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ABSTRACT

Background

About one in every three adults are affected by lower limb muscle cramps. For some people, these cramps reduce quality of life, quality of sleep and participation in activities of daily living. Many interventions are available for lower limb cramps, but some are controversial, no treatment guidelines exist, and often people experience no benefit from the interventions prescribed.

Objectives

To assess the effects of non-drug, non-invasive treatments for lower limb cramp.

Search methods

We searched the Cochrane Neuromuscular Disease Group Specialized Register (13 September 2011) using the terms: cramp, spasm, contracture, charley horse and lower limb, lower extremity, foot, calf, leg, thigh, gastrocnemius, hamstring, quadriceps. We also searched CENTRAL (2011, Issue 3), MEDLINE (January 1966 to August 2011) and EMBASE (January 1980 to August 2011) and the reference lists of included studies. There were no language or publication restrictions.

Selection criteria

All randomised controlled trials of non-drug, non-invasive interventions trialled over at least four weeks for the prevention of lower limb muscle cramps in any group of people. We excluded, for example, surgery, acupuncture and dry-needling, as invasive interventions. We selected only trials that included at least one of the following outcomes: cramp frequency, cramp severity, health-related quality of life, quality of sleep, participation in activities of daily living and adverse outcomes.

Data collection and analysis

Two authors independently selected trials, assessed risk of bias and cross checked data extraction and analysis. A third author was to arbitrate in the event of disagreement. We asked the authors of five trials for information to assist with screening studies for eligibility and received four responses.
Main results

One trial was eligible for inclusion. All participants were age 60 years or over and had received a repeat prescription from their general practitioner of quinine for nighttime cramps in the preceding three months. This review includes data from only those participants who were advised to continue taking quinine. Forty-nine participants were advised to complete lean-to-wall calf muscle stretching held for 10 s three times per day. Forty-eight participants were allocated to a placebo stretching group. After 12 weeks, there was no statistically significant difference in recalled cramp frequency between groups. No “significant” adverse effect was reported. Limitations in the study’s design impede interpretation of the results and clinical applicability.

Authors’ conclusions

There is limited evidence on which to base clinical decisions regarding the use of non-drug therapies for the treatment of lower limb muscle cramp. Serious methodological limitations in the existing evidence hinder clinical application. There is an urgent need to carefully evaluate many of the commonly recommended and emerging non-drug therapies in well designed randomised controlled trials.

Plain Language Summary

Non-drug therapies for lower limb muscle cramps

Lower limb muscle cramps are a common problem that can affect any person, but cramps mostly occur during exercise, at nighttime in older people, in pregnant women, in people with a neurological disease and during kidney dialysis. Non-drug treatments are described as being effective for the treatment of muscle cramps. Non-drug treatments include muscle stretching, physical exercise, avoidance of physical fatigue, massage, relaxation, heat therapy, weight loss, sensory nerve stimulation, ankle splints worn while sleeping, and changes to sleeping and sitting positions. We did not include invasive interventions such as surgery, acupuncture or dry-needling in this review.

Only one randomised trial has assessed the effectiveness of a non-drug treatment for lower limb muscle cramp. This trial evaluated daytime calf muscle stretching to prevent nighttime muscle cramp in adults age 60 years and over who had received a repeat prescription of quinine for nighttime cramps in the preceding three months. Forty-nine participants were advised to complete lean-to-wall calf muscle stretching held for 10 s three times per day. Forty-eight participants were allocated to a placebo stretching group. After 12 weeks, there was no statistically significant difference in the frequency of cramps, as recalled by the participants, between groups. No “significant” adverse effect was reported. Owing to serious limitations in the design of the trial, it is impossible to determine from the available evidence whether or not calf muscle stretching can prevent recurrent lower limb muscle cramp.

Further research is required to determine the effectiveness of non-drug treatments for lower limb muscle cramp.
### SUMMARY OF FINDINGS FOR THE MAIN COMPARISON

#### Calf muscle stretching versus placebo stretching for lower limb muscle cramps

**Patient or population:** patients with lower limb muscle cramps  
**Settings:** general practice  
**Intervention:** lean-to-wall calf muscle stretching held for 10 s, three times per day versus placebo (sham) stretching

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<td><strong>Lower limb cramp frequency</strong></td>
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<tr>
<td>Participant recall</td>
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<td>⚫⚫⚫⚫ very low&lt;sup&gt;1,2&lt;/sup&gt;</td>
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<td>Follow-up: 12 weeks</td>
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<td><strong>Lower limb cramp pain severity</strong></td>
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<tr>
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<td>0 (0)</td>
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<td>See comment</td>
<td>Not estimable</td>
<td>94 (1 study)</td>
<td>⚫⚫⚫⚫ very low&lt;sup&gt;1,2&lt;/sup&gt;</td>
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*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
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.

---

1 Participants were allowed to swap intervention groups after six weeks. Trial authors provided only 12-week follow-up data. Allowed treatment changes included: discontinuation of quinine, discontinuation of any form of stretching exercise and, if the participant was initially allocated to placebo stretching group, beginning the stretching exercise program. It is not known how many of the participants within the stretching and placebo stretching groups included in this review modified their intervention at six weeks.

2 Total sample size is lower than the calculated optimal information size and the 95% CI includes the point of no effect and the upper limit crosses the minimal important difference for benefit.
Background

Description of the condition

Muscle cramps are sudden, involuntary, painful contractions of skeletal muscle (Norris 1957; Jansen 1991a; El-Tawil 2010) that are characterised electrically by repetitive firing of motor unit action potentials at high rates up to 150 per s (Miller 2005). Cramps can be excruciatingly painful (Kanaan 2001; Jansen 2002) and can cause delayed onset muscle soreness (a type of painful muscle damage that typically produces pain for 2 to 14 days) (Leung 1999; Jansen 2002). Muscle cramps can also interfere with sleep (Gootnick 1943; Gentili 1997; Gulich 1998) and activities of daily living (Matsumoto 2009); cause distress (Naylor 1994; Butler 2002); reduce quality of life (Kanaan 2001; Miller 2005; Shaker 2005; Kobrin 2007); limit sports participation and performance; and lead to chronic undertreatment in people undergoing haemodialysis (Canzanello 1992; Kobrin 2007). In one of the earliest case series reports, Gootnick 1943 reported three people whose sleep was so troubled by cramps during the night that they had been unable to sleep for months except in an armchair. Muscle cramp can occur at any time throughout the day or night (Abdulla 1999) and can affect people of any age (Jansen 1991; Miller 2005), although they are uncommon in children less than eight years of age (Leung 1999). Cramps usually affect muscles of the legs (Naylor 1994; Hirai 2000), particularly the calf muscles (Jansen 1991; Manjra 1996; Abdulla 1999; Leung 1999). Most cramps resolve spontaneously within 10 minutes (Naylor 1994; Abdulla 1999) but are reported to last up to one hour (Abdulla 1999). In most cases, cramps occur irregularly, often appearing and disappearing spontaneously for long periods, irrespective of treatment (Jansen 1999).

No large, randomly sampled population studies have been conducted to determine the prevalence of lower limb muscle cramps. In a nationwide, randomised study of general muscle cramps in nearly 2000 Dutch adults, approximately one in every three people reported at least one muscle cramp during the prior year (Jansen 1991a). Similarly, of over 1000 people surveyed in southern Germany, approximately one in every three people reported a history of nocturnal calf cramp and one in every four had experienced cramp within the four weeks prior to completing the questionnaire (Gulich 1998). In a separate study of 365 people aged 65 years and over, one in two people reported leg cramps at rest (for example while sleeping) (Abdulla 1999).

Compared to the general population, leg cramps are more prevalent in pregnant women (Jansen 1991; Jansen 1991a); people over 60 years of age (Naylor 1994; Gulich 1998; Hirai 2000); people undergoing haemodialysis (Canzanello 1992; Kobrin 2007); people with cirrhosis (Angeli 1996), fibromyalgia (Yunus 1996), arthritis (Yunus 1996; Abdulla 1999), lower motor neuron disease (for example amyotrophic lateral sclerosis) (Forshew 2003), lumbar spinal canal stenosis (Matsumoto 2009), peripheral neurological deficit (Haskell 1997), paraesthesia (Yunus 1996) or various veins (Gulich 1998; Hirai 2000); and children and adults with Charcot-Marie-Tooth disease (Burns 2009; Redmond 2009). There is no consistent relationship between cramp prevalence and gender (Jansen 1991; Naylor 1994; Manjra 1996; Gulich 1998; Abdulla 1999; Hirai 2000), serum electrolyte levels (Oboler 1991; Haskell 1997; Sulzer 2005), dehydration (Manjra 1996; Sulzer 2005; Schwellnus 2007), heart disease (Oboler 1991; Haskell 1997; Gulich 1998; Abdulla 1999) or diabetes mellitus (Oboler 1991; Gulich 1998).

Description of the intervention

Many interventions are available for lower limb muscle cramps. The most common interventions can be broadly categorised as drug or non-drug. Common drug interventions include quinine sulphate (El-Tawil 2010), gabapentin (Miller 2005), magnesium (Frussu 1999; Roffe 2002; Sohrabvand 2006), vitamin E (Connolly 1992; Burnakos 2000); calcium channel blockers (Baldodano 1988); nifedipine oxalate (Young 1993) and calcium (Sohrabvand 2006). Non-drug interventions reported in the literature include: muscle stretching (Daniell 1979; Jones 1983; Simchak 1991; Leclerc 1996; Leung 1999; Kanaan 2001; Miller 2005; Matsumoto 2009); massage (Jones 1983; Kanaan 2001; Matsumoto 2009); relaxation (Joes 1982); sensory nerve stimulation (Bentley 1996); foot wear changes (Roberts 1965); weight loss (Roberts 1965); physical exercise (Miller 2005); avoiding physical fatigue (Roberts 1965); heat therapy (Jones 1983); compression garments (Young 2009); night ankle dorsiflexion splints (Miller 2005); placebo (Butler 2002); and changes to sleeping (Gootnick 1943; Moss 1948; Cutler 1984; Warburton 1987; Abdulla 1999; Leung 1999; Kanaan 2001) and sitting (Roberts 1965) position. More controversial historical interventions include sleeping with a horseshoe (Simchak 1991), a magnet (Fowler 1973), corks (Warburton 1987) or potatoes (Warburton 1987) beneath the mattress.

Why it is important to do this review

Management of leg cramps can be frustrating (Riley 1995) and difficult (Butler 2002) for physicians and patients alike. Many people experience no benefit from the interventions prescribed (Sontag 1988; Abdulla 1999; Miller 2005) and many more receive no intervention at all (Naylor 1994; Gulich 1998; Abdulla 1999). Only two Cochrane systematic reviews have focused on interventions for muscle cramps. These reviews assessed the treatment of muscle cramps in pregnancy (Young 2002) and the effectiveness of quinine for muscle cramps (El-Tawil 2010). Two other reviews, currently in protocol stage, will evaluate intervention for muscle cramp in amyotrophic lateral sclerosis (Weber 2003) and magnesium for skeletal muscle cramps (Garrison 2011). The Cochrane systematic review of quinine for muscle cramps (El-Tawil 2010)
concluded that, despite moderate quality evidence that quinine significantly reduces cramp frequency, intensity and cramp days more than placebo, alternative interventions for muscle cramp should be investigated due to the risk of adverse effects. A systematic review of non-drug interventions for lower limb muscle cramps will serve two purposes. Firstly, it will enable healthcare providers, users, and policy decision makers to make better informed decisions about the treatment of lower limb muscle cramps. Secondly, it will identify for researchers interventions that require further evaluation in clinical trials.

OBJECTIVES

To evaluate the effectiveness of non-drug interventions for lower limb muscle cramps.

METHODS

Criteria for considering studies for this review

Types of studies

All randomised controlled trials (RCTs) of non-drug interventions for the prevention of lower limb muscle cramps. We excluded trials of acute treatment of muscle cramps. We accepted any diagnosis of cramp unless there was clear contradiction with the broad definition used in this review (cramp as a sudden, involuntary, painful contraction of skeletal muscle). We planned to report definitions of 'cramp' used in included studies in the table Characteristics of included studies. We contacted authors for clarification if: (1) 'cramp' was diagnosed but not defined, (2) authors diagnosed commonly used synonyms for cramp (for example, spasm) or (3) authors described the problem in such a way that we suspected muscle cramp (for example, if they described the problem as a sudden, painful contraction of skeletal muscle).

Types of participants

We included all people (including men, women and children) who experience lower limb muscle cramp. We included cramp during any activity or condition, including (but not limited to) cramp associated with exercise, pregnancy, haemodialysis, sleep and neurological conditions.

Types of interventions

We included all non-drug and non-invasive interventions used for at least four weeks. These may have included, but were not limited to, muscle stretching, splints, massage, warmth, change in sleeping position, placebo, relaxation, footwear, weight loss, compression garments, transcutaneous nerve stimulation, muscle strengthening, regular exercise, and changes to physical activity. We excluded all drug interventions and invasive interventions (including surgery and acupuncture/dry needling). For this review, drug interventions included any intervention taken orally (such as tablets, capsules, tonics), by injection, by rectal or vaginal suppository or by inhalation. We also excluded topically applied medicines (for example glyceryl trinitrate patches); however, topically applied preparations that are not thought to alter body function directly and independently (such as oils used in massage) were eligible for inclusion.

Types of outcome measures

Primary outcomes

• Cramp frequency, measured as number of cramps per week

When analysing cramp frequency, we planned to use change from baseline frequency where possible. If change scores with measures of variability were not available for all trials included in the same analysis, we planned to combine change scores and follow-up values using standardised mean differences (SMD).

Secondary outcomes

• Adverse outcomes

• Cramp severity, including pain severity (measured using any validated pain assessment tool, for example, a visual analogue scale) and duration of cramp (measured in seconds)

• Health-related quality of life (measured by any validated assessment tool, for example, SF-36 survey)

• Quality of sleep (measured by any validated assessment tool, for example, MOSleep survey)

• Participation in activities of daily living, including physical activity (measured by any validated assessment tool)

If data from only one study were included in an analysis, we measured the outcome at the time point used in the original study. If data from more than one trial were included in an analysis, we planned to group the outcomes in the time points 'four weeks to three months' and 'more than three months'. In this instance, the time point 'more than three months' was planned to be the primary outcome and the time point 'four weeks to three months'
was planned to be a secondary outcome. We planned to standardise all outcome data combined in meta-analysis to an appropriate follow-up period, for example, cramps per week after four months combined with cramps per week after six months.

'Summary of findings' tables were prepared for the outcomes cramp frequency, cramp pain severity, cramp duration and adverse effects.

**Search methods for identification of studies**

**Electronic searches**

We searched the Cochrane Neuromuscular Disease Group Specialized Register (13 September 2011) using the terms 'cramp', 'spasm', 'contracture', 'charley horse' and 'lower limb', 'lower extremity', 'foot', 'calf', 'leg', 'thigh', 'gastrocnemius', 'hamstring' or 'quadriceps'. We also searched CENTRAL (2011, Issue 3), MEDLINE (January 1966 to August 2011) and EMBASE (January 1980 to August 2011). The detailed search strategies are in the appendices: Appendix 1 (MEDLINE), Appendix 2 (EMBASE) and Appendix 3 (CENTRAL). There were no language or publication restrictions.

**Searching other resources**

We checked the references of the included study for other suitable trials. The first author of the included study was contacted via e-mail to assist in identifying relevant unpublished and published trials.

**Data collection and analysis**

**Selection of studies**

Two authors (FB and KW) independently assessed the titles and abstracts of all trials identified by the search. The same two authors then independently assessed for inclusion full-text copies of potentially relevant studies. Authorship and results were not masked. We resolved disagreements by discussion between review authors or, when necessary, arbitration by a third review author (JB). We planned to contact study authors if arbitration by the third reviewer did not resolve the dispute. If this was unsuccessful, we were to report the disagreement in the review. This was not required. We sought guidance from the Cochrane Neuromuscular Group regarding the eligibility of one trial (Coppin 2005) for inclusion. Guidance was provided and followed.

**Data extraction and management**

One review author (FB) extracted data from the published report using a standardised, pilot tested form and a second review author (KW) performed cross checks. We contacted the study authors as necessary to provide missing information or clarifications. We planned to resolve disagreements by discussion between review authors or, if necessary, arbitration by a third review author (JB). This was not required.

**Assessment of risk of bias in included studies**

Two review authors (FB and KW) independently rated risk of bias of the included trial using the following criteria as described in the *Cochrane Handbook of Systematic Reviews of Interventions* (Higgins 2011):

- sequence generation;
- concealment of allocation;
- blinding;
- incomplete data;
- selective outcome reporting;
- other sources of bias, such as financial conflicts of interest and single author or single centre trials.

Each criterion was assigned a judgement of 'high risk', 'low risk' or 'unclear risk' relating to the risk of bias for that entry. 'Unclear risk' indicates an unclear or unknown risk of bias; or if an entry is not relevant to the study at hand. Results are presented in the 'Risk of bias' table. A 'Risk of bias summary' figure was generated using Review Manager 5 to present all of the judgements in a cross-tabulation of study by entry.

If we were required to contact study authors for missing information regarding risk of bias, we planned to ask open-ended questions to reduce the risk of leading, for example the open-ended question 'How did you decide which treatment the next patient should receive?' was to be used in preference to 'Did you conceal the allocation sequence?'. This was not required.

**Measures of treatment effect**

Where possible, we analysed continuous data using mean differences (MD) and 95% confidence intervals (CI). If meta-analysis was required to pool data derived from different measurement scales, we planned to perform standardised mean difference (SMD) analyses. This was not required. We analysed count data (for example number of muscle cramps) as continuous data (mean number of cramps per person per week).

We planned to report results for dichotomous data as risk ratios (RR) with 95% CI.

**Unit of analysis issues**

We planned to include cross-over trials; however, only data from the first phase of intervention were to be considered, as clinical trials of interventions for muscle cramp have detected significant
carryover effects (particularly for quinine) (Dunn 1993; Jansen 1997; Coppin 2005) and the effectiveness of washout periods for interventions for muscle cramps is unknown. Due to the substantial and variable carryover effect of quinine, we excluded studies that discontinued quinine less than three months prior to intervention within the trial.

Dealing with missing data
Where available, we planned to extract data from intention-to-treat analyses. If the original researchers did not perform intention-to-treat analyses and sufficient raw data were available, we planned to complete intention-to-treat analyses before entering data into the Cochrane statistical package Review Manager (RevMan) 5.1.2 (RevMan 2011), to limit attrition bias.

Assessment of heterogeneity
We planned to assess clinical heterogeneity across trials and if trials were sufficiently clinically homogenous in terms of participants, interventions, and outcomes we would have included them in meta-analysis.
We planned to quantify inter-trial statistical inconsistency using I² (Higgins 2011).
The I² value was to be calculated by:
$I^2 = 100\% \left[ (Q-df)/Q \right]$ where Q is Cochran’s heterogeneity, Chi² statistic and df is the degrees of freedom. The Cochran’s Q is determined by summing the squared deviations (SDs) of each trial’s estimate from the overall meta-analytic estimate and a P value obtained by comparing the statistic with a x² distribution with k - 1 degrees of freedom (where k is the number of trials). We planned to use the following guide to interpret the I² values:
- 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 heterogeneity could not readily be explained, we planned to use a random-effects model to incorporate heterogeneous trials in meta-analysis.

Data synthesis
We used RevMan for statistical analyses. One author (FB) entered statistical data into RevMan and a second author (KW) checked the data.

Subgroup analysis and investigation of heterogeneity
Providing sufficient data were available, we planned to perform subgroup analysis according to cramp type/aetiology, for example, rest cramp versus exercise-associated cramp versus haemodialysis cramp versus cramp in people with neuromuscular disease.

Meta-analyses of subgroups were to follow the same methodological principles as the primary analysis. If meta-analyses were performed within multiple subgroups, we planned to compare the magnitude of effect estimates for subgroups informally between groups. We were to have considered non-overlap of summary estimate CIs to indicate statistical significance. If CIs overlapped to a small degree, we would have considered the possibility of a statistically significant difference (Higgins 2011).
We did not plan to compare the statistical significance of the pooled effect estimates for individual subgroups as the likely unequal information loading subgroups would have affected the power to detect effects (Higgins 2011).

Sensitivity analysis
We planned to perform sensitivity analyses by including only studies of high methodological quality (low to moderate risk of bias). If we had found all trials to have a high risk of bias, we planned to perform sensitivity analyses by excluding trials not concealing allocation or blinding the participants.
If one or more outliers were found to contribute to heterogeneity, and a reason for the outlying result was apparent, we planned to perform analyses both with and without outlying trials as a component of sensitivity analysis (Higgins 2011).

RESULTS

Description of studies
See: Characteristics of included studies; Characteristics of excluded studies.

Results of the search
Electronic searches retrieved a total of 1351 citations (247 from CENTRAL; 717 from EMBASE; 356 from MEDLINE; and 31 from the Cochrane Neuromuscular Disease Group Specialized Register). Of these, we identified 19 as potentially relevant. We obtained translation of published reports from French, Swedish, Chinese and German as necessary, to determine eligibility. Searching of reference list of the included trial did not retrieve any additional potentially relevant trials. We contacted authors of five trials to provide additional information to assist with screening; four replied (Vayssairat 2000; Coppin 2005; Warke 2006; Schyns 2009); one did not (Carpenter 1995).

Included studies
One trial with a total of 191 participants was eligible for inclusion (Coppin 2005). Details of this trial are presented in the table.
Characteristics of included studies. This trial was identified by electronic searching. All participants were age 60 years or over and had received a repeat prescription from their general practitioner of quinine for nighttime cramps in the preceding three months. Participants were allocated to one of four groups: (1) advised to discontinue taking quinine and begin a calf stretching program; (2) advised to discontinue taking quinine and begin a placebo stretching program; (3) advised to continue taking quinine and begin a calf stretching program; and (4) advised to continue taking quinine and begin a placebo stretching program. No washout period was provided for those participants advised to discontinue taking quinine, which excludes two groups from this review. This review includes only the 97 participants who were advised to continue taking quinine. Of these participants, 49 (51%) were male, the mean age was 74.3 years (standard error of the mean (SEM 0.6)) and at baseline the recalled number of cramps in the previous four weeks was 10.4 (SEM 1.76). Each participant was allocated to either a calf stretching or placebo stretching group. Calf stretches were performed by leaning into a wall for 10 s three times per day. The stretches and placebo stretches are described in detail in the table Characteristics of included studies. After six weeks, participants were told that they could continue the stretching exercises if they wished and also decide themselves whether or not to continue taking quinine tablets. Participants in the placebo stretching group were also given an instruction sheet describing the stretching exercise program. Published results in Coppin 2005 are only for 1, 2 and 12 week follow-up and there is no separation within the stretching and placebo stretching groups of those participants who continued quinine versus those who discontinued quinine.

In an attempt to avoid the confounding effects of changing interventions and to exclude participants who were advised to discontinue taking quinine, trial authors were contacted to provide six-week follow-up data for the participants who were advised to continue taking quinine in the stretching and placebo stretching groups. Only 12 weeks follow-up data were provided for the primary outcome 'number of cramps in the last four weeks'.

Excluded studies
We excluded 18 of the potentially relevant trials upon reading the full-text versions. Reasons for exclusion were: 14 did not isolate lower limb muscle cramp as an outcome measure (Carpenter 1995; Cramp 1995; Vayssairat 2000; Thaler 2001; Lawani 2003; Laessøe 2004; Warke 2006; Wei 2006; Xue 2006; Goodwin 2007; Miller 2007; Ippolito 2008; Liu 2009; Schyns 2009); four provided the intervention for less than four weeks (Laessøe 2004; Goodwin 2007; Miller 2007; Ippolito 2008); three assessed only drug therapy (Stern 1966; Mauss 1970; Liu 2009); two were not randomised or controlled clinical trials (as defined by Higgins 2011); one did not isolate data from participants who reported lower limb muscle cramp (Nilsson 2008); and one used a non-validated scale to measure health-related quality of life (Nilsson 2008). Details of individual trials are presented in the table Characteristics of excluded studies.

Risk of bias in included studies
Assessment of risk of bias for Coppin 2005 is described in the table Characteristics of included studies and is summarised in Figure 1.
**Allocation**

Allocation order was concealed using sealed, numbered, opaque envelopes containing intervention instructions.

**Blinding**

Participants were blinded to their group allocation by comparing calf muscle stretching to "placebo exercise", which "was devised to be of comparable duration and simplicity to the intervention stretching exercise, but was passive and involved negligible stretching of the muscles of the calf and foot". Researchers "sought to minimise intervention contamination by terming the stretching and placebo exercises as 'standing' and 'lying' exercises, respec-

**Incomplete outcome data**

In the 12-week data provided by the authors, data is missing for one participant from the intervention group and two participants from the control group. Reasons for missing data are not provided.

**Selective reporting**

The online protocol (ISRCTN26051665) states that outcomes would include duration of leg cramps and subsequent GP atten-
After 12 weeks, participants were asked to recall how many nighttime muscle cramps they had experienced in the preceding 4 weeks. Participants might be subject to recall bias. As discussed under Included studies, participants were allowed to swap intervention groups after six weeks. When contacted, trial authors provided only 12-week follow-up data and did not respond to further request for six-week follow-up data. Permitted treatment changes after six weeks included discontinuation of quinine, discontinuation of any form of stretching exercise and, if the participant was initially allocated to the placebo stretching group, beginning the stretching exercise program. The study does not report how many of the participants within the stretching and placebo stretching groups included in this review modified their intervention at six weeks. This seriously limits the interpretability and clinical applicability of results.

Effects of interventions

See: Summary of findings for the main comparison Calf muscle stretching versus placebo stretching for lower limb muscle cramps

In Coppin 2005, of those participants advised to continue taking quinine, 49 were allocated to the stretching group and 48 were allocated to the placebo stretching group. No participant was lost to follow-up, yet owing to incomplete data, we included only 48 participants from the stretching group and 46 participants from the placebo stretching group in the analysis. As change-from-baseline frequencies and SD were not available for data analysis, plain follow up scores and SD were used. After 12 weeks, participants were asked to recall how many cramps they had experienced in the previous four weeks. The difference in mean number of recalled cramps between the stretching group (mean 10.02, SD 17.67) and the placebo stretching group (mean 8.83, SD 17.23) was not statistically significant (MD 1.19, 95% CI -5.86 to 8.25, Analysis 1.1). There were no “significant adverse effects” reported. No definition of “significant adverse effects” was provided.

Discussion

This review included only one study. The included study aimed to assess the effect of (1) calf stretching exercises and (2) cessation of quinine on nighttime muscle cramp. Only the effect of calf stretching exercises was considered in this review. This study compared 49 people randomly allocated to a lean-to-wall calf stretching group with 48 people randomly allocated to a placebo (fake) stretching group. Both groups performed their stretches for 10 s three times per day. All participants were also taking quinine for their muscle cramps. After six weeks, participants could decide for themselves whether or not to continue with their allocated stretches and quinine tablets. Outcome data were provided by the authors, but only for 12-week follow-up. At the 12-week follow-up, participants were asked how many cramps they had experienced in the last four weeks.

Summary of main results

There is very low quality evidence that lean-to-wall calf muscle stretching held for 10 s three times per day does not reduce frequency of nighttime cramp (of unspecified muscles) after 12 weeks in people aged 60 years and over who have also been prescribed quinine for muscle cramping. See Summary of findings for the main comparison.

Overall completeness and applicability of evidence

The evidence for the use of non-drug therapies for lower limb muscle cramps is overwhelmingly incomplete. Of all types of lower limb muscle cramp (for example, cramp associated with exercise, pregnancy, haemodialysis, sleep and neurological conditions) and of all available non-drug therapies (for example, muscle stretching, massage, weight loss, physical exercise, avoiding physical fatigue, heat therapy, compression garments, night ankle dorsiflexion splints, sensory nerve stimulation, footwear changes, and changes to sleeping and sitting position), only one small study has investigated calf muscle stretching for nighttime cramp in people aged 60 years and over who had been given a repeat prescription of quinine for night cramps in the previous three months. The applicability of the available evidence is severely limited by deficiencies in study design. These limitations are discussed in more detail below.

Quality of the evidence

The quality of the evidence included in this review is very low. Figure 1. Some strengths of the trial include: (1) the allocation order was concealed (that is, researchers and participants did not know to which group they would be allocated before agreeing to participate) and (2) participants were blinded to their group allocation (that is, participants were not aware if they were in the true stretching or fake stretching group). Some weaknesses of the trial include: (1) there is unexplained missing data for one participant in the true stretching group and two participants in the fake stretching group and (2) outcomes reported in the published paper are different to those listed in the registered online protocol (research plan). Other weaknesses and limitations of the study are discussed below.
Limitations of the study design

After six weeks, participants in both the stretching and placebo stretching groups were allowed to swap or discontinue stretching treatments and to decide for themselves whether or not to continue taking quinine. As we included only those participants who were advised at baseline to continue taking quinine, no published data were suitable for use in analysis and trial authors provided only 12-week follow-up data when contacted. It is not known how many participants in the stretching and placebo stretching groups included in this review swapped or discontinued stretching exercises, or discontinued taking quinine after six weeks. This study design does not reflect clinical practice and seriously limits interpretation as it is impossible to isolate the effect of the intervention assigned at baseline.

The portion of the published study included in this review is underpowered to detect a clinically important difference between groups according to the included study author's power calculations. Coppin 2005 estimated a minimum sample size of 128 participants with complete outcomes, well above the 94 participants with complete outcomes included in this review.

Limitations of the participants

Coppin 2005 recruited participants aged 60 years and over who had been given a repeat prescription of quinine for night cramps in the previous three months. It is not known whether or not the quinine was effective. No inclusion criterion addressed the number of cramps affecting the participant in a defined period (for example, one week). It is possible that some of the included participants no longer experienced nighttime muscle cramp, perhaps due to the effect of the prescribed quinine tablets. Also, no inclusion criterion addressed the location of the nighttime cramp. While nighttime cramp often affects the calf muscle, other muscles of the lower limb and upper limb muscles can be affected. It is not known how many of the included participants experienced nighttime calf cramp. This limits the justifiability of offering a stretching program that targeted only the calf muscles.

Limitations of the intervention

The stretching intervention advice offered in Coppin 2005 is described in full in the table Characteristics of included studies. To summarise, participants were asked to stretch both calf muscles at the same time by leaning towards a wall for 10 s, three times throughout the day.

As muscle stretching has never before been evaluated in a RCT for treating muscle cramp, no direct comparisons in the literature can be found. In a systematic review of stretching before or after exercising to reduce delayed onset muscle soreness, risk of injury, and athletic or sporting performance, calf muscle stretching consisted of two separate stretches of soleus and gastrocnemius muscles held for 20 s on each limb, with a total stretch time of 160 s (Herbert 2002). The stretches prescribed in Coppin 2005 may be less intense (by stretching both limbs at the same time) and held for a shorter time than is ideal and than calf stretches routinely recommended in clinical practice.

As discussed above, this calf muscle stretching intervention was given to all participants, regardless of the location of their nighttime muscle cramp. This may not reflect standard clinical practice, where stretching therapies are targeted at the affected muscles.

Potential biases in the review process

None

Agreements and disagreements with other studies or reviews

The effect of non-drug therapies for lower limb muscle cramp has never before been systematically reviewed.

Authors’ Conclusions

Implications for practice

Evidence from RCTs is too limited in quantity and quality to provide clinical direction for the use of any type of non-drug therapy for any type of lower limb muscle cramps.

Implications for research

This review highlights the urgent need for rigorously designed RCTs to evaluate the effectiveness of non-drug therapies for lower limb muscle cramps. Future research should explore the effectiveness of a wide range of the available non-drug therapies for cramp
associated with exercise, pregnancy, haemodialysis, sleep and neurological conditions. Published reports of future trials should include high quality descriptions of all aspects of methodology to enable appraisal and interpretation of results. Outcomes should at least include cramp severity and frequency, and measures of frequency should not rely on participants retrospectively recalling the number of cramps experienced over a period of time.

More research is required to determine clinical important difference values for people experiencing nighttime muscle cramp to improve power calculations in future research.

**ACKNOWLEDGEMENTS**

We thank Dr Richard Coppin and Professor Paul Little for providing raw data for use in this review and Françoise Schyns, Professor Deirdre Walsh and Professor Michel Vaysairat for responding to requests for additional information about their research to determine eligibility for inclusion.

The Cochrane Neuromuscular Disease Group editorial base is supported by the MRC Centre for Neuromuscular Disease, the Motor Neurone Disease Association and the Muscular Dystrophy Campaign.

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References to studies excluded from this review

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Goodwin 2007  **[published data only]**


Ippolito 2008  **[published data only]**


Laessoe 2004  **[published data only]**


Lawani 2003  **[published data only]**


Liu 2009  **[published data only]**


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Miller 2007  **[published data only]**


Nilsson 2008  **[published data only]**


Schyns 2009  **[published data only]**


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Simchak 1991

Sohrabvand 2006

Sontag 1988

Sulzer 2005

Warburton 1987

Weber 2003

Young 1993

Young 2002

Young 2009

Yunus 1996

* Indicates the major publication for the study

Non-drug therapies for lower limb muscle cramps (Review)
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Characteristics of included studies  [ordered by study ID]

<table>
<thead>
<tr>
<th>Methods</th>
<th>Factorial design randomised control trial for the first 6 weeks. After 6 weeks, participants were told that they could continue the exercises if they wished and also decide themselves whether or not to continue or resume taking quinine tablets. Participants were also given an instruction sheet describing the exercise programme and confirming the advice. Data collection took place between January 1999 and September 2001.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>Participants were recruited through 28 general practices in southern England. Database searching at each practice identified potential participants, who were then contacted by letter. Inclusion criteria: people aged 60 years and over who had been given a repeat prescription by their general practitioner of quinine for nocturnal (night) cramps in the previous 3 months. Cramp was not defined in the published report. Exclusion criteria: People with conditions significantly limiting their ability to undertake the stretching exercises (for example, severe osteoarthritis); leg pain that might be confused with cramp (for example, severe peripheral vascular disease); inability to reliably report on symptoms (for example, dementia); and patients who described being previously taught leg exercises aimed at preventing cramp.</td>
</tr>
</tbody>
</table>
| Interventions | Participants were given advice to either continue or discontinue taking quinine tablets (this review includes data only from participants who were allocated to continue quinine use) and to undertake a daily programme of either stretching or placebo exercises for which they received training. The placebo exercise was devised to be of comparable duration and simplicity to the intervention stretching exercise, but was passive and involved negligible stretching of the muscles of the calf and foot. An attempt was made to minimise intervention contamination by terming the stretching and placebo exercises as 'standing' and 'lying' exercises, respectively. Patients were then shown how to complete a 6-week diary record. One week later the nurse telephoned the patient to check that they were coping with the diary recording. Intervention advice was not repeated. For the stretching exercise the nurse gave participants the following instruction, "First remove your shoes, then stand facing a wall 2 to 3 feet away so that the tips of your fingers are just touching the wall. Lean forward keeping your heels in contact with the floor, keep your body straight and use your arms to control your forward tilt until you feel a moderately intense, but not painful, 'pulling sensation' in your calf muscles. This stretching position should be held for 10 seconds (count 1 elephant, 2 elephants, etc.). Stand upright and relax for 5 seconds, then repeat the stretch for a further 10 seconds. Carry out this sequence three times spread through each day". For the non-stretching exercise the nurse advised participants as follows, "First lie face up on your bed or on the floor so that your legs are out straight. Gently bend your knees up and move them towards your chest keeping your feet flat on the bed or floor. When your feet are as near your bottom as you can get them without causing discomfort, hold that position for 10 seconds (count 1 elephant, 2 elephants, etc.). Straighten your legs out and relax for 5 seconds then repeat the exercise for a further 10 seconds. Carry out...
this sequence three times spread through each day"

| Outcomes | Baseline: The baseline questionnaire recorded cramp symptoms according to frequency over the previous 4 weeks and severity on a 4-point descriptive scale. Overall symptom burden was assessed using the mean of 3 7-point Likert scale scores assessing "disrupted activity", "sleep quality", and "overall problems with cramp in the past week". Each Likert scale used the same format as the MYMOP (Measure Yourself Medical Outcome Profile). Quinine usage over the previous 4 weeks was also recorded. A 6-week diary was used by participants to record daily cramp symptoms, intervention compliance, and weekly symptom burden scores. 12 weeks from baseline, participants were sent a self-completion postal questionnaire. Non-responders were telephoned to record their questionnaire responses. The 12-week self-completion postal questionnaire recorded cramp symptoms, quinine usage and frequency of taught exercises over the previous week. |
| Notes | NHS Executive SE Project Grant SPGS787/SW |

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>&quot;Randomised by the research assistant using random number tables&quot;</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>&quot;Randomisation to the trial took place when the nurse opened a sealed, numbered, opaque envelope containing intervention instructions&quot;</td>
</tr>
</tbody>
</table>
| Blinding of participants and personnel (performance bias) All outcomes | Low risk | "The placebo exercise was devised to be of comparable duration and simplicity to the intervention stretching exercise, but was passive and involved negligible stretching of the muscles of the calf and foot. We sought to minimise intervention contamination by terming the stretching and placebo exercises as "standing" and "lying’ exercises, respectively”.
Indicator of successful blinding: "By 12 weeks people performed the exercises 3 to 4 times per week, and although the non-stretch group did them slightly more frequently, this was not significant (placebo = 3.79 times per week, stretch = 3.07 times per week, t = 1.71, P = 0.09)" |
### Characteristics of excluded studies  [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cappy 1999</td>
<td>Not a RCT</td>
</tr>
<tr>
<td>Carpenter 1995</td>
<td>Lower limb muscle cramp was not an isolated measured outcome (reported the percentage of participants who reported muscle cramps; not specific to the lower limbs). Authors did not respond to requests for more information</td>
</tr>
<tr>
<td>Cramp 1995</td>
<td>Lower limb muscle cramp was not a measured outcome</td>
</tr>
<tr>
<td>Goodwin 2007</td>
<td>Lower limb muscle cramp was not a measured outcome. Intervention given for less than 4 weeks</td>
</tr>
<tr>
<td>Ippolito 2008</td>
<td>Intervention given for less than 4 weeks. Unclear if a RCT. Lower limb muscle cramp was not an isolated measured outcome (symptom clump: pain, heaviness, paraesthesia, nocturnal cramps, restless leg syndrome and pruritus. As a group, rated from 0 (never present) to 3 (always present) )</td>
</tr>
<tr>
<td>Reference</td>
<td>Details</td>
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<tr>
<td>---------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Laessøe 2004</td>
<td>Lower limb muscle cramp was not a measured outcome (spasm in people with spinal cord lesion). Intervention given for less than 4 weeks.</td>
</tr>
<tr>
<td>Lawani 2003</td>
<td>Lower limb muscle cramp was not a measured outcome.</td>
</tr>
<tr>
<td>Liu 2009</td>
<td>Lower limb muscle cramp was not an isolated measured outcome (symptom clump: leg pain or joint pain). Did not assess non-drug therapy (dietary advice).</td>
</tr>
<tr>
<td>Mauss 1970</td>
<td>Did not assess non-drug therapy (horse chestnut seed extract versus placebo).</td>
</tr>
<tr>
<td>Miller 2007</td>
<td>Lower limb muscle cramp was not a measured outcome (spasticity in people with multiple sclerosis). Intervention given for less than 4 weeks.</td>
</tr>
<tr>
<td>Nilsson 2008</td>
<td>Unable to isolate participants with nocturnal leg cramps from participants with restless leg syndrome. Authors did not respond to requests for more information. Lower limb muscle cramp pain was not a measured outcome. A 10-point visual analogue scale was used as a measure of quality of life, from 0 (absence of symptoms) to 10 (worst symptoms). It was considered that 0 = poor quality of life due to discomfort; 10 = condition did not diminish the quality of life. This scale is not validated.</td>
</tr>
<tr>
<td>Schyns 2009</td>
<td>Lower limb muscle cramp was not a measured outcome (spasm and spasticity in people with multiple sclerosis). Confirmed by study author Francoise Schyns.</td>
</tr>
<tr>
<td>Stern 1966</td>
<td>Did not assess non-drug therapy (papaverine hydrochloride).</td>
</tr>
<tr>
<td>Thaler 2001</td>
<td>Lower limb muscle cramp was not an isolated measured outcome (symptom clump: leg pain, discomfort or cramps; rated as a group as unchanged, better or worse).</td>
</tr>
<tr>
<td>Vayssairat 2000</td>
<td>Lower limb muscle cramp was not an isolated measured outcome (symptom clump: lower limb pain, cramps and heaviness; rated as a group). Confirmed with author Michel Vayssairat.</td>
</tr>
<tr>
<td>Warke 2006</td>
<td>Lower limb muscle cramp was not a measured outcome (spasm in people with multiple sclerosis). Confirmed by study author Deirdre Walsh.</td>
</tr>
<tr>
<td>Wei 2006</td>
<td>Lower limb muscle cramp was not a measured outcome.</td>
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<tr>
<td>Xue 2006</td>
<td>Lower limb muscle cramp was not a measured outcome.</td>
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(Continued)
DATA AND ANALYSES

Comparison 1. Calf muscle stretching versus placebo stretching for lower limb muscle cramps

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Number of cramps in the last four weeks</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
</tbody>
</table>

HISTORY

Protocol first published: Issue 4, 2010

Review first published: Issue 1, 2012

CONTRIBUTIONS OF AUTHORS

FB was responsible for selecting trials; contacting authors of studies; assessing risk of bias; extracting, entering and analysing data; interpreting results; and writing the review. KW was responsible for selecting trials; assessing risk of bias; checking data extraction and entry; and providing comments on the draft review. JB and VC were responsible for providing comments on the draft review. All authors agreed on the final text.

DECLARATIONS OF INTEREST

FB received a research grant from The University of Newcastle, Australia, to support completion this review. The University of Newcastle did not interfere with the independence of the authors in regard to the conduct of the review and did not delay or prevent publication of this review.

VC: none known

KW: none known

JB receives research funding from the National Health and Medical Research Council of Australia, NIH Rare Diseases Inherited Neuropathies Consortium, Charcot-Marie-Tooth Disease Association of Australia, Australian Podiatry Education & Research Foundation.

SOURCES OF SUPPORT

Non-drug therapies for lower limb muscle cramps (Review)

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Differences between protocol and review

‘Risk of bias’ assessment was updated to suit revision of the Cochrane Handbook for Systematic Reviews of Interventions guidelines.

A ‘Summary of Findings’ table was included to suit revision of the Cochrane Handbook for Systematic Reviews of Interventions guidelines.

We clarified the ‘Types of participants’ description in the Methods. The published protocol stated ‘All people (including men, women and children) who experience lower limb muscle cramp will be included.’ We added the following sentence to clarify: “We included cramp during any activity or condition, including (but not limited to) cramp associated with exercise, pregnancy, haemodialysis, sleep and neurological conditions”. This change did not alter our inclusion criteria.

Index terms

Medical Subject Headings (MeSH)

*Lower Extremity; Muscle Cramp [“therapy”]; Muscle Relaxants, Central [therapeutic use]; Muscle Stretching Exercises [*methods]; Quinine [therapeutic use]; Randomized Controlled Trials as Topic

MeSH check words

Aged; Humans; Middle Aged