

Interventions for the reduction of prescribed opioid use in chronic non-cancer pain (Review)

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[Intervention Review]

Interventions for the reduction of prescribed opioid use in chronic non-cancer pain

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ABSTRACT

Background

This is the first update of the original Cochrane Review published in 2013. The conclusions of this review have not changed from the 2013 publication. People with chronic non-cancer pain who are prescribed and are taking opioids can have a history of long-term, high-dose opioid use without effective pain relief. In those without good pain relief, reduction of prescribed opioid dose may be the desired and shared goal of both patient and clinician. Simple, unsupervised reduction of opioid use is clinically challenging, and very difficult to achieve and maintain.

Objectives

To investigate the effectiveness of different methods designed to achieve reduction or cessation of prescribed opioid use for the management of chronic non-cancer pain in adults compared to controls.

Search methods

For this update we searched CENTRAL, MEDLINE, and Embase in January 2017, as well as bibliographies and citation searches of included studies. We also searched one trial registry for ongoing trials.

Selection criteria

Included studies had to be randomised controlled trials comparing opioid users receiving an intervention with a control group receiving treatment as usual, active control, or placebo. The aim of the study had to include a treatment goal of dose reduction or cessation of opioid medication.

Data collection and analysis

Two review authors independently extracted data and assessed risk of bias. We sought data relating to prescribed opioid use, adverse events of opioid reduction, pain, and psychological and physical function. We planned to assess the certainty of the evidence using the GRADE approach, however, due to the heterogeneity of studies, we were unable to combine outcomes in a meta-analysis and therefore we did not assess the evidence with GRADE.

Main results

Three studies are new to this update, resulting in five included studies in total (278 participants). Participants were primarily women (mean age 49.63 years, SD = 11.74) with different chronic pain conditions. We judged the studies too heterogeneous to pool data in a meta-analysis, so we have summarised the results from each study qualitatively. The studies included acupuncture, mindfulness, and cognitive behavioral therapy interventions aimed at reducing opioid consumption, misuse of opioids, or maintenance of chronic pain management treatments. We found mixed results from the studies. Three of the five studies reported opioid consumption at post-treatment and follow-up. Two studies that delivered 'Mindfulness-Oriented Recovery Enhancement' or 'Therapeutic Interactive Voice Response' found a significant difference between groups at post-treatment and follow-up in opioid consumption. The remaining study found reduction in opioid consumption in both treatment and control groups, and between-group differences were not significant. Three studies reported adverse events related to the study and two studies did not have study-related adverse events. We also found mixed findings for pain intensity and physical functioning. The interventions did not show between-group differences for psychological functioning across all studies. Overall, the risk of bias was mixed across studies. All studies included sample sizes of fewer than 100 and so we judged all studies as high risk of bias for that category.

Authors' conclusions

There is no evidence for the efficacy or safety of methods for reducing prescribed opioid use in chronic pain. There is a small number of randomised controlled trials investigating opioid reduction, which means our conclusions are limited regarding the benefit of psychological, pharmacological, or other types of interventions for people with chronic pain trying to reduce their opioid consumption. The findings to date are mixed: there were reductions in opioid consumption after intervention, and often in control groups too.

PLAIN LANGUAGE SUMMARY

Reducing prescribed opioid use in adults with chronic non-cancer pain

Bottom line

Based on the available evidence, we do not know the best method of reducing opioids in adults with chronic pain conditions. We found mixed results from a small number of studies included in this review.

Background

This is an updated review. The first review was published in 2013. About one in five adults suffer from moderate or severe chronic pain that is not caused by cancer. Some people with this type of pain are treated with opioids (typically with drugs such as morphine, codeine, oxycodone, fentanyl, or buprenorphine, either as tablets or as patches placed on the skin). It is not unusual for this medication to be ineffective or to stop working over time, and, sometimes, effective pain relief is not achieved despite doses being increased. Stopping using opioid drugs is not easy, especially when they have been used for some time, because stopping abruptly can cause unpleasant side effects. This review looked for high-quality studies (randomised controlled trials) of treatments to help adults safely stop taking opioids prescribed for their pain.

Study characteristics

We searched for studies up to January 2017. We found five studies, and they investigated 278 people. Most people included in the studies were women, who were around 50 years of age, and reported a mixture of chronic pain conditions (e.g. headache, back pain, muscle pain). The studies included acupuncture, mindfulness, and cognitive behavioral therapy as strategies to decrease the amount of opioids taken by adults with chronic pain.

Key results

No conclusions can be drawn from this small amount of information. Therefore, it is not clear whether these treatments decrease the amount of opioids in adults with chronic pain (primary outcome) or reduce pain intensity, physical ability or mood (secondary outcomes). Three studies did include negative effects of their treatment, and two reported that the participants did not have anything negative happen to them because of the trial they were in. Non-randomised studies, not included in this review, do indicate that for many people intensive rehabilitation packages may bring about major reduction in opioid use. Reducing prescribed opioid use in chronic non-cancer pain is an important topic in need of more systematic research.

Quality of the evidence

We were not able to judge the quality of evidence included in this review because the studies were so different and could not be combined.

BACKGROUND

Chronic pain of moderate or severe intensity and lasting six months or longer affects around 20% of adults and imposes significant reduction in quality of life (Moore 2013). Opioids have long been used in the treatment of acute and cancer pain, and over the last two decades there has been a marked increase in their prescription for chronic non-cancer pain (CNCP), especially in the USA, Australia, and Europe. Estimates of the numbers of people with CNCP treated with opioids are not commonly available, but one estimate for the UK indicates that almost one million people may use some form of opioid (Gallagher 2009).

Several randomised controlled trials suggested that opioids provide modest pain relief in the short to medium term (typical trial duration is 12 weeks; Furlan 2006; Kalso 2004). However, there is much less evidence that opioids provide long-term pain relief in CNCP (ASIPP 2012; Noble 2010), especially when statistical imputation methods are used in which withdrawal for any reason is regarded as treatment failure (and relevant to clinical practice), in comparison to the common practice of carrying forward the last observed pain readings to the end of the trial and using that measurement to estimate efficacy (Moore 2012; Steiner 2011).

Adverse events, principally sedation, impaired cognitive function, depression, constipation, and bladder dysfunction, are also common during opioid therapy (Benyamin 2008), with up to 80% of users suffering at least one adverse event (Moore 2005). Long-term opioid use can be associated with immune system depression, hormonal disturbances, and hyperalgesia (Benyamin 2008), as well as fractures (Miller 2011), and increased all-cause mortality in older people compared with other analgesics (Solomon 2010). Opioid use also carries risks of tolerance, dependence, and abuse.

Practicalities of the real world like prescribing restrictions for nonopioid analgesics, or guidelines that suggest early use of opioids, can mean that many people are prescribed opioid drugs, sometimes inappropriately. The American Society of Interventional Pain Physicians suggests that the majority of patients who start chronic opioid treatment continue with the treatment throughout their life (ASIPP 2012). On occasion, opioid doses are increased as a result of insufficient analgesia or the development of tolerance (i.e. requiring a higher dose to obtain the same therapeutic benefit). This can lead to people being prescribed very high doses of opioids, but still without acceptable pain relief. American opioid sales quadrupled between 1999 and 2010 (ASIPP 2012). In the USA, increased prescribing is associated with higher rates of overdose and overdose death (Paulozzi 2011), but with an indication that 60% of CNCP opioid deaths occurred while opioids were used as directed (ASIPP 2012). However, there is some evidence to suggest that whilst the prescription of opioids remains high, it decreased between 2012 and 2015 in the USA (Guy 2017).

A number of professional societies worldwide have produced guidance advocating/promoting the judicious and careful use of opioids. The American Academy of Pain Medicine, the American Pain Society, and the American Society of Addiction Medicine jointly advise that healthcare providers should exercise caution when prescribing opioids, assessing circumstances and suitability on an individual basis (Chou 2009). Guidance in Washington State, USA (Washington State Agency Medical Group 2010) has passed into law (HB2876 2010). Current UK guidance is typical in that it recognises that prescription opioids can lead to problem use, and that there is considerable uncertainty in the literature about any long-term benefits of continued use.

There is growing concern that the widespread use of opioids has public health implications (Stannard 2012). The balance between benefit and risks generated during long-term therapy with opioids suggests that it may be neither clinically effective nor in patients' best interests to continue opioid prescription without adequate pain relief. There is, therefore, a potential need to facilitate and maintain opioid dose reduction. For many patients it is likely that long-term opioid treatment is continued even when benefit is not demonstrated, and greater patient benefits may accrue from opioid withdrawal. Patients who do not benefit from treatment in terms of pain, or who suffer unacceptable adverse events, should be helped to cease opioid treatment whilst concurrently addressing their pain (Ballantyne 2003).

There is a growing recognition that many patients will reach a state where the reduction of prescribed opioids is the desired and shared goal of both patient and clinician. This state is sometimes reached after a history of long-term, high-dose opioid use, making simple unsupervised cessation clinically challenging, if not impossible. This may occur, at least in part, because of the reluctance of patient and prescriber to reduce the opioid dose for fear of worsening pain, as well as issues of dependence and subsequent withdrawal symptoms.

There are many studies of methods of withdrawal from opioids; most, however, are undertaken in the context of addiction ser-

vices for people with an opioid abuse problem. Our interest here was in the planned reduction or total withdrawal of opioids prescribed for pain management. Common opioid reduction techniques in the addiction field are instructive and include opioid replacement stabilisation and dose tapering, and may involve psychological treatments (Amato 2011). Inducing withdrawal under sedation using opioid antagonists such as naloxone, naltrexone, or nalmefene is possible (Gowing 2009; Gowing 2010) but is not recommended owing to unacceptable risks of adverse events. It is unclear whether similar interventions are effective when adjusted to CNCP in which treatment aims differ, or if other approaches are more appropriate.

Description of the condition

Patients

• with chronic pain of a non-cancer-related origin (such as neuropathic, musculoskeletal, visceral, or head pain)

• who are prescribed opioid medication for pain management

• who have a treatment goal of dose reduction or cessation of opioid medicine

Description of the intervention

The interventions included any clinical method that aimed to facilitate opioid withdrawal or dose reduction as a compulsory or optional aspect of treatment, as either a primary or a secondary outcome. The intervention could be pharmacological, physiological, psychological, or another, as long as its methods are documented clearly within the study.

How the intervention might work

Different methods will have different mechanisms. In particular, we expect non-pharmacological treatment aimed at opioid reduction to operate principally through behaviour change, and pharmacological methods to operate principally by reducing or managing the adverse events of opioid use or opioid withdrawal.

Why it is important to do this review

Increased prescribing of opioids is a problem because of their potential to cause harm, along with issues of limited relief and tolerance. Given the known risks of opioid therapy, it is appropriate to continue to prescribe opioid medicines only to those people for whom the treatment produces acceptable benefits, weighed against any adverse events. Given evidence in many societies of huge increases in the use of medicinal opioids for CNCP, their limited effectiveness, and their adverse event profile, we can reasonably expect a large increase in people seeking clinical help to reduce or halt opioid consumption. An evidence summary of the most effective methods is needed, along with guidance on treatment development.

OBJECTIVES

To investigate the effectiveness of different methods designed to achieve reduction or cessation of prescribed opioid use for the management of chronic non-cancer pain in adults compared to controls.

METHODS

Criteria for considering studies for this review

Types of studies

Included studies had to be randomised controlled trials (RCTs) comparing opioid users receiving an intervention with a control group receiving treatment as usual, active control, or placebo. The aim of the study had to include a treatment goal of dose reduction or cessation of opioid medicine. We excluded studies that included fewer than 10 participants in each arm at post-treatment.

Types of participants

Participants were adults (18 years of age or older) using prescription opioids for management of CNCP with a duration of at least three months. Pain conditions could include but were not limited to: neuropathic pain, myofacial pain, back pain, fibromyalgia, headache, abdominal, neck or musculoskeletal pain.

We excluded studies involving only participants with issues of addiction, abuse, dependence, or non-prescribed opioid use, and involving participants using opioids for pain relief during palliative care. This is because the aims of treatment for these populations differ substantially from those for the population of interest.

Types of interventions

We planned to include in this review a large variety of intervention types. Interventions could be based in pharmacology, physiology, psychology, spirituality, or another approach, provided that the underpinning methodology was well documented in the study and was valid. Eligible intervention types could include opioid antagonist treatment, dose tapering, or opioid replacement with other pain-relieving medication. Interventions could also involve physical therapy, massage, disability management, complementary therapies, or psychological approaches such as cognitive behavioural therapy, counselling, and coping techniques.

We excluded studies encompassing only interventions specifically for opioid addiction, medication overuse, dependence, or withdrawal symptoms.

Types of outcome measures

We extracted relevant outcomes before treatment, immediately after treatment, and at follow-up, at least three months later but no longer than a year. If there were two follow-up time points, the later would be chosen. Where appropriate, we extracted means and standard deviations of outcomes assessed with psychometrically tested measures.

Primary outcomes

The primary outcomes of this review are:

- prescribed opioid use in adults;
- adverse events related to opioid reduction.

Secondary outcomes

Secondary outcomes are:

- pain intensity/severity;
- psychological functioning;
- physical functioning.

Search methods for identification of studies

Electronic searches

We have conducted two searches to date. For the original review we searched the Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE, and Embase from 1999 to April 2013. For this update we searched the same databases on 4 January 2017 to identify any further studies meeting the inclusion criteria:

• CENTRAL (via CRSO) - April 2013 to 4/1/17;

• MEDLINE (via OVID) - April 2013 to December week 1 2016;

• Embase (via OVID) - April 2013 to 2017 week 1.

We did not restrict the searches based on language. See Appendix 1 for search strategies.

Searching other resources

We searched the reference lists of retrieved papers and carried out a citation search to identify any potentially eligible papers not found through the electronic search. We also contacted the authors of studies identified for inclusion to obtain additional data relevant to this review and not included in the published articles. Finally, we searched Clinicaltrials.gov for additional trials that met eligibility.

Data collection and analysis

Selection of studies

We filtered search results initially by title and abstract, and obtained full copies of potentially eligible studies. Two review authors read the studies to confirm eligibility, with disagreements discussed and mediated by a third review author if necessary. In the first version of this review, we limited the selection of studies to those published from 2000 onwards, to reflect the major growth in opioid prescribing for CNCP after 2000.

Data extraction and management

Two review authors extracted data using a standard data extraction form to include details of participants, intervention method and duration, quantity and type of opioid used, study design, and treatment outcomes. We discussed any discrepancies with a third review author.

Assessment of risk of bias in included studies

We assessed risk of bias in the included studies using the Cochrane 'Risk of bias' tool (Higgins 2011) to assign judgements of high, low, or unclear risk of bias to sequence generation, allocation concealment, blinding, incomplete outcome data, selective reporting, and any other potential sources of bias in the included studies. Specifically for other sources of biases, we assessed the sample size as a risk of bias and coded any studies that included fewer than 100 participants as high risk of bias. Two authors (CE, EF) independently assessed risk of bias for each study and resolved disagreement by discussion. 'Risk of bias' assessments are included in the Characteristics of included studies.

We assessed the following for each study.

• Random sequence generation (checking for possible selection bias). We assessed the method used to generate the allocation sequence as: low risk of bias (any truly random process, e.g. random number table; computer random number generator); unclear risk of bias (method used to generate sequence not clearly stated); and high risk of bias (studies that included a biased randomisation procedure). We excluded studies that were not randomised.

• Allocation concealment (checking for possible selection bias). The method used to conceal allocation to interventions prior to assignment determines whether intervention allocation could have been foreseen in advance of, or during recruitment, or changed after assignment. We assessed the methods as: low risk of bias (e.g. telephone or central randomisation; consecutively numbered, sealed, opaque envelopes); unclear risk of bias (method not clearly stated).

• Blinding of outcome assessment (checking for possible detection bias). We assessed the methods used to blind outcome

assessors from knowledge of which intervention a participant received. We assessed the methods as: low risk of bias (study had a clear statement that outcome assessors were unaware of treatment allocation, and ideally described how this was achieved); unclear risk of bias (study stated that outcome assessors were blind to treatment allocation but lacked a clear statement on how it was achieved).

• Incomplete outcome data (checking for possible attrition bias due to the amount, nature and handling of incomplete outcome data). We assessed the methods used to deal with incomplete data as: low risk (attrition fully reported and no differences between completers and non-completers); unclear risk of bias (attrition unclear or unclear differences between completers and non-completers); high risk of bias (attrition not reported or differences between completers and non-completers).

• Selective reporting (checking for reporting bias). We assessed whether all outcomes were reported in the paper as low risk of bias. High risk of bias would be given when data were not reported and not provided on request.

• Size of study (checking for possible biases confounded by small size). We assessed studies as being at low risk of bias (200 or more participants per treatment arm); unclear risk of bias (50 to 199 participants per treatment arm); high risk of bias (fewer than 50 participants per treatment arm).

Measures of treatment effect

We planned to use risk ratio (RR) to establish statistical difference, and number needed to treat for an additional beneficial outcome (NNTB) and number needed to treat for an additional harmful outcome (NNTH) as absolute measures of benefit or harm. We defined a 'responder' to treatment as a participant who experienced at least a 50% reduction in opioid consumption, or achieved complete opioid withdrawal or a reduction of their intake to below 'high' dose, which we identified as 120 mg a day oral morphine equivalent. Trials have previously shown that dose-related harms of taking more than 120 mg a day of opioid drugs outweigh the benefits (Braden 2010; Morasco 2010; Sullivan 2010), and published guidelines, including those of the American Pain Society and the American Academy of Pain Medicine (Chou 2009), and by the Washington State Agency Medical Group 2010, recommend a cut-off at 120 mg a day. A responder also had to have, at worst, no increase in pain as a result of the intervention. Both aspects of improvement had to be maintained for at least three months post intervention.

Our planned analyses included:

- Treatment versus control opioid reduction (post-treatment);
- Treatment versus control opioid reduction (follow-up);
- Treatment versus control pain intensity (post-treatment);
- Treatment versus control pain intensity (follow-up);

• Treatment versus control psychological functioning (post-treatment);

• Treatment versus control psychological functioning (follow-up);

• Treatment versus control physical functioning (post-treatment);

• Treatment versus control physical functioning (follow-up).

We planned to describe adverse events reported in all studies.

Unit of analysis issues

The unit of analysis was the individual participant.

Dealing with missing data

We used the intention-to-treat approach to deal with missing data. We would include in the analysis all participants who were randomised to treatment, and we assumed that those for whom follow-up data were not available were non-responders.

Assessment of heterogeneity

We anticipated that there would be significant clinical heterogeneity between studies (participants, conditions, interventions), so we planned to pool data using a random-effects model.

Assessment of reporting biases

We planned to assess publication bias by estimating the number of unpublished null studies needed to make a clinical finding likely to be unstable or irrelevant (Moore 2008). Unfortunately, this was not possible because we were not able to calculate any effect sizes.

Data synthesis

We planned to combine data using Review Manager 5 (RevMan 5) (RevMan 2014). We planned to include dichotomous outcome analysis for the number of people who achieved a 50% reduction in opioid consumption. For this analysis, we planned to use Mantel-Haenszel statistics, and report risk ratio outcomes. We described any adverse events in the trials. For our secondary outcomes, we planned to analyse pain intensity, psychological functioning, and physical functioning in separate analyses using inverse variance methods in a random-effects model. We planned to summarise heterogeneity using I² statistics (Higgins 2003), which we planned to interpret following Deeks 2011 reference points of 0% to 40%: might not be important; 30% to 60%: may represent moderate heterogeneity; 50% to 90%: may represent substantial heterogeneity; 75% to 100%: considerable heterogeneity.

If data were homogeneous we planned to combine data in a metaanalysis, and if heterogeneous to describe the findings from the studies separately, focusing on our primary and secondary outcomes.

Quality of the evidence

We planned that two review authors (CE, EF) would independently rate the quality of the outcomes. We planned to use the GRADE approach to rank the quality of the evidence using RevMan 5 (RevMan 2014), and the guidelines provided in chapter 12 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Schünemann 2011).

The GRADE approach uses five considerations (study limitations, consistency of effect, imprecision, indirectness and publication bias) to assess the quality of the body of evidence for each outcome. The GRADE system uses the following criteria for assigning grade of evidence.

• High: we are very confident that the true effect lies close to that of the estimate of the effect;

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

• Low: our confidence in the effect estimate is limited; the true effect may be substantially different from the estimate of the effect;

• Very low: we have very little confidence in the effect estimate; the true effect is likely to be substantially different from the estimate of effect.

We planned to decrease our rating by one (-1) or two (-2) if we identified:

- serious (-1) or very serious (-2) limitation to study quality;
- important inconsistency (-1);
- some (-1) or major (-2) uncertainty about directness;
- imprecise or sparse data (-1);
- high probability of reporting bias (-1).

'Summary of findings' table

We planned to include a 'Summary of findings' table to present the main findings for comparison of interventions to reduce opioid consumption versus control in a transparent and simple tabular format. In particular, we planned to include key information concerning the quality of evidence, the magnitude of effect of the interventions examined, and the sum of available data on the outcomes of reduction of opioid consumption, adverse events, pain intensity, psychological functioning, and physical functioning. We plan to include a 'Summary of findings' table in the next update, providing sufficient evidence is available.

Subgroup analysis and investigation of heterogeneity

We planned subgroup analyses to assess the effect of entry dose on intervention efficacy, and to compare outcomes between pain conditions or intervention type if sufficient data were available.

Sensitivity analysis

We planned to perform a sensitivity analysis should we suspect that studies with high risk of bias were significantly skewing results of a comparison, removing studies from the analysis to assess their influence.

RESULTS

Description of studies

Results of the search

This is an updated search of a review previously published in 2013 (Windmill 2013). During the first search of databases from 1999 to April 2013, we included three papers that reported on two studies (Naylor 2010; Zheng 2008). We searched CENTRAL, MEDLINE and Embase for studies eligible for inclusion. In the second search from April 2013 to January 2017 we identified 3480 abstracts; 2878 abstracts after duplicates were removed (Figure 1). We identified two new studies that met eligibility for this update (Garland 2014; Sullivan 2017). We contacted all first authors of the included studies for additional studies that we had not captured in our search, and identified one further study that met eligibility (Jamison 2010). Therefore, three studies are new to this update (Garland 2014; Jamison 2010; Sullivan 2017), resulting in five studies overall.



Included studies

Five studies met the inclusion criteria (Garland 2014; Jamison 2010; Naylor 2010; Sullivan 2017; Zheng 2008) that included 278 participants with chronic pain (women = 184; men = 94; mean age = 49.63 years, SD = 11.74). Three studies included participants with mixed chronic pain conditions, Naylor 2010 exclusively included participants with chronic musculoskeletal pain, and participants in Jamison 2010 reported chronic neck or back pain. Jamison 2010; Naylor 2010, Sullivan 2017, and Zheng 2008 reported that the average pain duration was 12.68 years (SD = 10.72).

Of the five studies, two studies aimed to reduce opioid consumption via cognitive behavioural therapy (CBT) (Sullivan 2017) or electroacupuncture (Zheng 2008). Two studies aimed to reduce opioid misuse in people with chronic pain (Garland 2014; Jamison 2010). These studies delivered CBT or mindfulness treatments to the participants. One study aimed to increase treatment compliance and adherence in people with chronic pain receiving a painmanagement treatment (Naylor 2010). Here, the authors delivered CBT to everyone, and then provided a therapeutic interactive voice response to participants randomised to the treatment condition to improve maintenance of treatment. Two studies compared treatment to active controls and the remaining studies used standard care controls. Full details can be found in the Characteristics of included studies table.

In addition to published data, two author groups (Naylor 2010; Zheng 2008) provided additional data for the outcomes of pain and psychological functioning (Appendix 2). The remaining studies included data pertinent to this review meaning that we did not need to contact them for additional data.

Excluded studies

We excluded seven studies from this review, none new to this update. Three did not meet methodological standards (Crisostomo 2008; Krymchantowski 2003; Townsend 2008), while four did not have opioid reduction as a primary aim (Hale 2007; Potter 2010; Roland 2011; Weinstein 2006). Full details are available in the Characteristics of excluded studies table.

Risk of bias in included studies

We assessed risk of bias using the Cochrane 'Risk of bias' tool (Higgins 2011). Justification for judgements can be found in Characteristics of included studies and the figures of bias assessments in Figure 2 and Figure 3.





Figure 3.



Allocation

Random sequence generation

Three studies described sound randomisation procedures and we judged them to be low risk of bias, one study did not give a clear description, and we judged one study to have a high risk of bias as it randomised people based on the order that they were entered into the trial.

Allocation concealment

Three studies provided adequate descriptions of allocation concealment and we gave a low risk of bias rating, whilst two studies did not describe methods to conceal allocation and we marked them as unclear risk.

Blinding

Detection bias

We rated one study as low risk of bias for blinding outcome assessors, and the remaining four studies as unclear risk.

Performance bias

We excluded performance bias from this review as it is not possible to blind personnel who are delivering psychological treatments, or participants who are receiving them.

Incomplete outcome data

For attrition bias, one study did not have any dropouts and was marked as low risk of bias. The four further studies were judged to be unclear because they did not note differences between completers and non-completers.

Selective reporting

Most studies reported data for all outcomes and were marked as low risk of bias. However, one study had missing data and was rated as high risk of bias.

Other potential sources of bias

We rated all studies as high risk of bias for including small sample sizes (fewer than 50 participants per treatment arm).

Effects of interventions

All studies provided data to be analysed. However, due to the heterogeneity of the studies, we did not pool data in a meta-analysis. Therefore, we did not conduct GRADE analyses. We describe findings from each study.

Primary Outcomes

Opioid use

Garland 2014 compared a Mindfulness-Oriented Recovery Enhancement (MORE) treatment with a support group control. They did not assess objective opioid use before or after the trial. However, desire for opioids and opioid misuse was reported. Authors reported that the MORE group had a significantly lower desire for opioid consumption post treatment, but results were not maintained at follow-up. The MORE group also self-reported significantly lower opioid misuse (63%) compared to the support group (32%) at post-treatment. However, similar to the desire for opioids, participants did not differ at follow-up.

Jamison 2010 did not assess objective opioid use before or after the trial. A self-reported, prescription drug-use questionnaire was taken, but differences between groups at post-treatment were not reported.

Naylor 2010 compared Therapeutic Interactive Voice Response (TIVR) through a computer for four months with usual treatment, following CBT delivered to all participants for 11 weeks. The TIVR group (n = 26, 14 of whom were using opioids at baseline) reported a significant decrease in opioid use from baseline at both four- and eight-month follow-ups, with three participants stopping opioid use entirely. The standard care control (n = 25, 15 of whom were using opioids at baseline) significantly increased opioid consumption from baseline to the eight-month follow-up, and three more participants began opioid treatment. At post-treatment and eight-month follow-up, the difference in mean opioid dose was significant, with the TIVR group using less than the standard care control.

Sullivan 2017 randomised 35 participants to an opioid-tapering support intervention (n = 18) or usual care (n = 17). Both the treatment and control group reduced their intake of opioids at 22 weeks, and there was no significant difference between groups (morphine-equivalent doses (MED) taper support = 111.94, SD = 153.63; usual care = 169.85, SD = 201.31). Similarly at follow-up, no between-group differences were identified (MED) taper support = 99.51, SD = 151.99; usual care = 138.24, SD = 155.85). There was also no significant difference in the percent reduction

from baseline and 22 weeks or 34 weeks for either group. However, opioid consumption was reduced compared to baseline levels.

Zheng 2008 randomised participants to receive either real electroacupuncture (REA; n = 17) or sham electroacupuncture (SEA; n = 18) for 20 minutes twice a week for six weeks. Opioid consumption varied considerably within each group, and the mean consumption at baseline differed between groups, being 462 (± 463) mg a week in the REA group and 296 (± 288) mg a week in the SEA group. Participants in both groups who completed the six weeks of treatment (REA = 12; SEA = 14) reported a significant reduction in opioid consumption between baseline and the end of treatment at eight weeks, of 64% and 46% in the REA and SEA groups, respectively. In an intention-to-treat analysis, the reductions were 39% and 26%. The difference between groups was not statistically significant. At follow-up at 20 weeks (REA = 9; SEA = 14), opioid consumption had gradually increased in the REA group and was significantly higher at 20 weeks than at eight weeks, while in the SEA group there was no significant change.

Adverse events

Garland 2014 did not report the occurrence of adverse events. When we contacted the study authors, they reported that there had been no adverse events.

Jamison 2010 reported adverse events across all participants of dry mouth (44.9%), constipation (38.4%) sweating (37.5%), memory lapse (28.4%), weakness (24.1%), itching (23.9%), and headaches (28.4%). The treatment group reported lower rates of constipation and itching, but higher vision problems. The control group reported more severe constipation, sneezing, and nightmares than the treatment group.

Naylor 2010 did not report on adverse events, but contact with the study authors confirmed that there were no adverse events associated with treatment.

Sullivan 2017 reported one severe study-related adverse event in the taper-support group. The study psychiatrist prescribed nortriptyline during the participant's initial psychiatric evaluation, which the participant had a severe reaction to. This medication was discontinued and symptoms resolved.

Zheng 2008 reported a total of 33 adverse events during the treatment period with REA, and 19 with SEA, none of which were classed as serious adverse events. Opioid-based adverse events decreased from baseline to eight weeks after treatment by 40% in the REA group and 45% in the SEA group.

Secondary Outcomes

Pain intensity

These numbers are from data supplied by the study authors (Appendix 2) and differ slightly from the published data.

Garland 2014 used the Brief Pain Inventory, pain intensity subscale and found that the MORE group reported significantly lower pain intensity at post-treatment (Mean (M) = 4.86, SD = 1.38) that met the threshold for minimally clinically significant change, in comparison to the support group (SG) control group (M = 5.71, SD = 1.58). This between-group difference was maintained at follow-up (MORE M = 4.77, SD = 1.95; SG M = 6.10, SD = 1.48).

Jamison 2010 assessed pain using the Brief Pain Inventory, pain intensity subscale. However, post-treatment means and standard deviations were not reported.

Naylor 2010 analysed pain using the McGill Pain Questionnaire, reported in Naylor 2008 (Naylor 2010). The TIVR group reported a decrease in typical pain from baseline to eight months from 5.7/10 to 3.4/10, and the standard care control from 6.8 to 5.7. The difference between groups was statistically significant.

Sullivan 2017 assessed pain intensity using the Brief Pain Inventory, pain severity subscale. There was no significant difference between groups at 22 weeks (taper support = 4.72, SD = 1.62; usual care = 5.77, SD = 1.92) or at 34 weeks (taper support = 4.67, SD = 1.79; usual care = 6.16, SD = 2.64).

Zheng 2008 used the Visual Analogue Scale to assess pain intensity. Average pain at baseline was 4.9/10 in the experimental group and 5.6/10 in the control group, and post-treatment scores were 4.2 and 5.4, respectively. No differences were detected between groups. At 20 weeks average pain scores were 3.6 and 4.6.

Psychological function

Garland 2014 assessed depression using the Calgary Symptoms of Stress Inventory, depression subscale. There were no betweengroup differences at post-treatment (MORE = 8.20, SD = 7.09; SG = 10.76, SD = 6.44). Depression was not assessed at followup.

Jamison 2010 reported that the control group had significantly higher scores compared to the treatment group on anxiety and depression measures (treatment = 6.06, SD = 3.55; standard care = 9.00, SD = 3.39; treatment = 8.1, SD = 4.8; standard care = 9.06, SD = 4.11) respectively.

Naylor 2010 reported a decrease in scores of depression throughout the study, using the Beck Depression Inventory. There were significant between-group differences, favouring lower scores in the experimental group compared to the control group at posttreatment (TIVR = 10.4, SD = 6.4; standard care = 16.7, SD = 11.2) and eight-month follow-up (TIVR = 8.1, SD = 4.8; standard care = 14.9, SD = 8.7).

Sullivan 2017 assessed depression using the Patient Health Questionnaire-9. There were no significant group differences at 22 weeks (taper support = 8.88, SD = 7.49; usual care = 11.27, SD = 6.58), or at 34 weeks (taper support = 9.00, SD = 5.80; usual care

= 11.13, SD = 7.53).

Zheng 2008 reported a significant decrease in depression scores (Beck Depression Inventory) from baseline to post-treatment at eight weeks in the REA group, from 18 to 17. Scores in the SEA group also decreased, from 19 to 15. At final measurement at 20 weeks, the REA group mean score was 14, and the SEA group mean score was 15. There was no significant difference between the groups at either time point.

Physical function

Garland 2014 assessed functional interference with the Brief Pain Inventory. The MORE group reported a significantly lower painrelated functional interference compared to the SG at post-treatment (MORE M = 5.22 SD = 1.88; SG M = 6.90, SD = 1.50), and the between-groups difference was maintained at follow-up (MORE M = 4.60 SD = 2.66; SG M = 6.75, SD = 1.86).

Jamison 2010 reported that there were no differences between treatment and control at six-month post-treatment on the Pain Disability Index.

Naylor 2010 reported physical function using the SF-36 Physical Function composite scale. The experimental group showed a small increase (from 31/100 to 40/100) in functioning over eight months, while the control group did not (29/100 to 31/100). The difference between groups was statistically significant.

Sullivan 2017 assessed functional interference with the Brief Pain Inventory, interference subscale. Significant group differences were identified at 22 weeks (taper support = 4.55, SD = 2.39; usual care = 6.38, SD = 2.11). There were also group differences at 34 weeks, but these were only trending in favour of the taper support group (taper support = 4.49, SD = 2.08; usual care = 6.05, SD = 2.72).

Zheng 2008 did not measure physical function.

DISCUSSION

Summary of main results

We included three new studies in this update. However, there remain no adequate data from which to draw any conclusions from five small studies with different interventions and only 278 treated participants. Therefore, our conclusions for this review have not changed.

Overall completeness and applicability of evidence

Because of the very small number of included studies, for the previous version of the review we decided to additionally investigate methods of prescription opioid reduction that were not randomised controlled trials, in case this was a more commonly used study design. In 2013, we looked at papers from the previous search results and additional reference searching. Inclusion criteria remained the same as in the main search, excepting the criteria of randomised controlled design. The overall completeness and applicability from the previous update are still relevant for this update, due to the lack of evidence in this area and the complexity of designing and conducting an RCT to investigate this problem. In contrast to the randomised evidence, there was a much larger body of evidence from observational studies. A three-week, outpatient, intensive, multidisciplinary pain rehabilitation programme conducted at the Mayo Clinic Pain Rehabilitation Center demonstrated large reductions in medication use, particularly in use of opioids, in a number of publications in recent years. The threeweek programme included stretching, goal setting, stress management, physical therapy, pain management, relaxation, and occupational therapy (Mayo 2017). Typical opioid use in patients at admission was high, often above 40% and as high as 100%, and at discharge and follow-up was low, often below 10%. The analyses were retrospective or longitudinal, and not randomised, but represented an interesting body of additional data. Results like these were obtained for 159 patients with fibromyalgia (Hooten 2007), for 383 patients after fusion or non-fusion spinal surgery, or no surgery (Crisostomo 2008), in a group of 411 patients with a wide range of age and non-cancer pain conditions (Darchuk 2010), and for 634 chronic pain patients of different smoking status (Hooten 2009). In a group of 213 patients all taking opioids on admission, the rate of opioid use at discharge was 7% and remained low for as long as six months after admission (Townsend 2008).

Change in medication use, including opioid medication use, is a common feature of multimodal and multi-component programmes of cognitive behavioural therapy for chronic pain. The evidence for such programmes in improving disability status and reducing the impact of mental health outcomes is promising (Williams 2012). At present, however, it is not possible to extract and describe the components of such programmes for their effectiveness on medication consumption outcomes, although individual trials report positive effects. A challenge will be to determine methods of analysis, if possible, of such treatment packages with multiple components addressing multiple outcomes.

Others have sought evidence from literature reviews to prevent opioid over-use, and have put forward what is claimed to be an evidence-based algorithmic approach (Atluri 2012).

Legislation (HB2876 2010) has had a major effect on opioid prescribing in Washington State, where a de-facto limit of 120 mg oral morphine equivalent a day is suggested, with higher doses available after consultation with a specialist. An interim assessment showed that about half of physicians followed guidance on opioid prescribing, and that about 90% of them found it useful (DLI

2009). A survey has shown large falls in opioid prescribing (27%), and in the proportion taking more than 120 mg a day oral morphine equivalents (35%), as well as in opioid-related deaths (50%) (Franklin 2012). The benefits of treating patients with chronic pain continue to show a lack of long-term benefit. A systematic review assessing opioid consumption comparing to placebo or no opioids did not find studies assessing long-term follow-up (i.e. one year). Opioids were found to be frequently abused and misused in chronic pain samples, and associated with dependence, fractures, and myocardiac infarctions (Chou 2015). Further, recent recommendations from the Centres for Disease Control state that opioid therapy should not be given to people with chronic pain (not associated with a life-limiting condition) due to potential harm and lack of medium- and long-term benefit (Dowell 2016).

Quality of the evidence

The evidence base identified by this review is small and limited and we were unable to perform a GRADE assessment of the certainty of evidence in this area. The individual studies have small numbers of participants, and overall we have evidence of the experience of only 278 chronic pain patients. There is a heterogeneity of interventions and outcome reporting. Poor reporting is common, meaning that the risk of bias was often unclear or high.

Potential biases in the review process

We were not aware of any biases in the review process, although there was a potential for bias in searching for studies. While the intention to reduce opioid use may have been clear, possible interventions may have been disparate.

Agreements and disagreements with other studies or reviews

We found no other similar reviews.

AUTHORS' CONCLUSIONS

Implications for practice

There were too few data in this review to permit any comments about implications for practice.

Implications for research

General implications

There is an urgent need for more research. There is a growing population of people with chronic pain, who also have chronic use of opioids, which are thought to be untherapeutic and for whom reduction is a primary clinical goal. We are unable to reduce our uncertainty around any treatment offered to these people for this purpose.

Design

We need more randomised controlled trials of theoretically grounded behaviour-change interventions that focus on opioid medication use in the context of medically treated chronic pain. There should be no fewer than 100 participants in each trial arm.

Measurement (endpoints)

More work is needed to agree the best endpoints for treatments of medication reduction. Measures of the type of opioid and the median daily opioid dose in morphine equivalents consumed in a particular time period are critical to report. In addition, measures of patient-relevant outcomes such as mood, social functioning, and personal role functioning are also important to assess.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Garland 2014

Methods	Randomised, 8-week, active-control RCT Assessments at pre-post treatment and 3-month follow-up
Participants	Chronic pain condition: mixed (including fibromyalgia, arthritis, cervicalgia, lumbago, and other) 115 participants were randomised, MORE (n = 57), support group (n = 58) Female 78; male 37 Mean age = 48 years, SD = 13 years
Interventions	Mindfulness-oriented recovery enhancement (MORE): delivered to participants in 8 two-h sessions by a Master's level clinical social worker with > 10 years of clinical experience. 15-min homework sessions were given to be practiced daily Support group: 8 two-h sessions for participants to discuss topics pertinent to chronic pain and long-term opioid use were discussed. Topics were designed to match sessions in the MORE group. Lead by a Master's level clinical social worker
Outcomes	Prescribed medication use: no measure of prescription. Measures of desire for opioids and Current Opioid Misuse Measure were included Pain: BPI, pain severity subscale Psychological function: CSQ; Calgary Symptoms of Stress Inventory Physical function: BPI, pain interference subscale
Notes	Funded by National Institute on Drug Abuse and a grant from Fahs-Beck Fund for Research and Experimentalation

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Participants were randomly allocated to MORE or to the SG. Order of randomisa- tion was computer generate via simple ran- domisation blocks of varying sizes (6-8) to preserve unpredictability of allocation."
Allocation concealment (selection bias)	Low risk	"Order of randomisation was computer generated via simple randomisation blocks of varying sizes (6-8) to preserve unpre- dictability of allocation."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Assessments were conducted by project staff blind to each respondent's group as- signment, which was concealed through-

Garland 2014 (Continued)

		out the study"
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition is described but analyses of dif- ferences between completers and non-com- pleters was not conducted
Selective reporting (reporting bias)	Low risk	Comprehensive reporting of outcomes
Size	High risk	42/46 received at least 1 session
Jamison 2010		
Methods	Randomised trial, standard medical care co Assessments at baseline and 6 months	ontrol
Participants	Chronic pain patients with a history or risk of opioid misuse. All participants had chronic back or neck pain 42 patients were randomised; CBT (n = 21), standard medical care (n = 21) Female 20; male 22 Mean age = 46.79 years, SD = 7.27 years	
Interventions	CBT for prevention of opioid misuse: group and individual sessions were delivered to participants. Treatment was composed of 5 components; electronic monthly diaries, monthly urine screens, monthly completion of the Opioid Compliance Checklist, monthly group education sessions and worksheets, individual motivational compliance counselling. Groups and individual sessions included enhancing and maintaining motivation to avoid illicit substance use, coping with urges, problem solving, and lifestyle balance. Participants were provided with discussions round risk factors, motivational counselling, homework, and completed a monthly opioid compliance checklist. Treatment lasted 6 months Standard medical care: no description provided	
Outcomes	Prescribed medication use: no measure of prescription. Measures of desire for opioids and Current Opioid Misuse Measure were included Pain: BPI, pain severity subscale Psychological function: HADS Physical function: BPI, pain interference subscale, Pain Disability Index	
Notes	The study was supported in part by an investigator-initiated grant from End Pharmaceu- ticals, Chadds Ford, PA and Grants R21 DA024298, Jamison PI and K23DA020682 Wasan PI from the National Institute on Drug Abuse of the National Institutes of Health, Bethesda, MD, and the ARthritis Foundindation (Wasan)	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Jamison 2010 (Continued)

Random sequence generation (selection bias)	High risk	"assignment to treatment group based on a randomized number list created before the start of the study. Subjects were assigned to their group in the order that they entered into the study"
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No description provided
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition is described but analyses of dif- ferences between completers and non-com- pleters was not conducted
Selective reporting (reporting bias)	High risk	Post-treatment means and SDs are not pro- vided for all measures
Size	High risk	21 per group
Naylor 2010		
Methods	11 weeks plus 120 days' duration, random ments at baseline, post intervention, 4 mor	ised, standard care, controlled trial. Assess- ths, and 8 months post intervention
Participants	Chronic pain condition: chronic musculosl 55 participants randomised, 51 participants Female 44, male 7 Mean age 46 (SD ± 11.5) years	xeletal pain s received allocated intervention
Interventions	All participants received 11 90-min session: Therapeutic Interactive Voice Response (n Standard care (n = 25)	s of CBT pre-randomisation = 26)
Outcomes	 Prescribed medication use: dose and frequency of opioid analgesics, non-steroidal anti- inflammatory drugs, benzodiazepines, and antidepressants Pain: Short Form MPQ, Pain Symptoms sub scale from the TOPS Psychological function: BDI, SF-36 Mental Function Scale, CSQ Physical function: SF-36 Physical Function Scale, TOPS Total Pain Experience Scale 	
Notes	4 participants were excluded following rand This research was supported by grants fror (NIDA) R21 DA016115, National Institut AMS) R01 AR052131, and National Instit AAA) R01 AA014270	domisation. n the National Institute of Drug Addiction e of Arthritis, Musculoskeletal Diseases (NI- rute on Alcohol Abuse and Alcoholism (NI-

Naylor 2010 (Continued)

Risk of bias

5		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomized using a stratified block de- sign"
Allocation concealment (selection bias)	Low risk	"consecutively numbered, sealed envelopes were prepared for each gender group by the statistician"
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not described
Incomplete outcome data (attrition bias) All outcomes	Low risk	No dropouts during the study
Selective reporting (reporting bias)	Low risk	Comprehensive reporting of outcomes
Size	High risk	25/26 per group

Sullivan 2017

Methods	Non-blinded RCT. 18-week intervention. Assessments at baseline, 22 weeks, and 34 weeks
Participants	Chronic pain condition: chronic non-cancer pain 35 participants randomised, 18 to taper support; 17 to usual care Female 25, male 10 Mean age 54.4 years (SD = 10.1)
Interventions	All participants were shown a 14-min video of patients who had successfully tapered off opioids before randomisation Opioid tapering group: participants underwent motivational interviewing and then 17 30-min weekly sessions with Physician Assistant delivering CBT and occasional motivational interviewing. Booster phone calls were scheduled at 24, 39, and 32 weeks. Sessions were conducted in person and over the phone. Participants received workbooks and CDs throughout treatment. Participants' opioid medication was tapered 10% for the first 3 weeks, then reassessed and reduced 10% for the following weeks. Participants could pause tapering, but were not allowed to remain in the study if they increased opioid prescription Usual care: participants in the control group received usual care from physicians, with no restrictions on medications during the study period
Outcomes	Prescribed medication use: mean daily opioid dose Pain:BPI, pain severity subscale Psychological function: Patient Health Questionnaire-9 (depression)

Sullivan 2017 (Continued)

	Physical function: BPI, pain interference subscale
Notes	This work was supported by Grant R34DA033384 from the National Institute on Drug Abuse to Mark Sullivan Dr. Sullivan reports consulting with Chrono Therapeutics. Other authors report no conflicts of interest

Risk	of	bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Study participants were randomized 1:1 to receive either the opioid taper interven- tion or usual care according to a computer- generated randomisation list in sealed en- velopes"
Allocation concealment (selection bias)	Low risk	"Study participants were randomized 1:1 to receive either the opioid taper interven- tion or usual care according to a computer- generated randomisation list in sealed en- velopes"
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not described
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition is described but analyses of dif- ferences between completers and non-com- pleters was not conducted
Selective reporting (reporting bias)	Low risk	All outcomes reported
Size	High risk	17/18 per treatment arm

Zheng 2008

Methods	20 weeks, randomised, single-blind, sham-controlled trial Assessments at baseline and at weeks 5, 8, 12, 16, and 20
Participants	35 participants with non-malignant pain for > 3 months, using opioid medication Female 17, male 18 Mean age 50 years (SD = 11 years)
Interventions	Electroacupuncture (n = 17) for 20 min twice/week for 6 weeks Sham electroacupuncture (n = 18) for 20 min twice/week for 6 weeks

Zheng 2008 (Continued)

Outcomes	Prescribed opioid use: dosage of opioid-like medications and adverse events (type and frequency) Pain: pain intensity- visual analogue scale; MPQ Psychological function: BDI Physical function: none SF-36 v2 Health Survey
Notes	The study was supported by a research grant provided by the then Faculty of Life Sci- ences, RMIT University. Ms. Jessica Guo, a research candidate, was supported by an Aus- tralian Postgraduate Award (APA) and an Australian Acupuncture and Chinese Medicine Association (AACMA) Research Grant 2005

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Block randomisation code was computer generated and stored in a password pro- tected computer"
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Single blind
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition is described but analyses of dif- ferences between completers and non-com- pleters was not conducted
Selective reporting (reporting bias)	Low risk	All data reported. Last observation carried forward used for 9/35
Size	High risk	17/18 per treatment arm

BDI: Beck Depression Inventory BPI: Brief Pain Inventory CBT: cognitive behavioural therapy CSQ: Coping Strategies Questionnaire HADS: Hospital Anxiety and Depression Scale MPQ: McGIII Pain Questionnaire RCT: randomised controlled trial TOPS: Treatment Outcomes in Pain Survey

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Crisostomo 2008	Not randomised
Hale 2007	Primary aim of study was not opioid reduction
Krymchantowski 2003	< 10 participants in each arm at post-treatment
Potter 2010	Primary aim of study was not opioid reduction
Roland 2011	Primary aim of study was not opioid reduction
Townsend 2008	Not randomised
Weinstein 2006	Primary aim of study was not opioid reduction

Characteristics of ongoing studies [ordered by study ID]

NCT02602535

Trial name or title	Mindfulness-oriented recovery enhancement for chronic pain and prescription opioid misuse in primary care
Methods	RCT, single-blind
Participants	People with chronic pain
Interventions	Mindfulness-oriented recovery enhancement vs support group
Outcomes	Primary outcomes (baseline to 6 months) Current Opioid Misuse and/or Addiction Behaviors Checklist and/or urine screen (triangulation aggregate) - change in opioid misuse BPI - pain severity and interference Secondary outcomes Change in opioid craving Depression Anxiety Stress Scale Opioid doses converted into morphine equivalents Other outcomes Change in nonreactivity CSQ Cognition Emotion Regulation Questionnaire Ways of Savoring Checklist
Starting date	2016
Contact information	Dr Eric Garland, University of Utah

NCT02602535 (Continued)

Notes	clinicaltrials.gov/ct2/show/NCT02602535?term=Mindfulness-Oriented+Recovery+Enhancement&rank=1
NCT02935621	
Trial name or title	Behavioral interventions for active duty service members and veterans with chronic pain
Methods	RCT, single-blind
Participants	Active duty service members and veterans with chronic pain conditions
Interventions	Mindfulness-oriented recovery enhancement vs support group
Outcomes	Primary outcomes (baseline to 6 months) Current Opioid Misuse and/or Addiction Behaviors Checklist and/or urine screen (triangulation aggregate) - change in opioid misuse BPI - pain severity and interference Secondary outcomes Change in opioid craving Depression Anxiety Stress Scal Opioid doses converted into morphine equivalents Post-traumatic checklist - military version Response to Stressful Events Scale Other outcomes Change in nonreactivity CSQ Cognition Emotion Regulation Questionnaire Ways of Savoring Checklist
Starting date	2016
Contact information	Dr Eric Garland, University of Utah
Notes	clinicaltrials.gov/ct2/show/NCT02935621?term=Mindfulness-Oriented+Recovery+Enhancement&rank=4

BPI: Brief Pain Inventory CSQ: Coping Strategies Questionnaire RCT: randomised controlled trial

DATA AND ANALYSES

This review has no analyses.

WHAT'S NEW

Last assessed as up-to-date: 4 January 2017.

Date	Event	Description
17 February 2017	New search has been performed	This review has been updated to include the results of a new search in January 2017. Three new studies were identified
17 February 2017	New citation required but conclusions have not changed	We included three new studies including 192 partici- pants in this update. Conclusions have not changed

HISTORY

Protocol first published: Issue 1, 2013

Review first published: Issue 9, 2013

Date	Event	Description
30 March 2016	Amended	Contact details amended.

CONTRIBUTIONS OF AUTHORS

Draft the protocol	EF/CE
Develop a search strategy	CE
Search for studies (usually 2 authors)	EF/LH
Obtain copies of studies	EF/LH
Select which studies to include (2 + 1 arbiter)	EF/LH/CE
Extract data from studies (2 authors)	EF/LH
Enter data into RevMan	EF

(Continued)

Carry out the analysis	N/A
Interpret the analysis	All authors
Draft the final write-up of the review	EF/CE
Update the review	CE
Methodologist name	CE
Statistician name	Gavin Stewart

DECLARATIONS OF INTEREST

CE: none known.

EF: none known.

KHT: none known. KHT is a public health specialist in training who is involved in the planning of health services for patients with chronic pain.

LH: none known.

SD: none known.

CS: none known. CS is a consultant anaesthetist working in pain management who manages chronic pain patients who are attempting to reduce prescribed opioids.

RK has attended advisory board meetings, received honoraria or held research grants for Grunenthal (2012 to 2015), Mundipharma Research (2016 to 2019) and Actavis (2017), who market opioid analgesics.

RAM has received grant support from Grünenthal relating to individual patient-level analyses of trial data regarding tapentadol in osteoarthritis and back pain (2015). He has received honoraria for attending boards with Menarini concerning methods of analgesic trial design (2014), with Novartis (2014) about the design of network meta-analyses, and RB on understanding pharmacokinetics of drug uptake (2015). He has received honoraria from Omega Pharma (2016) and Futura Pharma (2016) for providing advice on trial and data analysis methods.

SOURCES OF SUPPORT

Internal sources

• Oxford Pain Relief Trust, UK. Institutional support

External sources

• No sources of support supplied

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We did not search PsycLIT in either the first version of this protocol (Windmill 2013) or in this 2017 update, as stated in our protocol, due to the low yield and duplication between that database and the other databases searched. We contacted the authors of studies identified for inclusion to obtain additional data relevant to this review not included in the published articles. The original search for studies was intended to be completed without a time limit, but we limited inclusion to studies published in 2000 and later to reflect major changes since 2000 in prescribing of opioids to large numbers of people with chronic non-cancer pain; in this way, we worked to ensure that the review would have contemporary relevance. Although not explicitly stated in our protocol, we excluded studies of fewer than 10 participants in each treatment arm at post-treatment.

For the 2017 update, it was not possible to undertake planned data synthesis, sub-group analyses, or sensitivity analyses. We added GRADE assessment methods as is mandatory with Cochrane Reviews, however, it was not possible to conduct these assessments due to the heterogeneity of the studies. We have also removed 'blinding of personnel and participants' from the 'Risk of bias' assessment, as it is not possible to blind participants or personnel to psychological treatments. We searched trials registries for further trials not captured by our search method. We amended the first secondary outcome of 'pain intensity' to 'pain intensity/severity' for clarity.

INDEX TERMS

Medical Subject Headings (MeSH)

Analgesics, Opioid [*administration & dosage]; Chronic Pain [drug therapy; etiology; *therapy]; Cognitive Therapy [*methods]; Drug Tolerance; Electroacupuncture [*methods]; Observational Studies as Topic; Randomized Controlled Trials as Topic; Therapy, Computer-Assisted [methods]

MeSH check words

Humans