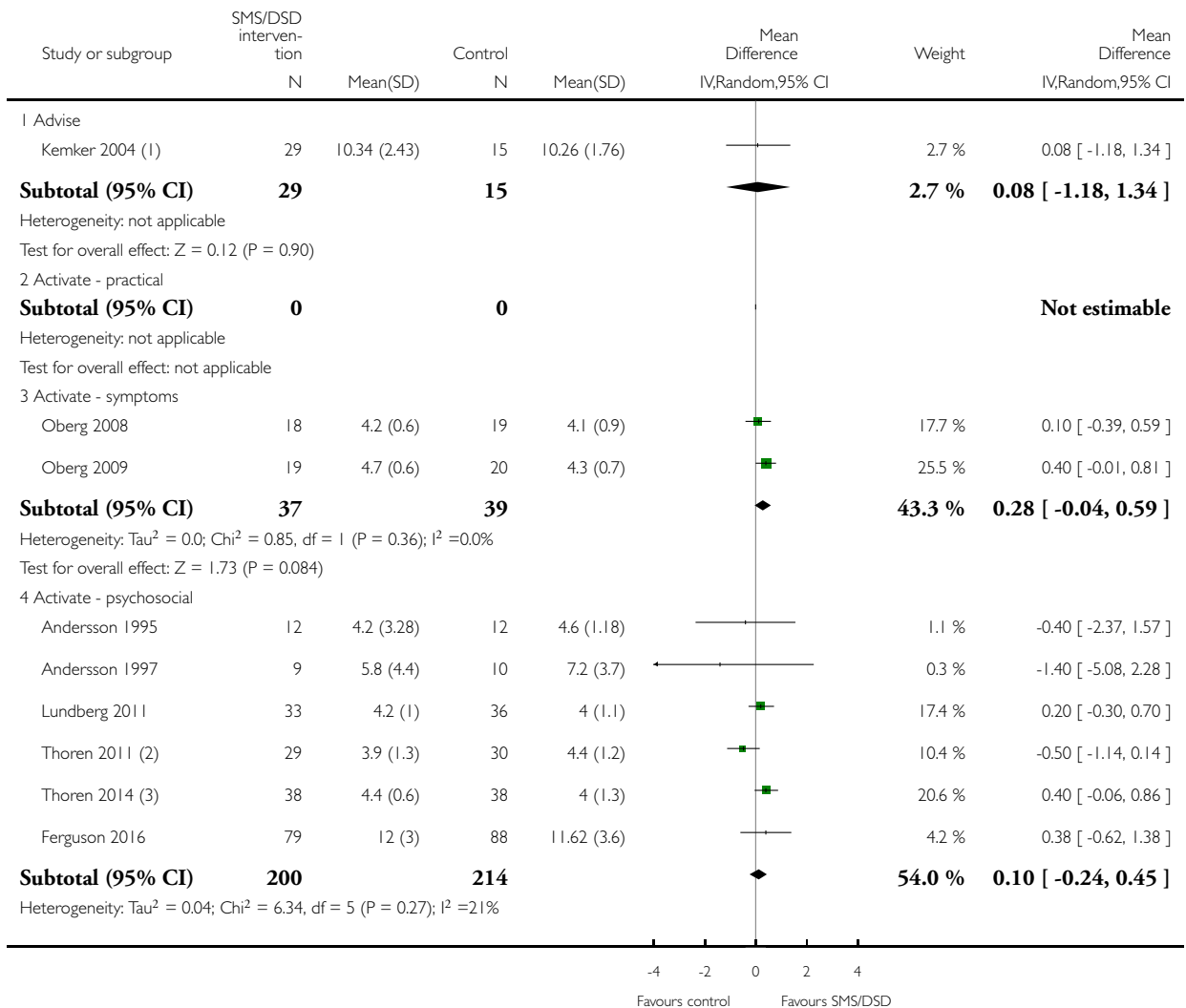


Analysis 3.3. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 3 Daily hours of hearing aid use - short/medium-term - SMS content.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

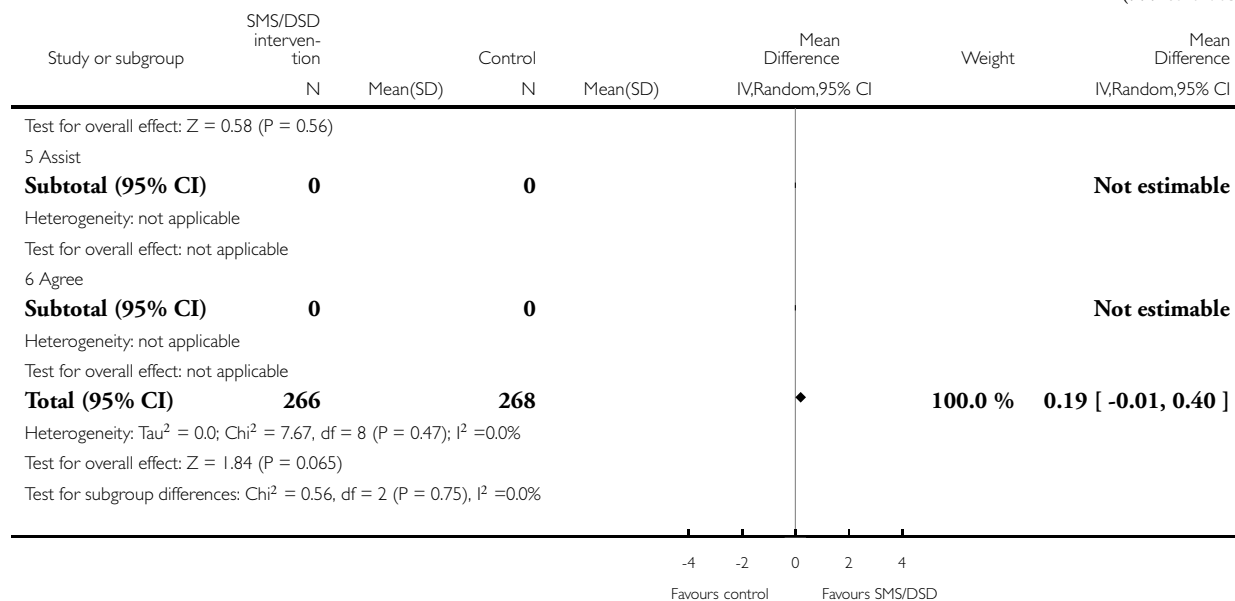
Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 3 Daily hours of hearing aid use - short/medium-term - SMS content



(Continued ...)

(... Continued)



(1) Combined pre and post fitting orientation, converted from % day worn based on a 12 hour day

(2) Medium term data

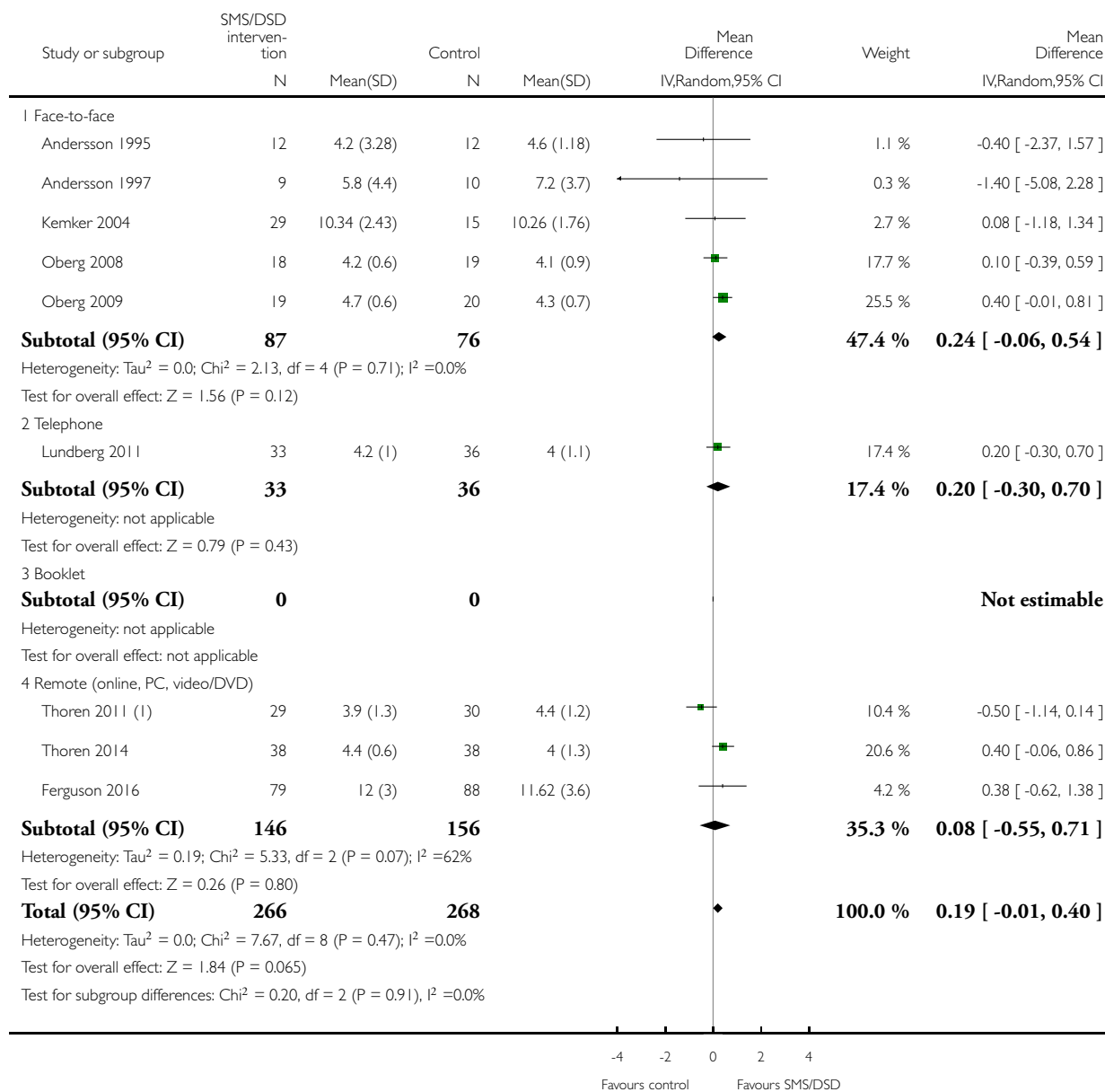
(3) Medium term data

Analysis 3.4. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 4 Daily hours of hearing aid use - short/medium-term - DSD format.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 4 Daily hours of hearing aid use - short/medium-term - DSD format



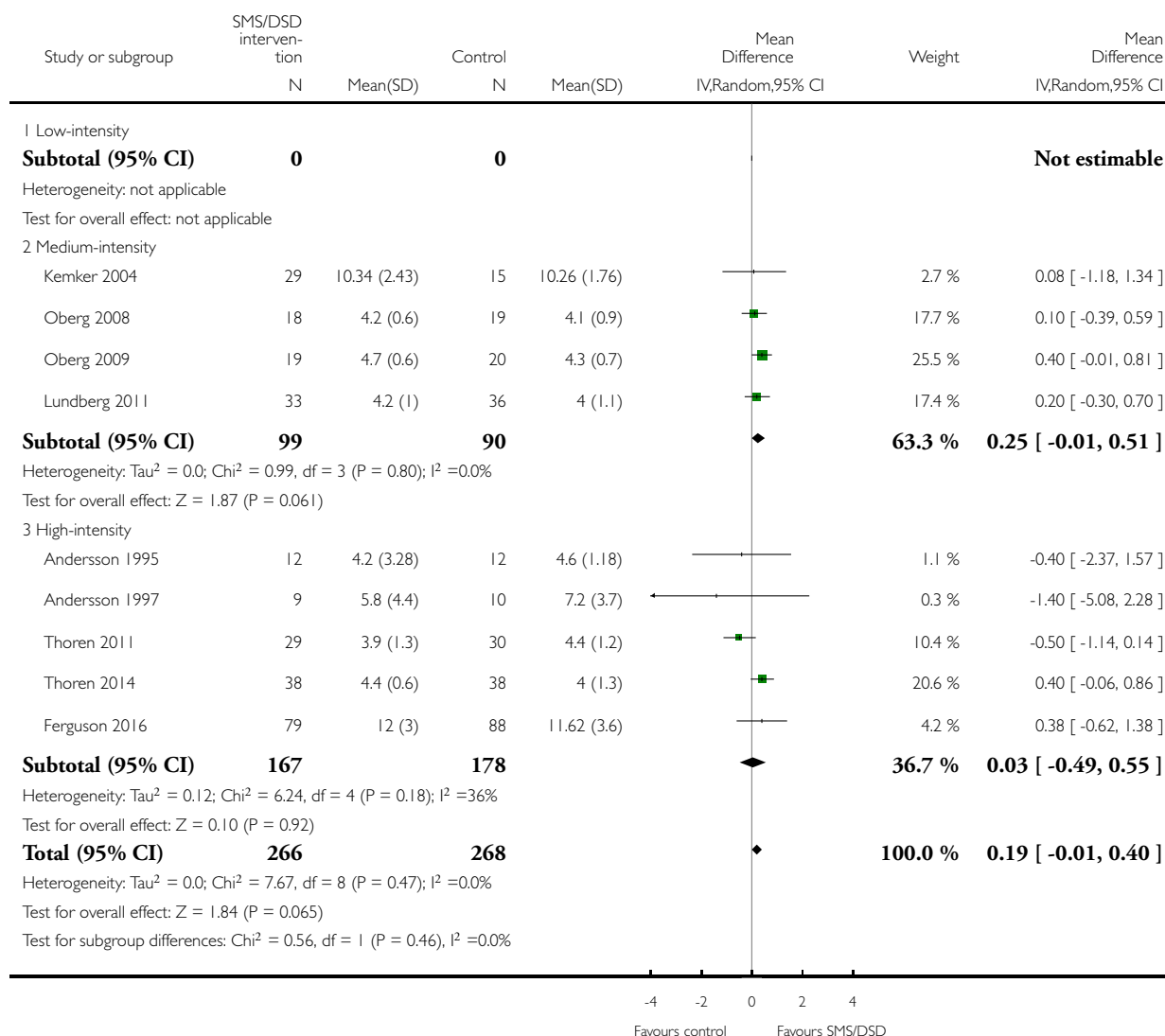
(1) Medium term data

Analysis 3.5. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 5 Daily hours of hearing aid use - short/medium-term - DSD intensity.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 5 Daily hours of hearing aid use - short/medium-term - DSD intensity

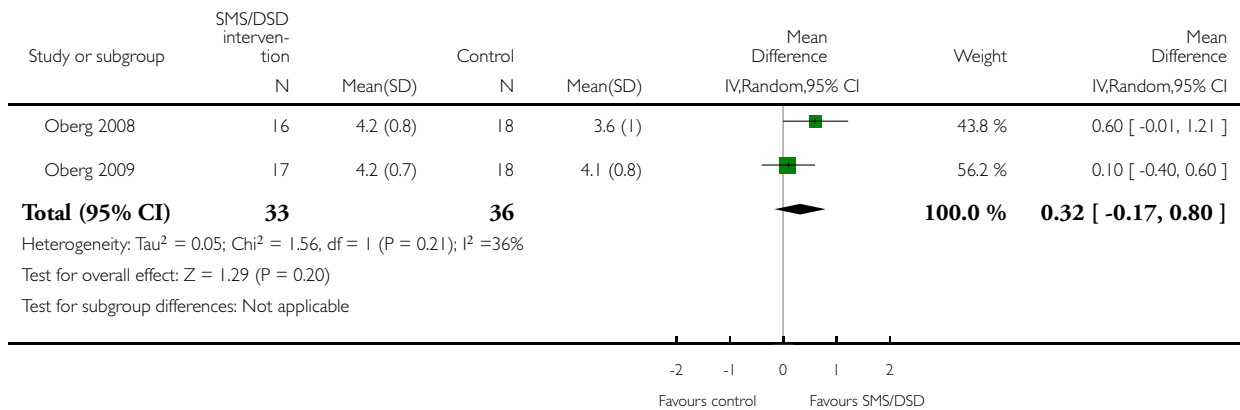


Analysis 3.6. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 6 Quality of life - long-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 6 Quality of life - long-term

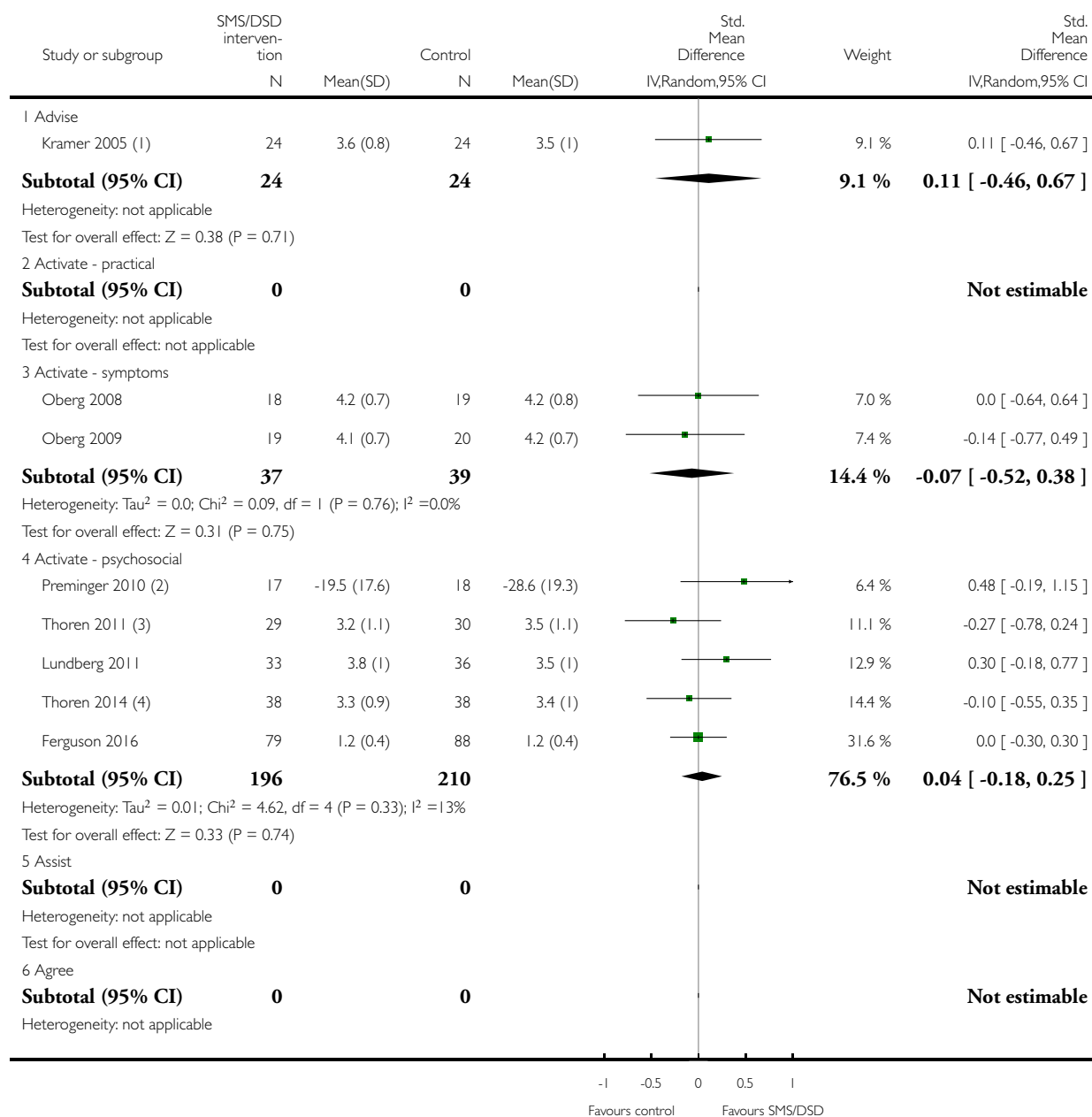


Analysis 3.7. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 7 Quality of life - short/medium-term - SMS content.

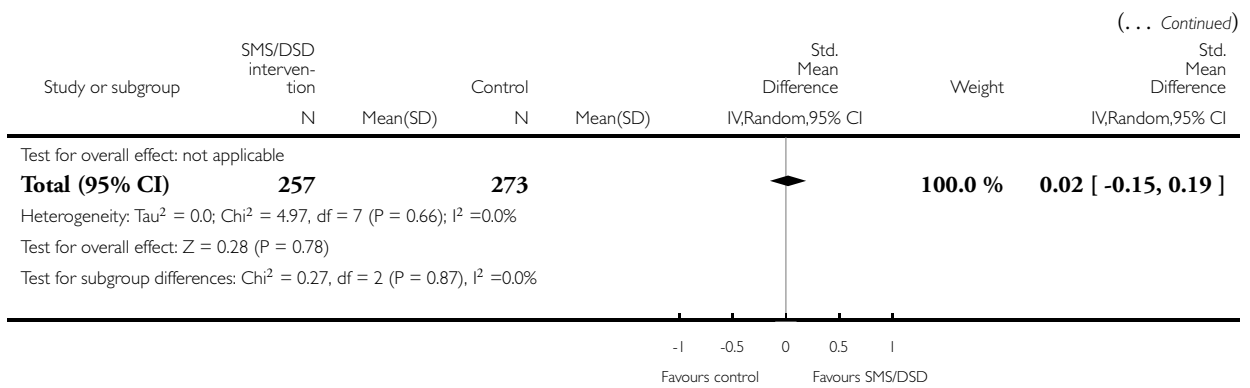
Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 7 Quality of life - short/medium-term - SMS content



(Continued ...)



(1) Medium term data

(2) Medium term data

(3) Medium term data

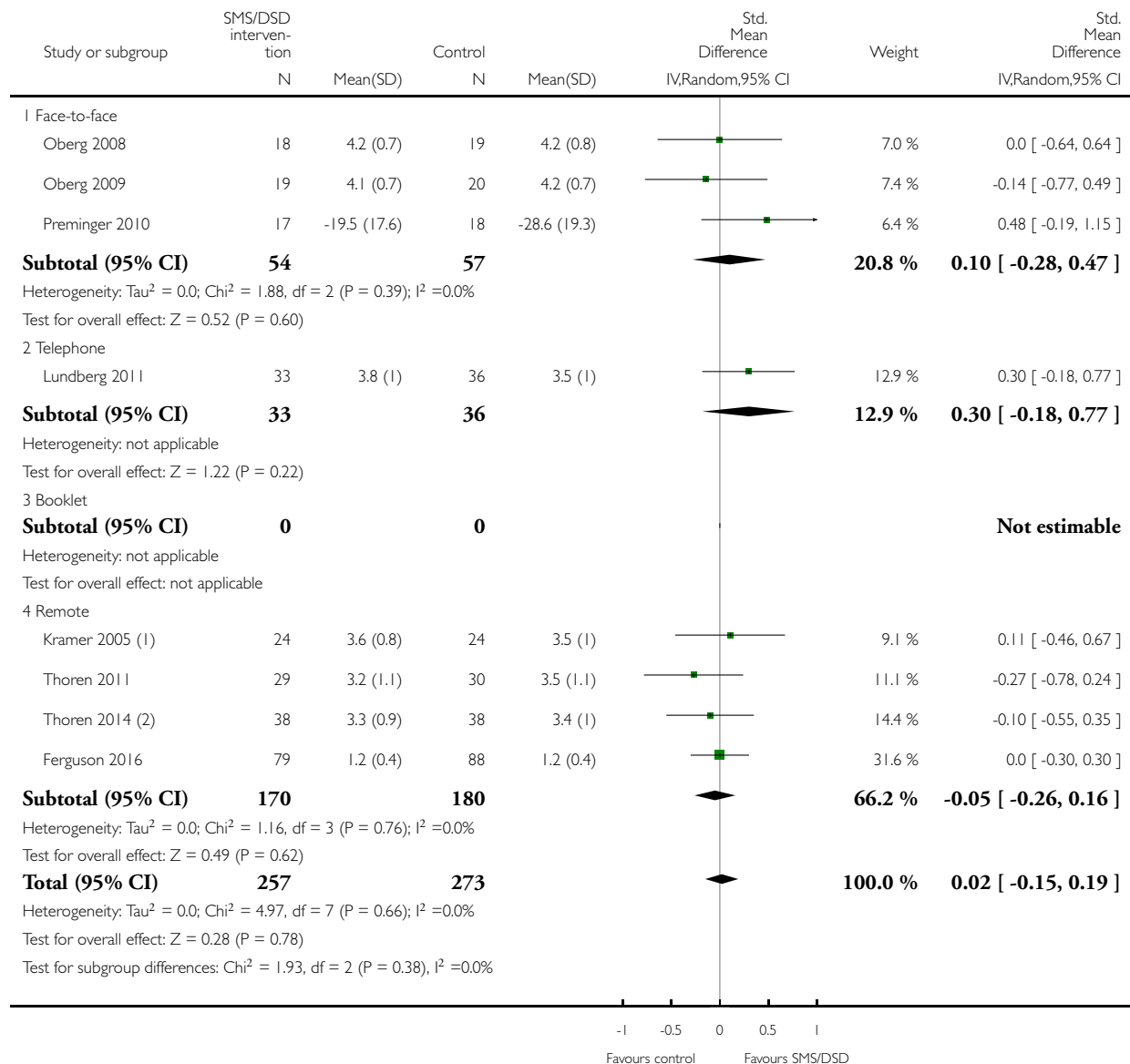
(4) Medium term data

Analysis 3.8. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 8 Quality of life - short/medium-term - DSD format.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 8 Quality of life - short/medium-term - DSD format



(1) Medium term

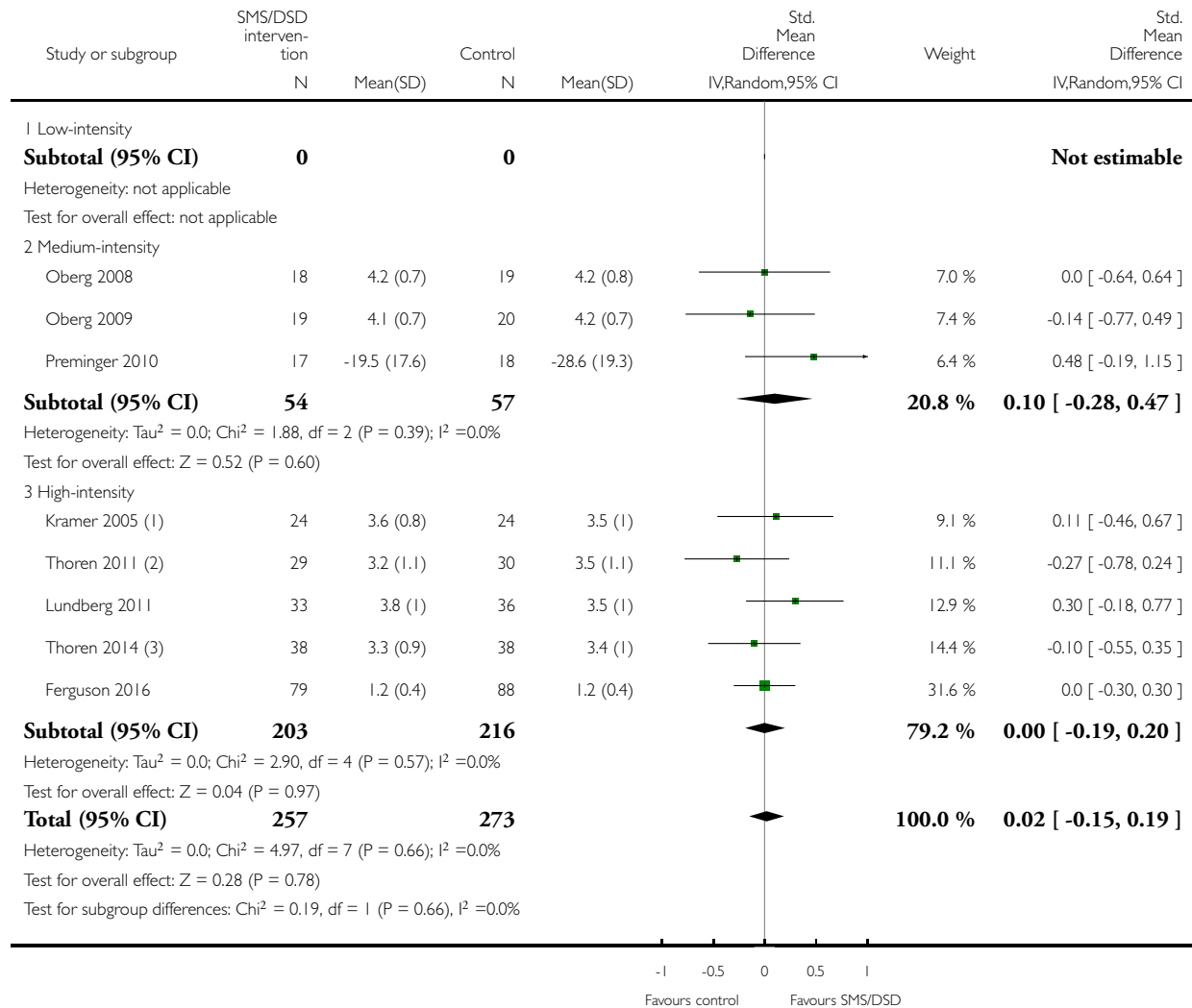
(2) Medium term data

Analysis 3.9. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 9 Quality of life - short/medium-term - DSD intensity.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 9 Quality of life - short/medium-term - DSD intensity



(1) Medium term

(2) Medium term data

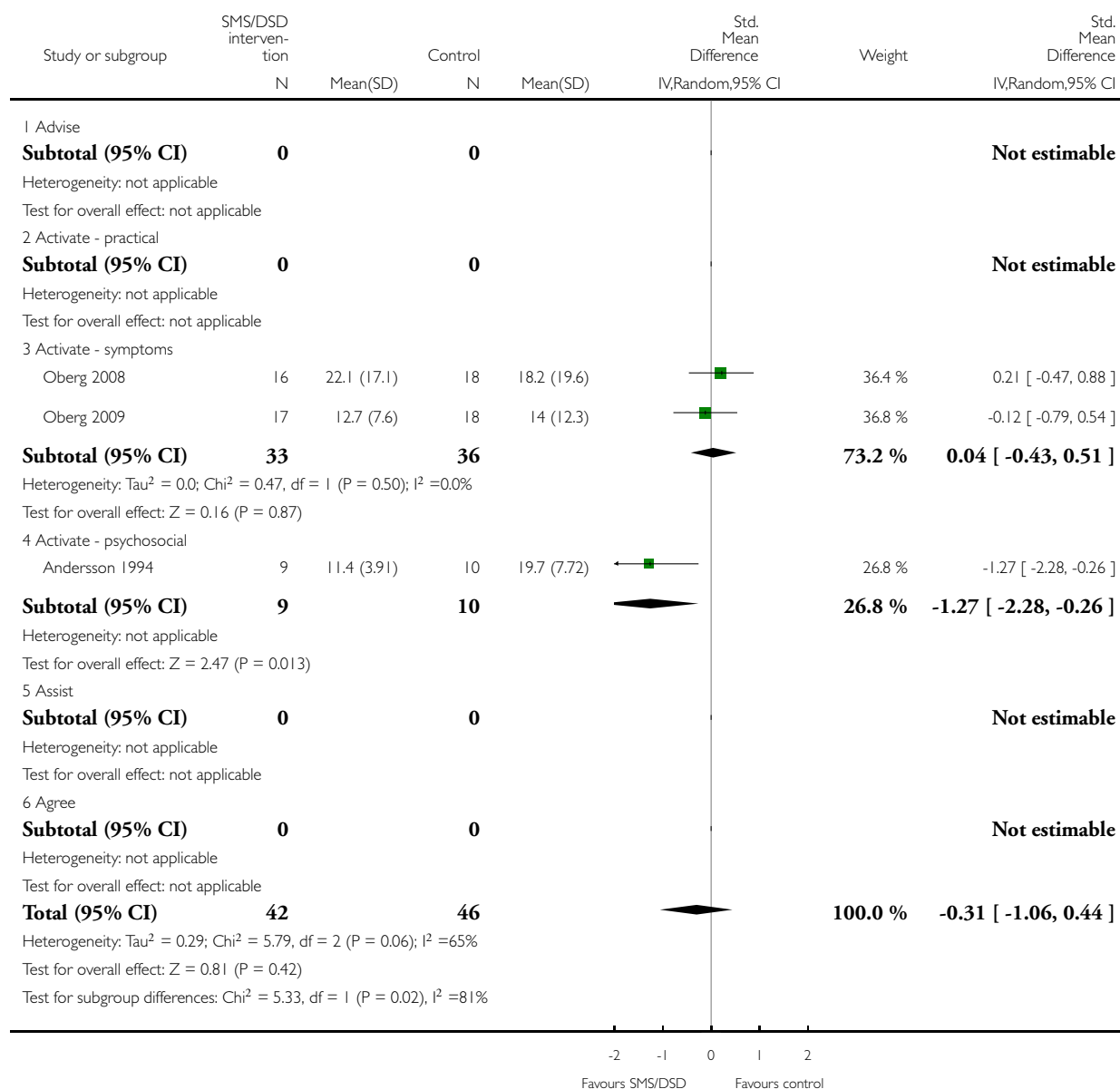
(3) Medium term data

Analysis 3.10. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 10 Self-reported hearing handicap - long-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 10 Self-reported hearing handicap - long-term

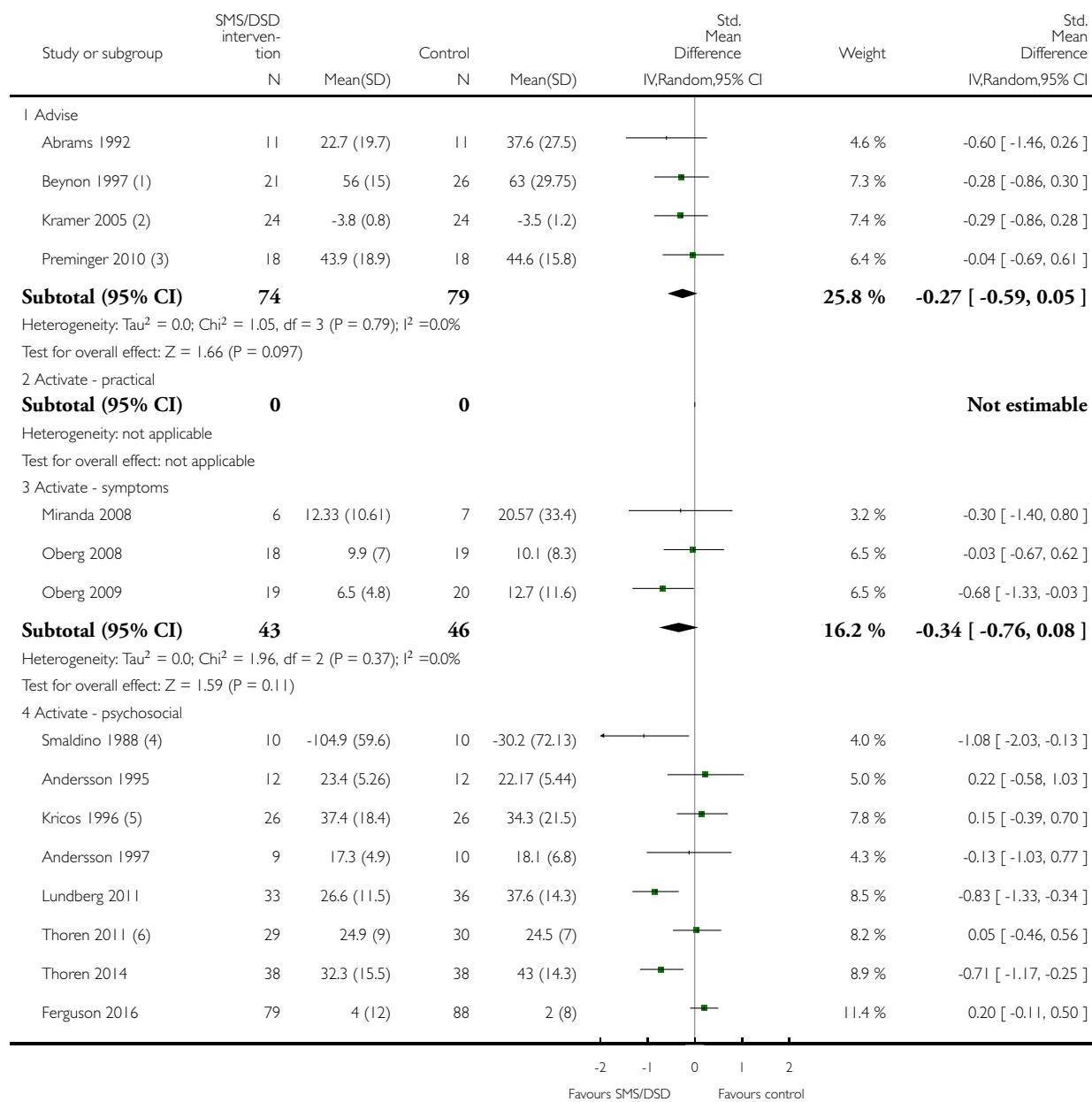


Analysis 3.1.1. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 11 Self-reported hearing handicap - short/medium-term - SMS content.

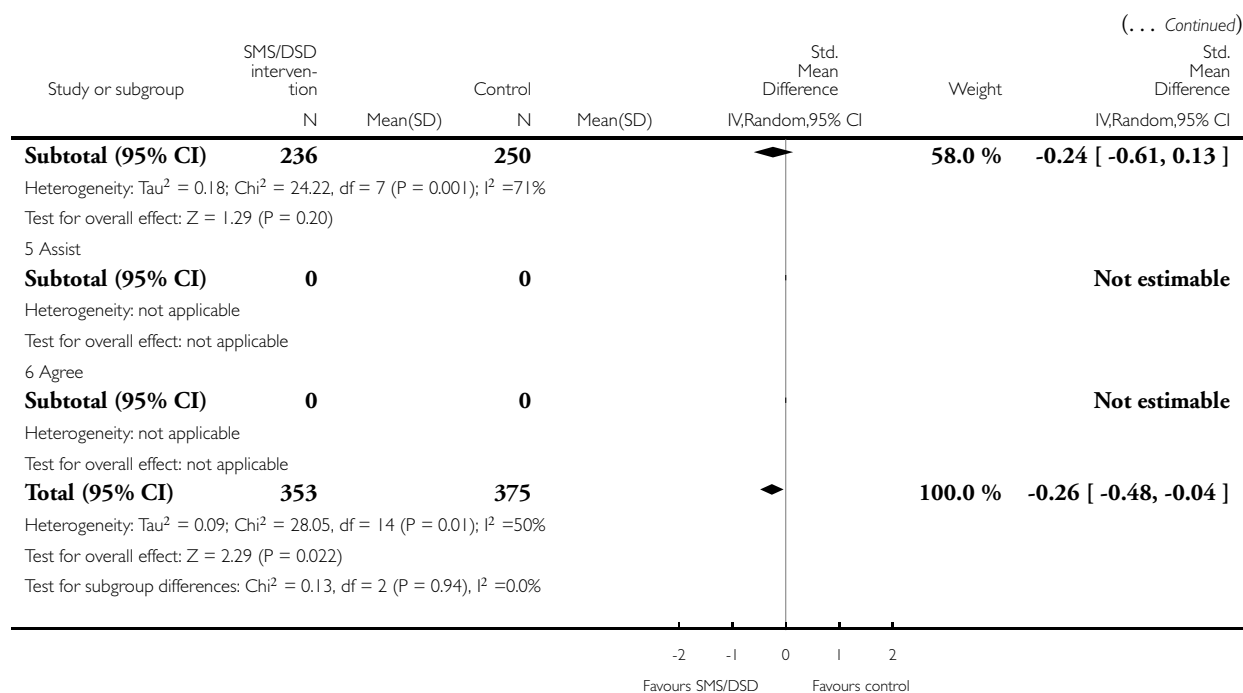
Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 11 Self-reported hearing handicap - short/medium-term - SMS content



(Continued ...)



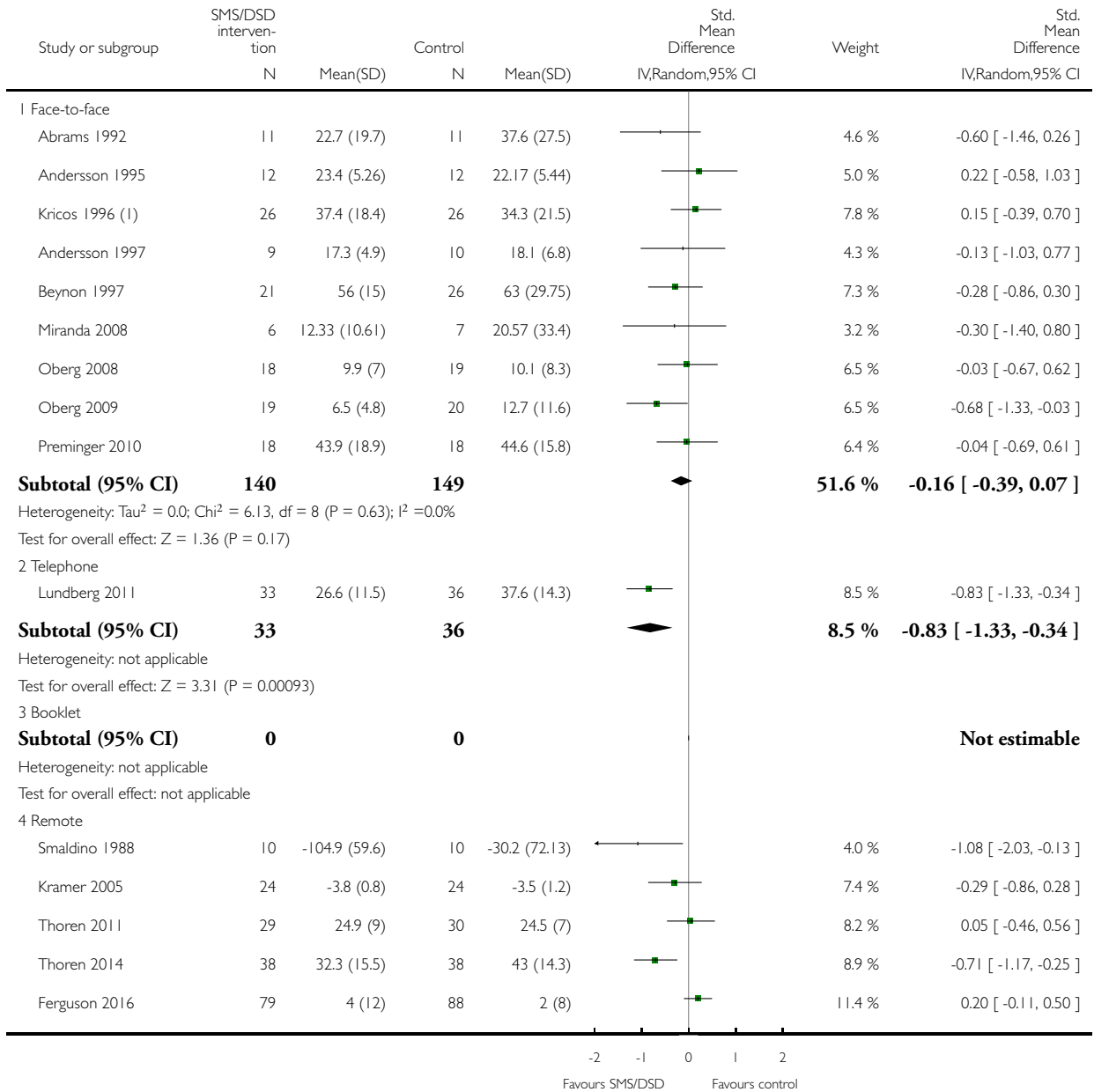
- (1) Medium term data
- (2) Medium term data
- (3) Medium term data
- (4) Change scores
- (5) Active listening intervention versus standard care
- (6) Medium term data

Analysis 3.12. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 12 Self-reported hearing handicap - short/medium-term - DSD format.

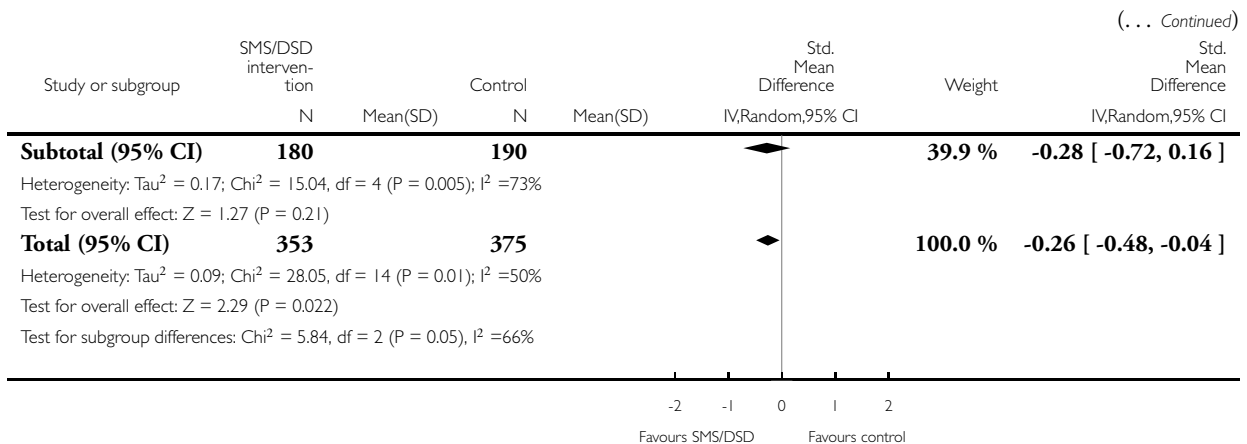
Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 12 Self-reported hearing handicap - short/medium-term - DSD format



(Continued ...)



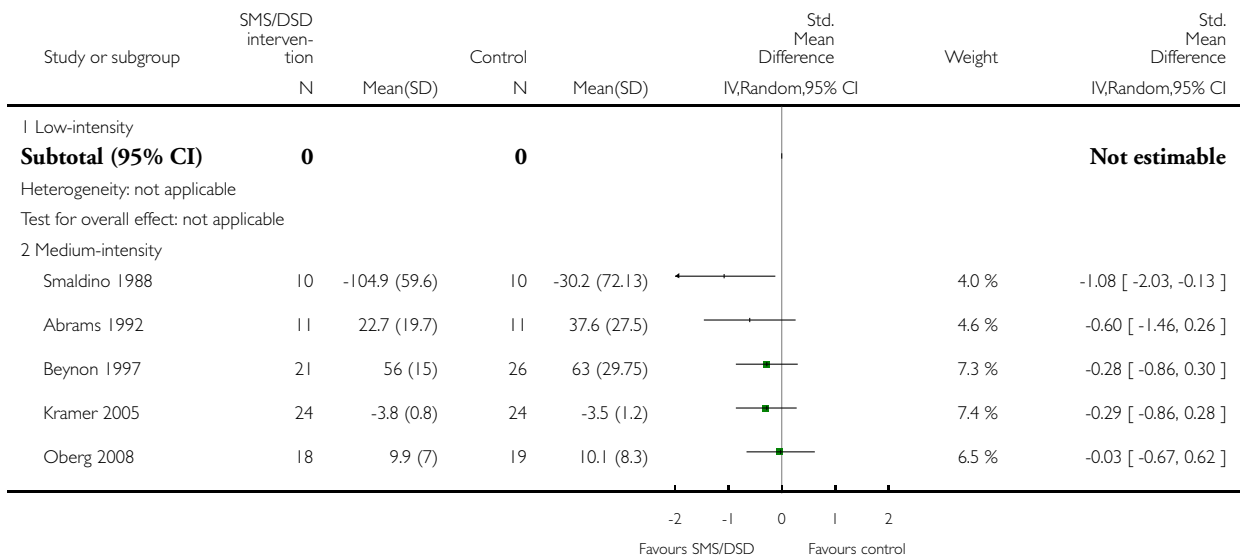
(I) Active listening intervention

Analysis 3.13. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 13 Self-reported hearing handicap - short/medium-term - DSD intensity.

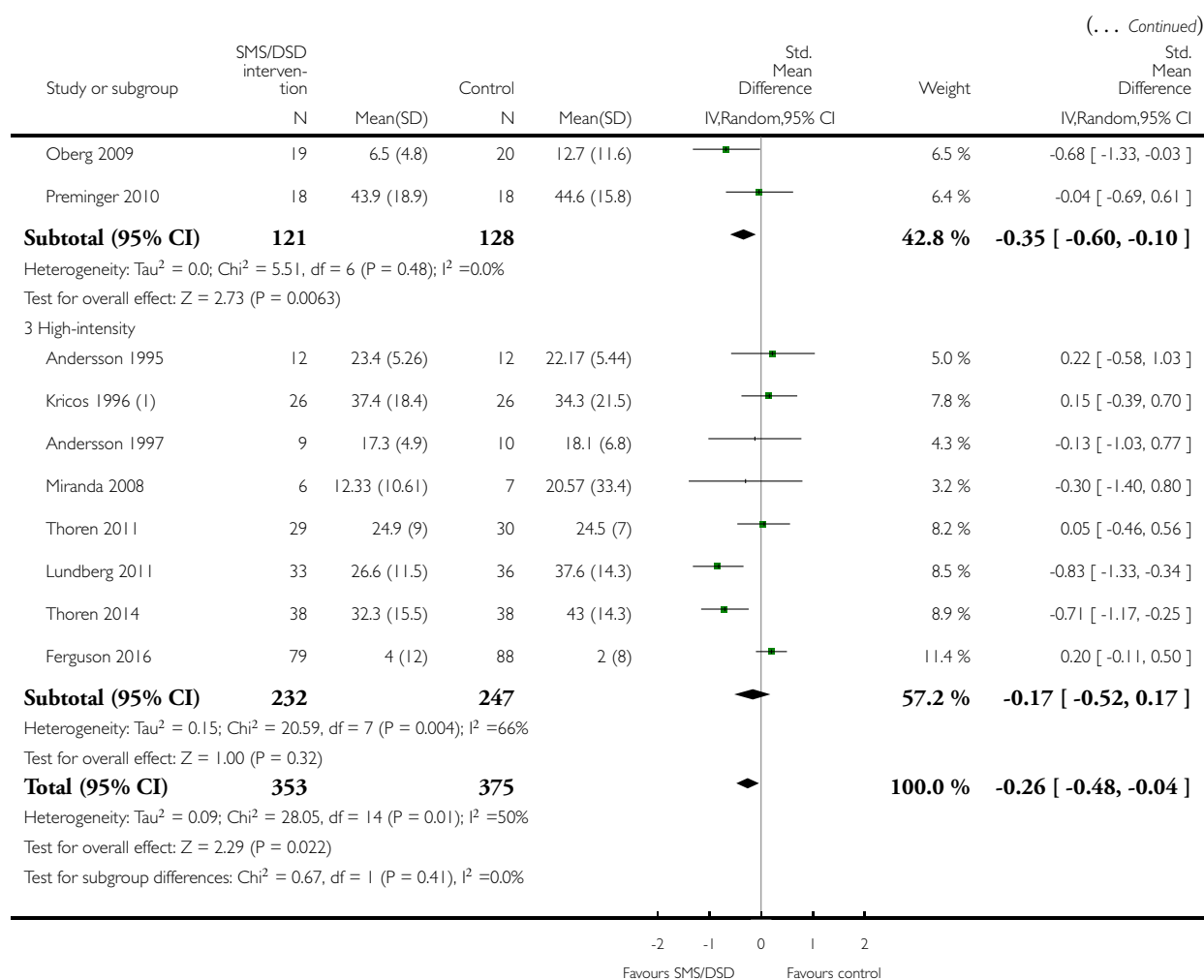
Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 13 Self-reported hearing handicap - short/medium-term - DSD intensity



(Continued ...)



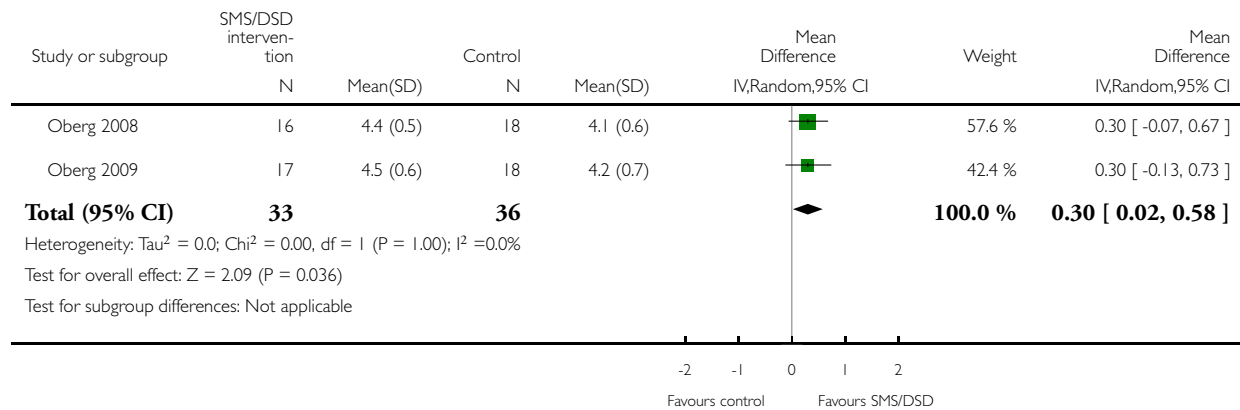
(1) Active listening intervention

Analysis 3.14. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 14 Hearing aid benefit - long-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 14 Hearing aid benefit - long-term

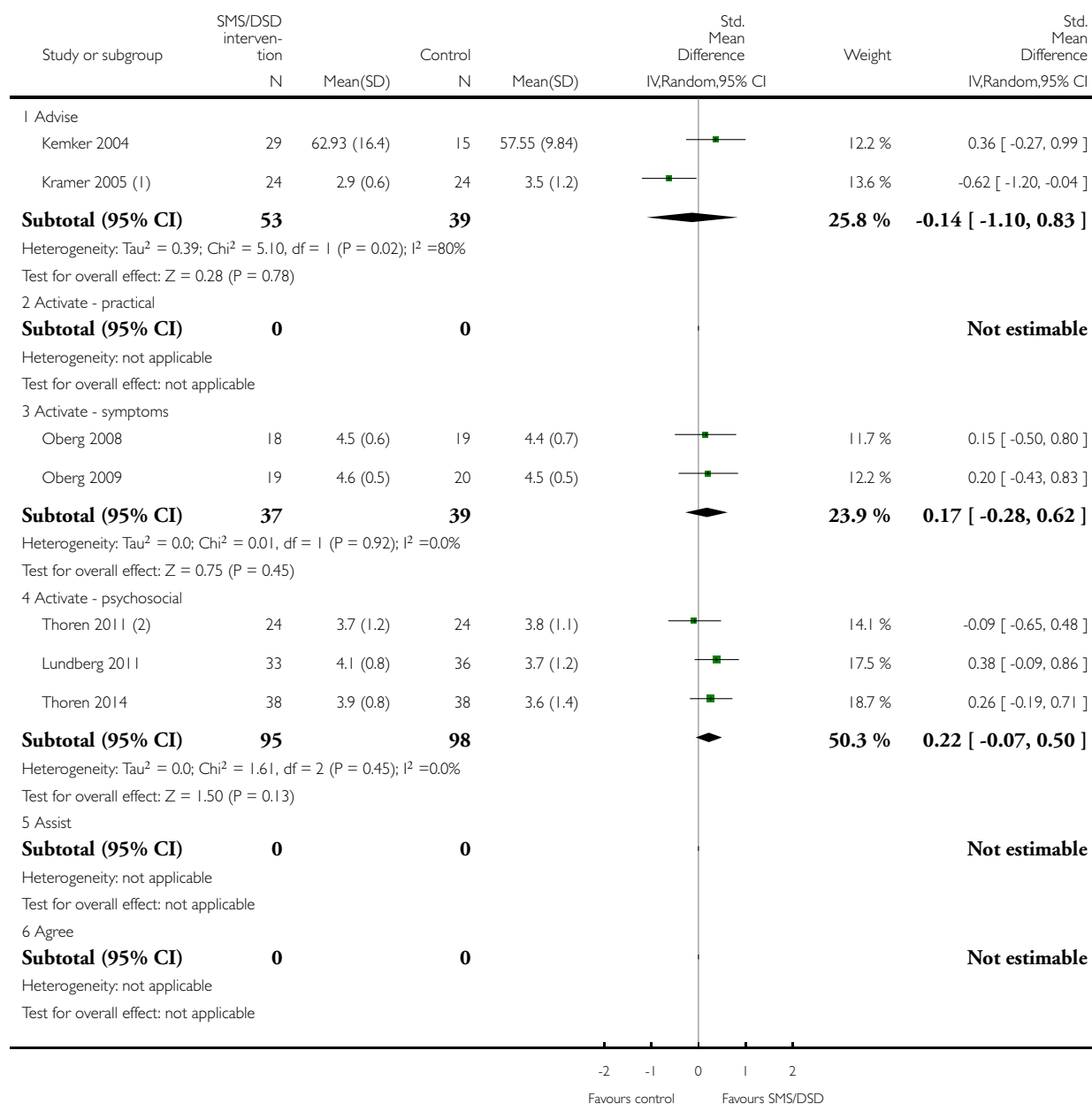


Analysis 3.15. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 15 Hearing aid benefit - short/medium-term - SMS content.

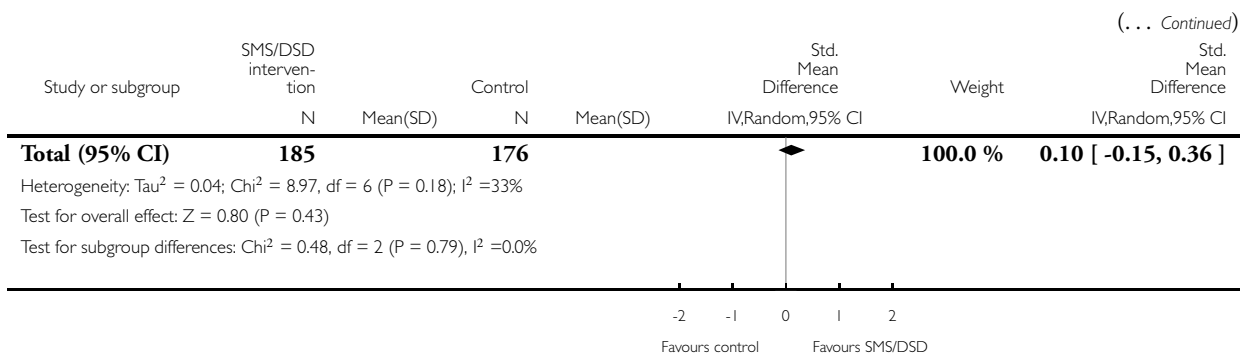
Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 15 Hearing aid benefit - short/medium-term - SMS content



(Continued ...)



(1) Medium term data

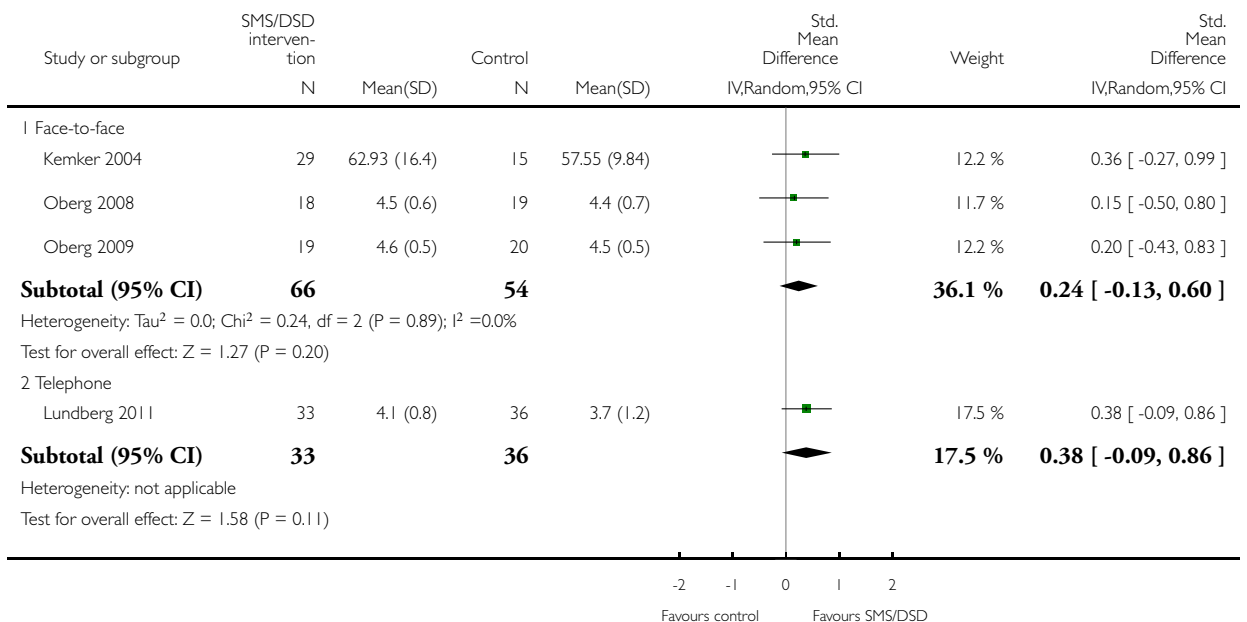
(2) Medium term data

Analysis 3.16. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 16 Hearing aid benefit - short/medium-term - DSD format.

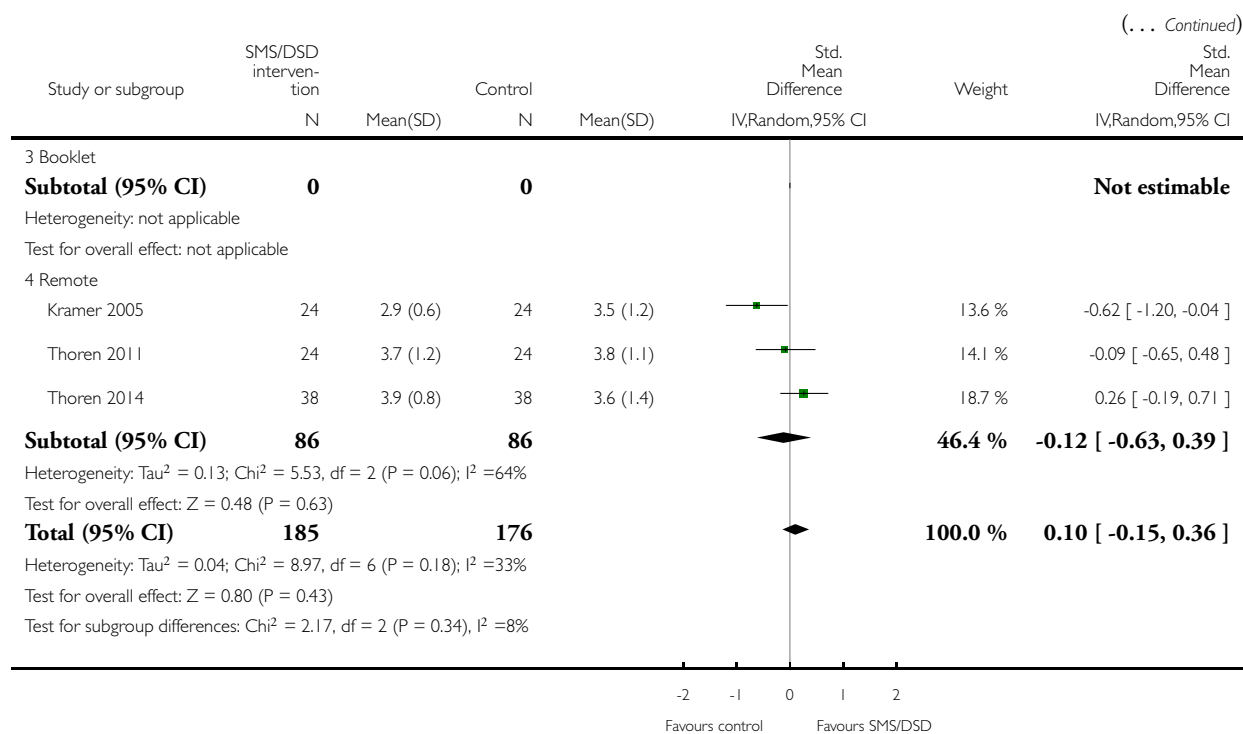
Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 16 Hearing aid benefit - short/medium-term - DSD format



(Continued ...)

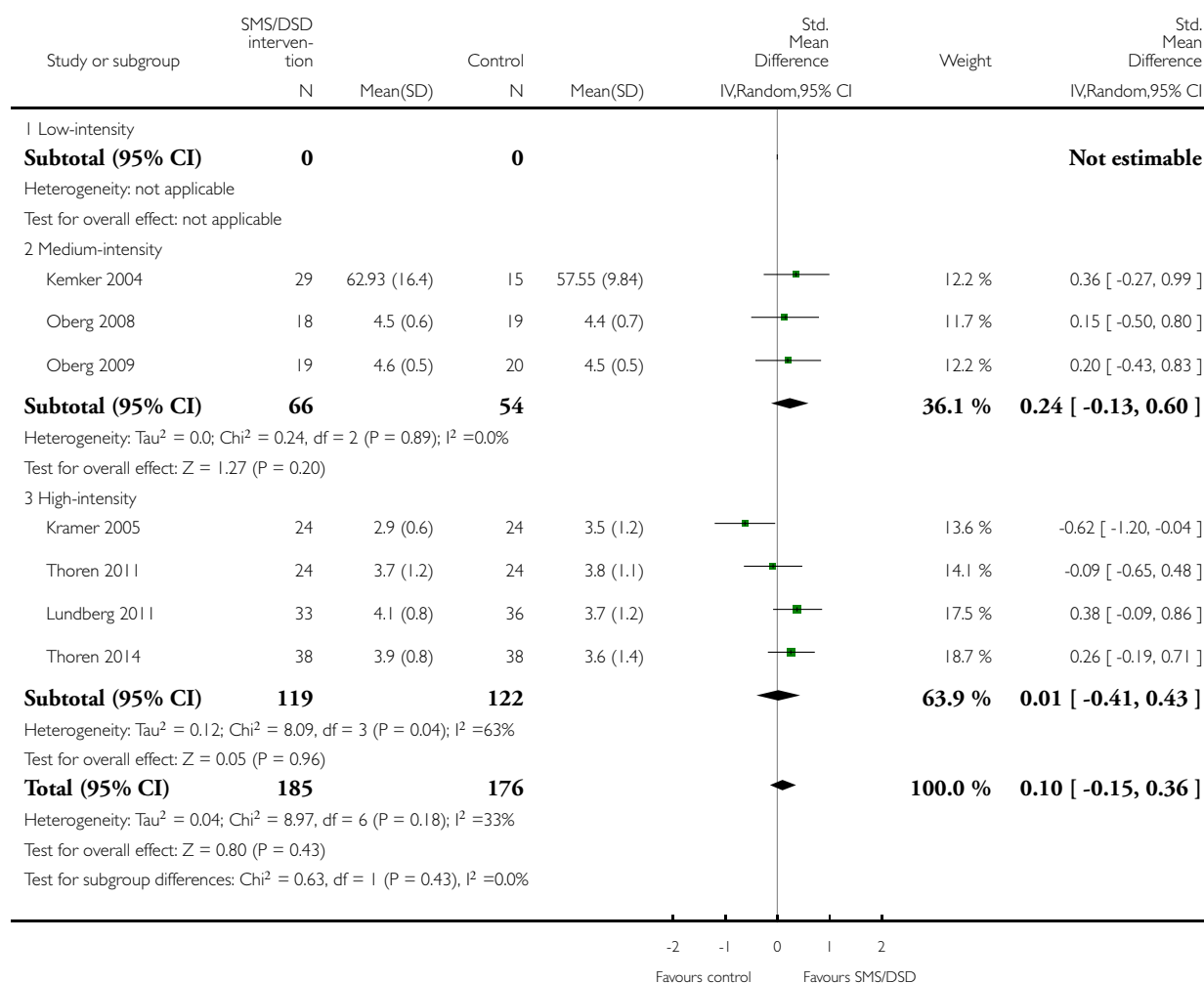


Analysis 3.17. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 17 Hearing aid benefit - short/medium-term - DSD intensity.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 17 Hearing aid benefit - short/medium-term - DSD intensity

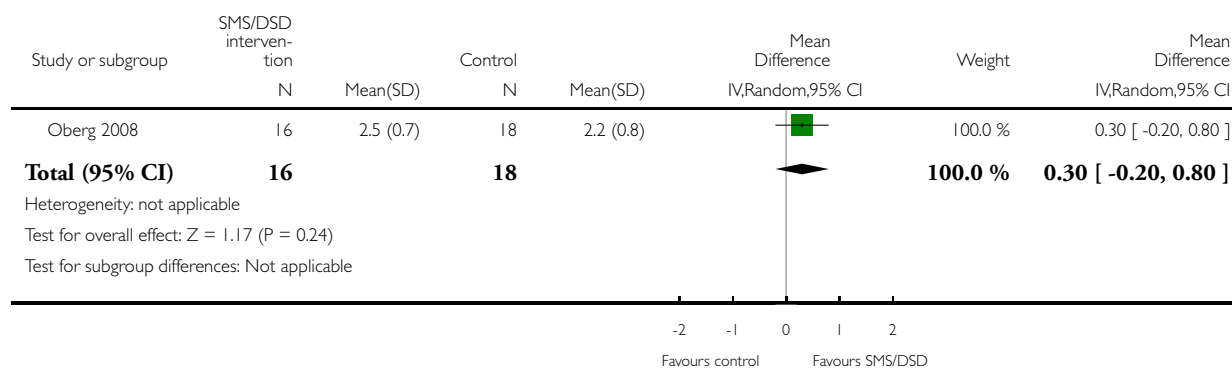


Analysis 3.18. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 18 Use of verbal communication strategy - long-term.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 18 Use of verbal communication strategy - long-term

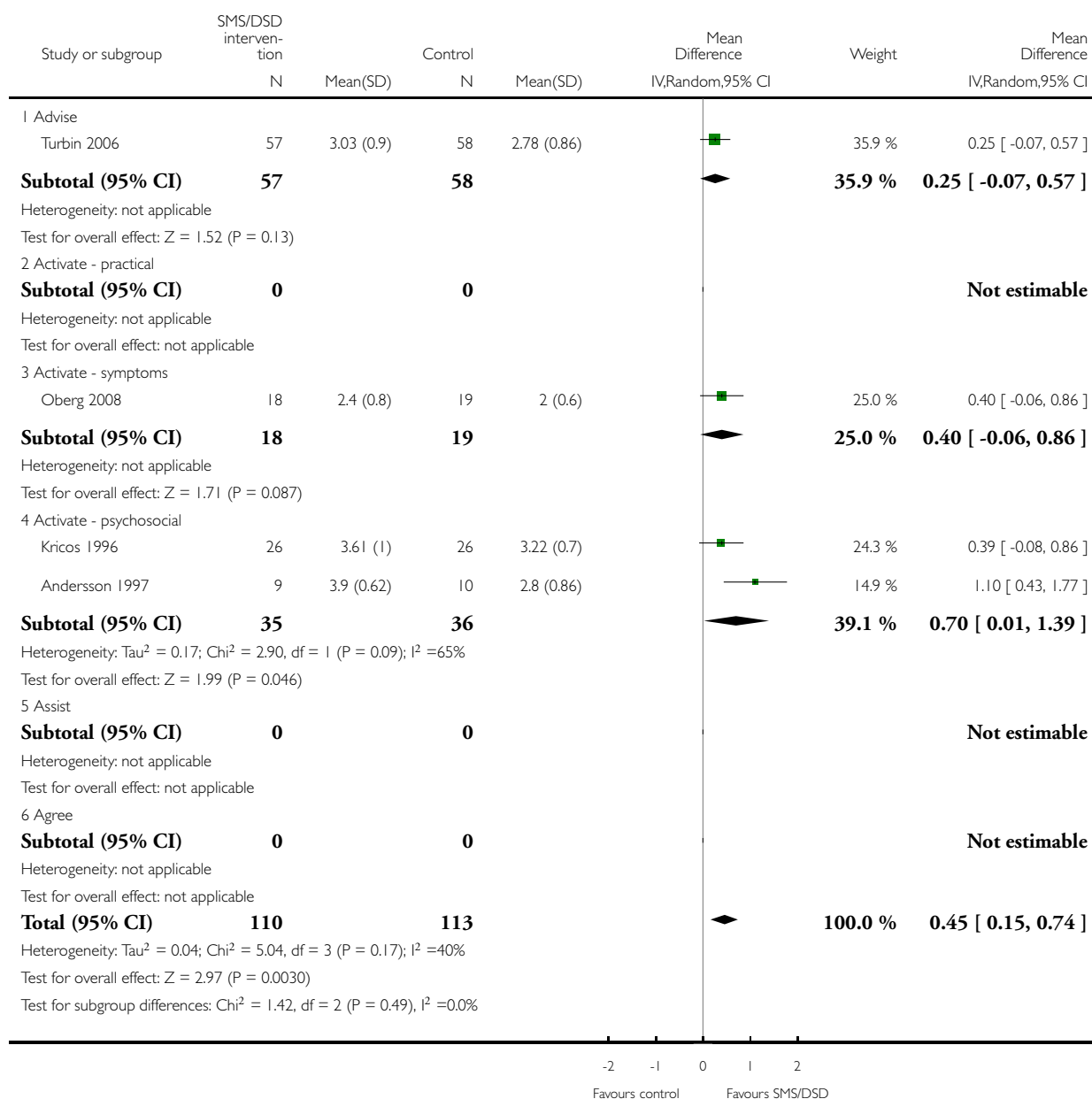


Analysis 3.19. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 19 Use of verbal communication strategy - short/medium-term - SMS content.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 19 Use of verbal communication strategy - short/medium-term - SMS content

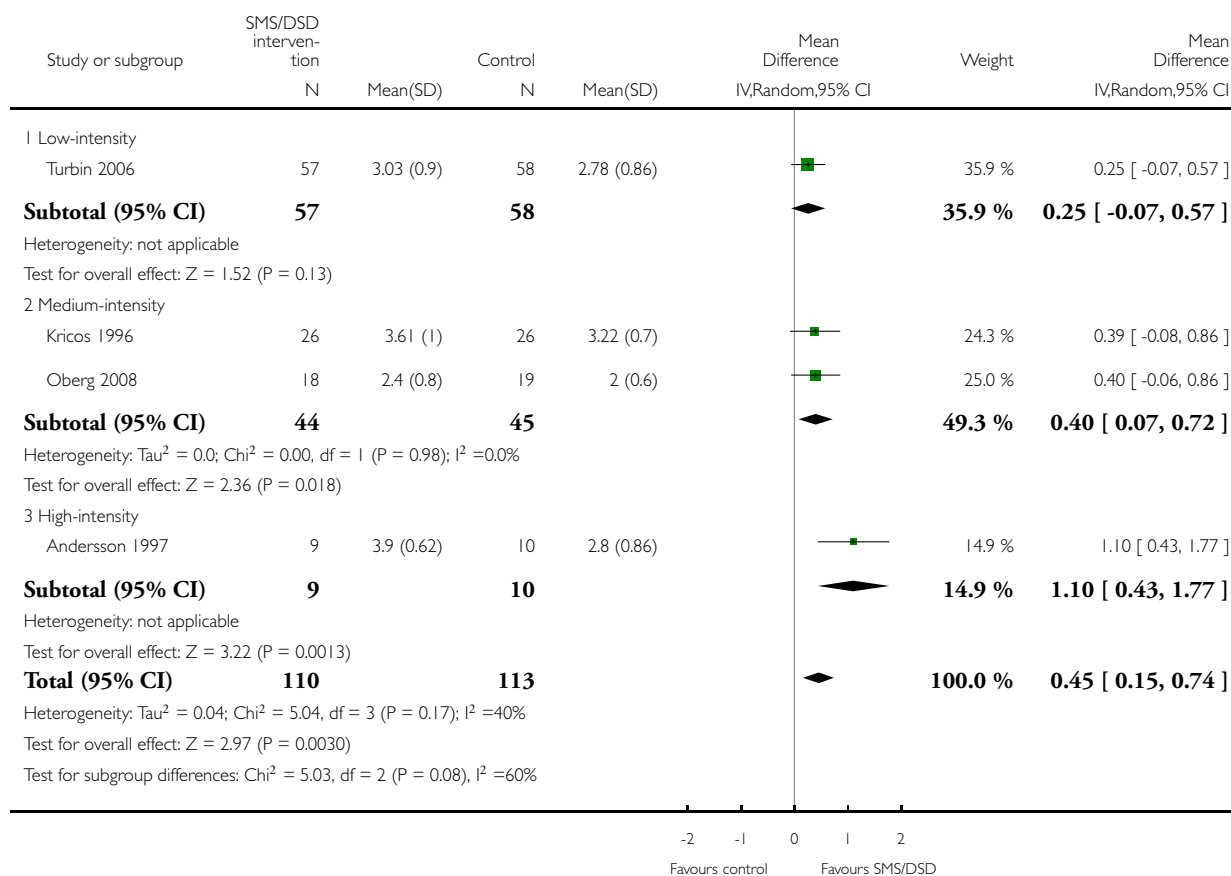


Analysis 3.20. Comparison 3 Combined SMS/DSD interventions versus control, Outcome 20 Use of verbal communication strategy - short/medium-term - DSD intensity.

Review: Interventions to improve hearing aid use in adult auditory rehabilitation

Comparison: 3 Combined SMS/DSD interventions versus control

Outcome: 20 Use of verbal communication strategy - short/medium-term - DSD intensity



ADDITIONAL TABLES

Table 1. Intervention range and type

CCM element	Study reference	Hearing health-care intervention	Control intervention	Self-management support (SMS) sub-type	Delivery system design (DSD) format	Delivery system design (DSD) intensity	Delivery system design (DSD) mode	Subgroup (s) compared
Health system	None found	-						
Community resources	None found							
Decision support	None found							
Clinical information system	None found							
Delivery system design	Campos 2013	Remote on-line fitting	Face-to-face fitting	Activate practical	- Remote (on-line) versus face-to-face	Low	Individual	DSD format
	Cherry 1994	Telephone follow-up at 6, 9 and 12 weeks post-fitting - questions answered, troubleshooting and counselling	Face-to-face follow-up on request	Activate symptom	- Telephone versus face-to-face	Medium versus low	Individual	DSD format and intensity
	Collins 2013	60-minute group orientation with PowerPoint presentation covering use, care and maintenance of the hearing aid	30-minute individual orientation with handout of same PowerPoint presentation	Advise	Face-to-face	Low	Group versus individual	DSD mode
	Cunningham 2001	As many post-fitting adjustments	No post-fitting adjustments	Activate symptom	- Face-to-face	Medium versus low	Individual	DSD intensity

Table 1. Intervention range and type (Continued)

		justments as patients requested						
	Lavie 2014	Simultaneous binaural fitting	Sequential binaural fitting	Activate - practical	Face-to-face but simultaneous versus sequential	Low	Individual	DSD format
	Ward 1981	Self-help book on hearing tactics	Single session face-to-face advice on hearing tactics	Advise	Booklet versus face-to-face	Low	Individual	DSD format
Self-management support	Fitzpatrick 2008	Auditory training - phoneme discrimination in single words, then sentences and then in presence of background noise. 13 x 1 hour	13 x 1-hour lectures on hearing loss, hearing aids and communication	Activate - symptom versus advise	Face-to-face	High	Individual	SMS content
	Kricos 1996	4-week communication training programme 8 x 1-hour including information and practice in communication skills and coping strategies for communication	8 x 1-hour analytic auditory training	Activate - psychosocial versus symptom	Face-to-face	High	Individual	SMS content
	Preminger 2010a	6 x 1-hour group communication	6 x 1-hour group communication	Activate - psychosocial+ versus	Face-to-face	High	Group	SMS content

Table 1. Intervention range and type (Continued)

		tion strategy training plus psychosocial exercises addressing emotional and psychological impact of hearing loss	tion strategy training	psychosocial				
	Saunders 2009	Pre-fitting counselling including demo	Pre-fitting counselling with no demo	Activate - symptom versus none	Face-to-face	Low	Individual	SMS content
	Saunders 2016	20 x 30-minute sessions auditory training (LACE) over a 4-week period on PC at home	20 x 30-minute sessions over a 4-week period listening to an audio book (placebo)	Activate - symptom versus none	Remote	High	Individual	SMS content
Combined SMS/DSD	Abrams 1992	Group AR 90 minutes once a week for 3 weeks post-fitting. Each week lectures covering different topics relating to hearing loss and communication	No intervention post-fitting	Advise	Face-to-face	Medium	Group	SMS content DSD format DSD intensity DSD mode
	Andersson 1994	60-minute individual behavioural counselling session then 3 consecutive weeks of group or in-	No intervention post-fitting	Activate - psychosocial	Face-to-face	Medium	Group or Individual	SMS content DSD format DSD intensity DSD mode

Table 1. Intervention range and type (Continued)

		dividual sessions where hearing tactics and coping strategies were taught and practised						
	Andersson 1995	60-minute individual behavioural counselling session then 4 x 2-hour sessions including video feedback on role play, applied relaxation, information and homework	No intervention	Activate - psychosocial	Face-to-face	High	Individual	SMS content DSD format DSD intensity
	Andersson 1997	Self-help manual supplied with 1-hour face-to-face training session including relaxation training followed by telephone contact over 4 consecutive weeks	No intervention	Activate - psychosocial	Face-to-face	High	Individual	SMS content DSD intensity
	Beynon 1997	4-week communication course - information and discussion regarding	No intervention	Advise	Face-to-face	Medium	Group versus individual	SMS content DSD intensity DSD mode

Table 1. Intervention range and type (Continued)

		hearing loss, hearing aids and communication						
	Chisolm 2004	4-week course AR - 2 hours per week with lectures covering different aspects relating to hearing loss and communication	No intervention	Advise	Face-to-face	Medium	Group versus Individual	SMS content DSD intensity DSD mode
	Eriksson-Mangold 1990	5 visits including fitting - structured guidance, use of diary with specific homework tasks, restricted HA use during first month	Standard fitting	Activate - psychosocial	Face-to-face	High	Individual	SMS content DSD intensity
	Ferguson 2016	Interactive DVD to use at home following fitting including information and exercises on hearing aid management and communication	Standard fitting	Activate - psychosocial	DVD	Medium	Individual	SMS content DSD format DSD intensity
	Gil 2010	8 x 1-hour twice a week for 4 weeks - synthetic -	No intervention	Activate - symptom	Face-to-face	High	Individual	SMS content DSD inten-

Table 1. Intervention range and type (Continued)

		pointing to words, figures, digits and verbal repetition						sity
	Kemker 2004	2 x 1-hour sessions of hearing aid orientation - could be pre- or post-fitting. In the review we combined these groups	No intervention	Advise	Face-to-face	Medium	Individual	SMS content DSD intensity
	Kramer 2005	5 sequential videos showing listening situations and coping tactics	No intervention	Advise	Remote (video)	High	Individual	SMS content DSD format DSD intensity
	Kricos 1992	4-week communication training programme 8 x 1-hour including information and practice in communication skills and coping strategies for communication	No intervention	Activate - psychosocial	Face-to-face	High	Individual	SMS content DSD intensity
	Kricos 1996	4-week communication training programme 8 x	No intervention	Activate - psychosocial	Face-to-face	High	Individual	SMS content DSD intensity

Table 1. Intervention range and type (Continued)

		1-hour including information and practice in communication skills and coping strategies for communication						
	Lundberg 2011	Weekly topic-based reading tasks based on an information booklet plus 5 x 10- to 15-minute telephone calls with an audiologist to discuss the tasks	Information booklet	Activate - psychosocial versus advise	Telephone	High	Individual	SMS content DSD format DSD intensity
	Miranda 2008	7 x 50-minute weekly session of auditory training - mix of synthetic and analytic	No intervention	Activate - symptom	Face-to-face	High	Individual	SMS content DSD intensity
	Oberg 2008	Pre-fitting sound awareness training. 3 visits with different listening exercises. 1 visit without amplification and 2 with an ex-	No intervention	Activate - symptom	Face-to-face	Medium	Individual	SMS content DSD intensity

Table 1. Intervention range and type (Continued)

		per- imental ad- justable aid							
	Oberg 2009	Pre-fitting use of an experimental adjustable hearing aid - 3 clinic visits to adjust the aid a week apart and experience at home in between	No intervention	Activate symptom	-	Face-to-face	Medium	Individual	SMS content DSD intensity
	Olson 2013	20 x 30-minute sessions at home over 4 weeks using interactive DVD delivering synthetic auditory tasks	No intervention	Activate symptom	-	Remote (DVD)	High	Individual	SMS content DSD format DSD intensity
	Preminger 2008	6 x 1-hour speech training classes including auditory and audiovisual analytic and synthetic tasks	No intervention	Activate symptom	-	Face-to-face	High	Group versus None	SMS content DSD intensity DSD mode
	Preminger 2010	Group AR plus separate group for SPs 4 x 90 minutes	Group AR without group for SPs	Advise		Face-to-face	Medium	Group	SMS content DSD intensity
	Saunders 2016	10 x 30-minute auditory training sessions	No intervention	Activate symptom	-	Re- mote (DVD or PC based)	High	Individual	SMS content DSD intensity

Table 1. Intervention range and type (Continued)

		delivered by DVD at home over a 2-week period OR 20 x 30-minute auditory training sessions delivered by PC at home over a 4-week period						
	Smaldino 1988	4 sessions of rehabilitation including information on hearing and hearing aids, practice and problem-solving regarding communication and role play	No intervention	Activate - psychosocial	Remote (PC-based)	Medium	Individual	SMS content DSD intensity
	Sweetow 2006	30 minutes 5 days a week for 4 weeks at home analytic and synthetic auditory training, information on communication strategies	No intervention	Activate - symptom	Remote (PC-based)	High	Individual	SMS content DSD format DSD intensity
	Thoren 2011	5-week online education programme including in-	Online discussion forum with 5 weekly topics but no	Advise versus Activate - psychosocial	Remote (email follow-up)	High	Individual	SMS content DSD format DSD intensity

Table 1. Intervention range and type (Continued)

		formation, tasks assignments and professional contact via email	task assignments and no professional guidance					
	Thoren 2014	5-week on-line rehabilitation programme including self-study, training and professional coaching in hearing physiology, hearing aids, and communication strategies as well as on-line contact with peers	No intervention	Activate - psychosocial	Remote	High	Individual	SMS content DSD format DSD intensity
	Turbin 2006	Single session of group AR - length not clear	No intervention	Advise	Face-to-face	Low	Group versus Individual	SMS content DSD intensity DSD mode
	Vreeken 2015	Weekly home visits for 3 to 5 weeks. Participants received a handbook with background information and a checklist accompanied with exercises covering:	No intervention	Activate - psychosocial	Face-to-face plus booklet	High	Individual	SMS content DSD format DSD intensity

Table 1. Intervention range and type (Continued)

		hearing aid use, maintenance and handling; living environment; hearing assistive devices; communication strategies						
	Ward 1978	2 treatment groups - 1 received 2 x 2-hour AR sessions, the other 4 x 2-hour sessions. Sessions including physical practice with aids and communication advice and practice. Also psychosocial aspects	No intervention	Activate - psychosocial	Face-to-face	Medium	Group	SMS content DSD intensity DSD mode
	Ward 1981	Self-help book on hearing tactics	No intervention	Advise	Booklet	Low	Individual	SMS content DSD format DSD intensity

AR: auditory rehabilitation

CCM: chronic care model

DSD: delivery system design

HA: hearing aid

SMS: self-management support

SP: spouse

APPENDICES

Appendix I. Search strategy

CENTRAL	PubMed	EMBASE (Ovid)	CINAHL (EBSCO)
<p>#1 MeSH descriptor: [Hearing Loss] explode all trees</p> <p>#2 MeSH descriptor: [Hearing Impaired Persons] explode all trees</p> <p>#3 hearing near (loss or impair*)</p> <p>#4 Hypoacusis or Hypoacus</p> <p>#5 #1 or #2 or #3 or #4</p> <p>#6 MeSH descriptor: [Adult] explode all trees</p> <p>#7 older or elderly or aged or aging or "middle age*" or "age related" or acquir* or adult*</p> <p>#8 #6 or #7</p> <p>#9 #5 and #8</p> <p>#10 MeSH descriptor: [Presbycusis] explode all trees</p> <p>#11 Presbycusis or Presbycuses</p> <p>#12 #9 or #10 or #11</p> <p>#13 MeSH descriptor: [Hearing Aids] this term only</p> <p>#14 MeSH descriptor: [Prosthesis Fitting] explode all trees</p> <p>#15 "hearing aid*"</p> <p>#16 "ear mold*" or earmold*</p> <p>#17 "ear mould*" or earmould*</p> <p>#18 amplif*</p> <p>#19 #13 or #14 or #15 or #16 or #17 or #18</p> <p>#20 #12 and #19</p> <p>#21 MeSH descriptor: [Health Behavior] this term only</p> <p>#22 MeSH descriptor: [Patient Compliance] this term only</p> <p>#23 MeSH descriptor: [Treatment Refusal] explode all trees</p> <p>#24 MeSH descriptor: [Patient Acceptance of Health Care] explode all trees</p> <p>#25 MeSH descriptor: [Counseling] this term only</p> <p>#26 MeSH descriptor: [Patient</p>	<p>#1 Search "Hearing Loss"[Mesh]</p> <p>#2 Search "Hearing Impaired Persons"[Mesh]</p> <p>#3 Search ("hearing loss" OR "hearing impair*")</p> <p>#4 Search (Hypoacusis or Hypoacus)</p> <p>#5 Search (#1 OR #2 OR #3 OR #4)</p> <p>#6 Search "Adult"[Mesh]</p> <p>#7 Search (older or elderly or aged or aging or "middle age*" or "age related" or acquir* or adult*)</p> <p>#8 Search (#6 OR #7)</p> <p>#9 Search (#5 AND #8)</p> <p>#10 Search "Presbycusis"[Mesh]</p> <p>#11 Search (Presbycusis or Presbycuses)</p> <p>#12 (#9 OR #10 OR #11)</p> <p>#13 Search "Hearing Aids"[Mesh:NoExp]</p> <p>#14 Search "Prosthesis Fitting"[Mesh]</p> <p>#15 Search "hearing aid*"</p> <p>#16 Search ("ear mold*" or earmold* or "ear mould*" or earmould* or amplif*)</p> <p>#17 (#13 OR #14 OR #15 OR #16)</p> <p>#18 (#12 AND #17)</p> <p>#19 Search "Health Behavior"[Mesh:NoExp]</p> <p>#20 Search "Patient Compliance"[Mesh:NoExp]</p> <p>#21 Search "Treatment Refusal"[Mesh]</p> <p>#22 Search "Patient Acceptance of Health Care"[Mesh]</p> <p>#23 Search "Counseling"[Mesh:NoExp]</p>	<p>1. exp hearing impairment/</p> <p>2. (hearing adj (loss or impair*)).tw.</p> <p>3. (Hypoacusis or Hypoacus)</p> <p>.tw.</p> <p>4. 1 or 2 or 3</p> <p>5. exp adult/</p> <p>6. (older or elderly or aged or aging or "middle age*" or "age related" or acquir* or adult*).tw</p> <p>7. 5 or 6</p> <p>8. 4 and 7</p> <p>9. exp presbycusis/</p> <p>10. (Presbycusis or Presbycuses)</p> <p>.tw.</p> <p>11. 8 or 9 or 10</p> <p>12. hearing aid/</p> <p>13. exp prosthesis/</p> <p>14. "hearing aid* ".tw.</p> <p>15. ("hearing aid*" or "ear mold*" or earmold* or "ear mould*" or earmould* or amplif*).tw</p> <p>16. 12 or 13 or 14 or 15</p> <p>17. 11 and 16</p> <p>18. patient compliance/</p> <p>19. health behavior/</p> <p>20. exp treatment refusal/</p> <p>21. exp patient attitude/</p> <p>22. counseling/</p> <p>23. exp patient education/</p> <p>24. behavior therapy/</p> <p>25. exp behavioral medicine/</p> <p>26. exp adaptive behavior/</p> <p>27. ((patient* or healthcare or "health care") and (compliance or cooperat* or co-operat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior)</p>	<p>S1 (MH "Deafness+") OR (MH "Hearing Loss, Partial+")</p> <p>S2 TX "hearing loss" or "hearing impair*"</p> <p>S3 TX Hypoacusis or Hypoacus</p> <p>S4 S1 OR S2 OR S3</p> <p>S5 (MH "Adult+")</p> <p>S6 TX older or elderly or aged or aging or "middle age*" or "age related" or acquir* or adult*</p> <p>S7 S5 OR S6</p> <p>S8 S4 AND S7</p> <p>S9 (MH "Presbycusis")</p> <p>S10 TX Presbycusis or Presbycuses</p> <p>S11 S8 OR S9 OR S10</p> <p>S12 (MH "Hearing Aids")</p> <p>S13 (MH "Prosthetic Fitting")</p> <p>S14 TX "hearing aid*" OR "ear mold*" OR earmold* OR "ear mould*" OR earmould* OR amplif*</p> <p>S15 S12 OR S13 OR S14</p> <p>S16 S11 AND S15</p> <p>S17 (MH "Health Behavior")</p> <p>S18 (MH "Patient Compliance")</p> <p>S19 (MH "Treatment Refusal")</p> <p>S20 (MH "Counseling")</p> <p>S21 (MH "Patient Education")</p> <p>S22 (MH "Audiology/MT")</p> <p>S23 (MH "Behavior Therapy")</p> <p>S24 (MH "Adaptation, Psychological")</p> <p>S25 (MH "Patient Attitudes")</p> <p>S26 TX (patient* or healthcare or "health care") and (compliance or cooperat* or co-op</p>

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Education as Topic] explode all trees #27 MeSH descriptor: [Audiology] explode all trees and with qualifiers: [Methods - MT] #28 MeSH descriptor: [Choice Behavior] this term only #29 MeSH descriptor: [Behavior Therapy] this term only #30 MeSH descriptor: [Behavioral Medicine] explode all trees #31 MeSH descriptor: [Adaptation, Psychological] explode all trees #32 (patient* or healthcare or "health care") and (compliance or cooperat* or co-operat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior) #33 Any MeSH descriptor with qualifier(s): [Psychology - PX, Rehabilitation - RH, Utilization - UT] #34 educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or prefitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or ((audio* or aural or auditory) near rehab*) or "hearing tactic*" or "active fitting" #35 ("take up" or "take-up" or use or utilis* or utiliz* or "non-use") and #19 #36 #21 or #23 or #22 or #24 or #25 or #26 or #27 or #28 or #29 or #30 or #31 or #32 or #33 or #34 or #35 #37 #36 and #20 #38 MeSH descriptor: [Hear-	#24 Search "Patient Education as Topic"[Mesh] #25 Search "Audiology/methods"[Mesh] #26 Search "Choice Behavior"[Mesh:NoExp] #27 Search "Behavior Therapy"[Mesh:NoExp] #28 Search "Behavioral Medicine"[Mesh] #29 Search "Adaptation, Psychological"[Mesh] #30 Search ((patient* or healthcare or "health care") and (compliance or cooperat* or co-operat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior) #31 Search (PX OR RH OR UT[MeSH Subheading]) #32 Search (educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or prefitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or "hearing tactic*" or "active fitting") #33 Search ("audio* rehab*" OR "aural rehab*" OR "auditory rehab*") #34 Search (("take up" or "take-up" or use or utilis* or utiliz* or "non-use") AND #26) #35 (#19 OR #20 OR #21 OR #22 OR #23 OR #24 OR #25 OR #26 OR #28 OR #29 OR #30 OR #31 OR #32 OR #33 OR #34) #36 (#35 AND #18)).tw 28. (educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or prefitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or ((audio* or aural or auditory) adj rehab*) or "hearing tactic*" or "active fitting").tw 29. ("take up" or "take-up" or "use" or utilis* or utiliz* or "non-use").tw 30. 16 and 29 31. 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 30 32. 17 and 31	erat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior) S27 TX educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or prefitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or "audio* rehab*" or "aural rehab*" or "auditory rehab*" or "hearing tactic*" or "active fitting" S28 TX ("take up" or "take-up" or use or utilis* or utiliz* or "non-use") S29 S15 AND S28 S30 S17 OR S18 OR S19 OR S20 OR S21 OR S22 OR S23 OR S24 OR S25 OR S26 OR S27 OR S29 S31 S16 AND S30
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ing Aids] explode all trees and with qualifiers: [Utilization - UT, Therapy - TH, Psychology - PX] #39 #37 or #38	#37 Search (("Hearing Aids/psychology"[Mesh] OR "Hearing Aids/utilization"[Mesh])) #38 (#36 OR #37)		
CAB Abstracts (Ovid)	AMED (Ovid)	Web of Science (Web of Knowledge)	Trial Registries
1. exp hearing impairment/ 2. (hearing adj (loss or impair*)).tw. 3. (Hypoacusis or Hypoacusis) .tw. 4. (Presbycusis or Presbycusis) .tw. 5. exp people with hearing impairment/ 6. 1 or 2 or 3 or 4 or 5 7. ("hearing aid*" or "ear mold*" or earmold* or "ear mould*" or earmould* or amplif*).tw 8. 6 and 7 9. exp patient compliance/ 10. exp counselling/ 11. exp patient education/ 12. health behaviour.sh. 13. ((patient* or healthcare or "health care") and (compliance or cooperat* or co-operat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior)).tw 14. (educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or prefitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or ((audio* or aural or auditory) adj	1. exp Deafness/ 2. (hearing adj (loss or impair*)).tw. 3. (Hypoacusis or Hypoacusis) .tw. 4. (Presbycusis or Presbycusis) .tw. 5. 1 or 2 or 3 or 4 6. exp Hearing aids/ 7. exp Prosthesis/ 8. ("hearing aid*" or "ear mold*" or earmold* or "ear mould*" or earmould* or amplif*).tw 9. 6 or 7 or 8 10. 5 and 9 11. exp Patient compliance/ 12. exp Health behavior/ 13. exp Treatment refusal/ 14. counseling/ 15. exp Patient education/ 16. behavior therapy/ 17. exp Adaptation psychological/ 18. ((patient* or healthcare or "health care") and (compliance or cooperat* or co-operat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior)).tw 19. (educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or pre-	#1 TS=(hearing NEAR/6 (loss or impair*)) #2 TS=(Hypoacusis or Hypoacusis) #3 TS=(Presbycusis or Presbycusis) #4 #3 OR #2 OR #1 #5 TS=("hearing aid*" OR "ear mold*" OR earmold* OR "ear mould*" OR earmould* OR amplif*) #6 #5 AND #4 #7 TS=((patient* or healthcare or "health care") and (compliance or cooperat* or co-operat* or adherence or "non-compliance" or noncompliance or "non-adherence" or nonadherence or accept* or nonaccept* or behaviour or behavior)) #8 TS=(educat* or train* or counsel* or "self manag*" or "management plan*" or "care plan*" or "support tool*" or "chronic care mode" or ccm or promot* or psycholog* or psychosocial or teach* or motivat* or prefitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or ((audio* or aural or auditory) NEAR/6 rehab*) or "hearing tactic*" or "active fitting") #9 TS=("take up" or "take-up" or use or utilis* or utiliz* or "non-use") #10 #5 AND #9	Clinicaltrials.gov "hearing aid" OR "hearing aids" ICTRP hearing aid*

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rehab*) or "hearing tactic*" or "active fitting").tw 15. ("take up" or "take-up" or "use" or utilis* or utiliz* or "non-use").tw 16. 7 and 15 17. 9 or 10 or	fitting or Postfitting or "fitting protocol" or ghabp or "hearing aid orientat*" or HAO or "pre-fitting" or "post-fitting" or ((audio* or aural or auditory) adj rehab*) or "hearing tactic*" or "active fitting").tw 20. ("take up" or "take-up" or "use" or utilis* or utiliz* or "non-use").tw 21. 9 and 20 22. 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18 or 19 or 21 23. 10 and 22	#11 #10 OR #8 OR #7 #12 #11 AND #6	
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WHAT'S NEW

Last assessed as up-to-date: 13 June 2016.

Date	Event	Description
11 July 2016	New citation required but conclusions have not changed	We included five new studies (Ferguson 2016 ; Lavie 2014 ; Saunders 2016 ; Thoren 2014 ; Vreeken 2015), and excluded a further four studies (Aazh 2016 ; Cardemil 2014 ; Kuk 2014 ; Lavie 2013). We identified three new ongoing studies (ISRCTN77340339 ; NCT02233361 ; NCT02264314), and two studies are awaiting classification (Henshaw 2013 ; Malmberg 2015). One new combined self-management support/delivery system design study showed a significant effect on adherence to hearing aid use in the short term. Otherwise the results and conclusions of the review are unchanged
11 July 2016	New search has been performed	The review has been updated to include data from an updated search (June 2016)

CONTRIBUTIONS OF AUTHORS

FB and SdeL conceived the review question. FB wrote the protocol, review and co-ordinated comments from the other authors.

EM and LE provided clinical advice, content relating to audiology and participated in study selection and data extraction.

SdeL and SJ provided general feedback on the protocol, review text and analyses.

DECLARATIONS OF INTEREST

Fiona Barker: none known

Emma Mackenzie: none known

Lynette Elliott: none known

Simon Jones: none known

Simon de Lusignan: none known

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Internal sources

- University of Surrey, PhD programme, UK.

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External sources

- National Institute for Health Research, UK.

Infrastructure funding for Cochrane ENT

- National Institute for Health Research, UK.

NIHR-Cochrane Incentive Award 2015 (for the update of the review)

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We made some revisions to the method regarding definitions of intervention types to provide more theoretical support for our approach using the chronic care model. In particular, we provided additional evidence and rationale for subgroup analyses within the elements of self-management support and delivery system design.

Other changes to the methods included the following:

We decided to use a random-effects model for all analyses and included our rationale in the full review text.

We had originally intended to enter skewed endpoint data from studies of fewer than 200 participants into additional tables rather than into the analyses. In fact there was a considerable quantity of potentially skewed data with high standard deviations relative to the means. We considered that the risk of not including these data and being able to draw only limited conclusions outweighed the statistical risk of including these data. Where there was a high risk of skew we noted this and downgraded the quality of the evidence if necessary. This allowed us to combine data that would otherwise have been excluded from the meta-analyses.

We revised the outcomes of interest. Originally we had intended to scope the range of outcomes addressed in adult hearing healthcare research. However, we were advised that this introduced additional complexity into an already complex review. We therefore redefined (post-protocol publication but pre-analysis) the primary and secondary outcomes we judged to be of most interest to patients, clinicians and policy-makers. Adherence remained our main primary outcome but we combined self-reported daily hours of hearing aid use and data-logged hearing aid use into a single outcome of daily hours of hearing aid use. Adverse effects were included in the original review

protocol as a secondary outcome. During the peer review process we were advised to include this as a primary outcome. We then specified the secondary outcomes of interest.

Some situations anticipated in the protocol did not occur and so some types of analysis could not be performed (such as funnel plots).

INDEX TERMS

Medical Subject Headings (MeSH)

Communication; Hearing Aids [*utilization]; Hearing Loss [*rehabilitation]; Patient Compliance [statistics & numerical data]; Quality of Life; Randomized Controlled Trials as Topic; Self Care [methods]; Time Factors

MeSH check words

Adult; Aged; Humans; Middle Aged