Effect of magnesium therapy on nocturnal leg cramps: a systematic review of randomized controlled trials with meta-analysis using simulations

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Abstract

Background and objective. Nocturnal leg cramps (NLC) are common in primary care and may cause severe pain and sleep disturbance. We systematically reviewed the effectiveness of magnesium in treating NLC and the side-effect profile of magnesium compared to placebo.

Methods. We searched Medline, Embase, Cochrane Library, ClinicalTrials.gov, the International Standard Randomised Controlled Trial Number and the International Clinical Trials Registry Platform registries until August 2012. All randomized controlled trials (RCTs) comparing magnesium therapy for NLC in adults with any other comparator were eligible. Two investigators independently selected, extracted data from and rated the risk of bias of relevant studies. To compensate for the heterogeneity in outcome measures, simulations were used to summarize the data.

Results. Seven RCTs were included in the review (n = 361), all comparing magnesium to placebo. Three of these trials included only pregnant women. The difference in the median number of leg cramps per week between the placebo and the intervention groups was 0.345 (quantile 2.5%: −0.133, quantile 97.5%: 0.875). This difference was 0.807 (quantile 2.5%: 0.015, quantile 97.5%: 1.207) in the three studies involving only pregnant women and 0.362 (quantile 2.5%: −0.386, quantile 97.5%: 1.148) in the others. Overall gastrointestinal side effects were slightly more common with magnesium therapy than with placebo. The strength of this evidence was weak, mainly due to small study sizes and short follow-up.

Conclusions. Magnesium therapy does not appear to be effective in the treatment of NLC in the general population, but may have a small effect in pregnant women. Further research using better designed RCTs is necessary.

Key words: Magnesium, nocturnal leg cramps, primary care, systematic review.

Introduction

Nocturnal leg cramps (NLC) are defined as painful involuntary contractions of the lower limbs occurring during prolonged periods of rest, typically during the night (1). They may cause severe pain and sleep disturbance, and are particularly common among older adults, though they can occur in all decades of life (1–3). They are frequently unreported to physicians (2,4,5). In a general population survey carried out in the UK (n = 233), the overall prevalence of NLC was 37%, but the disorder was more
the objectives were to assess the effectiveness of this therapy, RCTs, including new studies undertaken since the last review, in NLC. Through a critical appraisal and synthesis of relevant information about any other published or unpublished trial that they might be aware of. The identification of relevant studies and the extraction of the data were carried out independently by two investigators (PS, DMH) in duplicate. Doubts and disagreements were resolved by discussion and consensus.

Methods

Design and search strategy

We conducted a systematic review of the literature through the electronic databases Medline, Embase and Cochrane Library (from inception to the end of August 2012), using the key words ‘magnesium’ and (leg or legs) and (cramp or cramps’), without any limits. The archives of the ClinicalTrials.gov registry, the International Standard Randomised Controlled Trial Number registry and the International Clinical Trials Registry Platform of the World Health Organization were consulted to identify unpublished clinical trials (using the key words ‘magnesium’ and ‘cramps’) and authors contacted to obtain additional data where necessary. The reference lists of all relevant papers were also scanned and local specialists in internal medicine, neurology and nutrition were contacted for information about any other published or unpublished trial that they might be aware of. The definition of relevant studies and the extraction of the data were carried out independently by two investigators (PS, DMH) in duplicate. Doubts and disagreements were resolved by discussion and consensus.

Study eligibility

All RCTs, assessing the effect of magnesium against any other comparator on NLC in pregnant women or in the general adult population (i.e. men and/or non-pregnant women) were eligible for inclusion. No language, publication date (or completion date for unpublished studies) or setting restrictions were applied, but studies in prepubertal children were excluded. We based the definition of NLC on the diagnostic criteria of the American Academy of Sleep Medicine (see panel).

Panel: Diagnostic criteria for NLC as published by the American Academy of Sleep Medicine (2005) (1):

1. The painful sensation in the leg or foot is associated with sudden muscle hardness or tightness, indicating a strong muscle contraction.
2. The painful muscle contractions in the legs or feet occur during the sleep period, though they may arise from either wakefulness or sleep.
3. The pain is relieved by forceful stretching of the affected muscles, releasing the contraction.
4. The sleep-related leg cramps are not better explained by another current sleep disorder, medical or neurologic disorder, medication use or substance use disorder.

Trials assessing treatments for exertion cramps, restless legs syndrome, periodic limb movements, vascular claudication,
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peripheral neuropathy and dystonia, as well as trials of treatments for cramps not located in the legs were excluded.

Data extraction
The following data were obtained from all the retrieved studies: design (parallel or cross over), details of the population (country, setting, pregnant women or general population, mean age and range, gender, number of subjects followed up, number lost to follow-up and number having refused to participate), type and dose of magnesium as well as length of treatment, criteria for eligibility, outcome measures used to assess the effectiveness of magnesium compared to placebo and side effects.

Data quality assessment
The two reviewers independently assessed the quality of the papers using Crombie’s check list and the Cochrane collaboration standard criteria (31,32). A summary assessment was prepared for each study across the following domains: sequence generation; allocation concealment; selection and attrition bias; blinding of participants and research staff; blinding of outcome assessors; sample size estimate; description of study population, design and measurements; analysis by intention to treat and assessment of statistical significance; role of funding body or industry; reporting of side effects of treatment; main study limitations. Doubts and disagreements were resolved by discussion and consensus.

Data analysis
A critical appraisal and synthesis of all relevant RCTs were performed. As NLC were measured in a large variety of ways (mean or median; number of days and/or nights with cramps or number of cramps), outcome data could not be summarized using conventional meta-analytic methods. In addition, the differences between the studies were important regarding the study population (pregnant women or general population), the study design (parallel or cross-over RCTs), the type of magnesium (oral or intravenous magnesium), the dose and length of treatment.

Monte Carlo simulations were therefore used in an attempt to overcome this heterogeneity in designs, primary outcomes and methods of summarizing these outcomes. When little information was available regarding data distribution, and because the distribution of leg cramps was usually highly asymmetric, we simulated the data from a log normal distribution so that the mean or the median, and the standard deviation fitted the reported data. Details of these simulations for each study can be found in the appendix (See online supplementary Table S1).

The cycle of simulation was repeated 10 000 times, and the main outcome was difference between the median number of cramps per week in the magnesium group and the median in the placebo group after the intervention. For each of the 10 000 simulations, the difference in the medians of each group was stored for each study, for the subset of studies dealing with pregnant women, for the subset of studies dealing with non-pregnant patients and for the entire set of patients. This allowed the estimation of 2.5%, 50% and 97.5% quantiles of the difference between the intervention and the control groups. All the analyses were undertaken using TIBCO Spotfire S+® 8.2 for Windows, TIBCO Software Inc., Palo Alto, CA.

This systematic review was carried out in accordance with the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guidelines (33). Assessment of publication bias was not performed because of the great heterogeneity and the overall small sample size of the studies included in the review.

Results
The study selection process is summarized in Figure 1. Six RCTs, involving a total of 321 patients, were retrieved through the electronic databases search (9,19,24,34–36), and one additional unpublished RCT (n = 40) was identified in the archives of the ClinicalTrials.gov registry (Table 1) (37). Table 2 presents the inclusion and exclusion criteria for the population involved in the trials.

The study population was restricted to pregnant women in three studies, whereas in the remaining four studies the outcomes were assessed in general populations. Two studies used a cross-over design. The sample sizes for all studies were small: the highest total number of subjects followed up was 80. In comparison to the number followed up attrition was particularly high in two studies [number lost to follow-up/number followed up (%) = 27/46 (59%) in Roffe’s and 7/38 (18%) in Nygaard’s trials]. The number declining to participate was not stated in four studies and was high in two studies in comparison to the number followed up (number declining participation/number followed up (%)) = 47/42 (112%) in Frusso’s and 17/46 (37%) in Garrison’s trials).

Type of magnesium treatment
One trial assessed magnesium therapy in infusion (4 hours for five consecutive days), whereas the others evaluated oral magnesium therapy, all against placebo. The type, dose and length of magnesium therapy (5 days to 6 weeks) were very different from one trial to another.

Quality of the studies
Overall, the quality of the evidence was poor with all studies presenting high risks of bias and lack of precision due to small sample sizes (Table 3). In particular, risks of selection, allocation and attrition bias were present or suspected to some extent in all studies. Many details were missing from the available reports in three
studies, including the unpublished trial, the author of which did not respond to our request for additional information (19, 24, 37). All the studies had only short-term follow-ups. Only the two studies carried out by Garrison and by Supakatisant could be considered as being at low risk of bias in terms of the role of the funding body or industry. Note that two studies used a cross-over design providing more precise comparisons and therefore requiring fewer subjects to achieve the same precision (because each individual is considered as his/her own control), but making the analyses more difficult to be interpreted because of residual effects (carry-over effects) (24, 34, 38). Support for the reviewers’ judgements is presented in the appendix (See online supplementary Table S2).

Effect of magnesium treatment on the number or severity of NLC
As the data presented in the four older papers were insufficient to allow calculation of effect measures with 95% confidence interval (CI), we present summary measures for the two groups and P values (Table 4). Only two studies, both involving pregnant women, showed a statistically significant effect of magnesium therapy with a larger reduction in the number and severity of NLC in the intervention group compared to the placebo group (19, 36). The third study in a pregnant population (n = 38) found no difference in outcomes between women receiving magnesium and those exposed to placebo (9).

Effect on other outcomes
Two studies reported patient self-evaluation of the effectiveness of treatment as an outcome. In both studies, the proportion of patients reporting that the treatment was effective was significantly higher in the group receiving magnesium (Table 4) (19, 34). Only one study examined cramp-related sleep disturbances and it found no effect of magnesium treatment on this outcome (24).

Figure 1. Flow chart for selection of studies, with indication of main reasons for exclusion.
Table 1. Characteristics of RCTs of magnesium in nocturnal leg cramps (Mg = magnesium group, Plac = placebo group, CO = crossover design, NK = information not provided in the paper)

<table>
<thead>
<tr>
<th>References</th>
<th>Study participants with nocturnal leg cramps, and setting</th>
<th>Design</th>
<th>Mean age (range or SD)</th>
<th>Number followed up, total (women)</th>
<th>Loss to FU, total (%)a</th>
<th>Number refusing to participate</th>
<th>Type and dose of magnesium per day</th>
<th>Length of treatment and FU</th>
<th>Outcome measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dahle (1995) (19)</td>
<td>Pregnant women referred from two prenatal care units in Sweden (n = 73)</td>
<td>Parallel</td>
<td>NKb</td>
<td>34 (34), 35 (35), 4 (6)</td>
<td>NK</td>
<td>Mg citrate lactate 360 mg (15 mmol)</td>
<td>3 weeks, FU 3 weeks</td>
<td>Number of cramps, severity, subjective assessment</td>
<td></td>
</tr>
<tr>
<td>Frusso (1999) (24)</td>
<td>Patients referred by family doctors at a University-based ambulatory clinic in Argentina (n = 45)</td>
<td>CO</td>
<td>62 (28–87)</td>
<td>NKc, NKc, 3 (7)</td>
<td>47</td>
<td>Mg citrate 1800 mg (72 mmol)</td>
<td>Two periods of 30 days separated by washout period of 30 days, FU after the two periods</td>
<td>Number of cramps, severity, duration, sleep disturbances</td>
<td></td>
</tr>
<tr>
<td>Roffe (2002) (34)</td>
<td>Subjects recruited by advertisements in the UK (n = 73)</td>
<td>CO</td>
<td>Mg: 61 (11), Plac: 64 (10)</td>
<td>17 (11), 29 (14), 27 (59)</td>
<td>NK</td>
<td>Mg citrate 300 mg (12 mmol)</td>
<td>Two periods of 6 weeks without washout period, FU after the final two periods of 4 weeks</td>
<td>Number of cramps, severity, duration, subjective assessment</td>
<td></td>
</tr>
<tr>
<td>Nygaard (2008) (9)</td>
<td>Pregnant women attending a routine ultrasound examination in Norway (n = 45)</td>
<td>Parallel</td>
<td>Mg: 32 (4)d, Plac: 30 (3)d</td>
<td>21 (21), 17 (17), 7 (18)</td>
<td>NK</td>
<td>Mg citrate lactate 360 mg (15 mmol)</td>
<td>5 days, FU 30 daysb</td>
<td>Number of cramps, severity</td>
<td></td>
</tr>
<tr>
<td>Garrison (2011) (35)</td>
<td>Patients recruited by advertisements in the offices of family doctors and in newspapers in Canada (n = 46)</td>
<td>Parallel</td>
<td>69 (8)</td>
<td>24 (19), 22 (13), 0</td>
<td>17</td>
<td>Mg sulphate 5g (20 mmol), 4 hours infusion</td>
<td>4 weeks, FU 4 weeks</td>
<td>Number of cramps, severity</td>
<td></td>
</tr>
<tr>
<td>Supakatisant (2012) (36)</td>
<td>Pregnant women attending an antenatal care clinic in Thailand (n = 86)</td>
<td>Parallel</td>
<td>Mg: 29 (6)e, Plac: 29 (5)e</td>
<td>41 (41), 39 (39), 6 (8)</td>
<td>9</td>
<td>Mg bis-glycinate chelate 300 mg</td>
<td>4 weeks, FU 4 weeks</td>
<td>Number of cramps, severity</td>
<td></td>
</tr>
<tr>
<td>Rosenbaum (unpublished, completed in 2009) (37)</td>
<td>Subjects recruited by radio advertisements in the USA (n = 40)</td>
<td>Parallel</td>
<td>67 (10)</td>
<td>20 (10), 20 (13), 0</td>
<td>NK</td>
<td>Mg lactate 336 mg</td>
<td>6 weeks, FU 4 weeks</td>
<td>Number of cramps, severity, duration, sleep disturbances</td>
<td></td>
</tr>
</tbody>
</table>

FU, follow-up.

aNumber lost to FU/number followed up.
bMean length of pregnancy of the 73 included subjects who started the study; 29 weeks (range 22–36 weeks); mean age not stated.
cTotal number of subjects followed up: 42; 73% women among the 45 included subjects who started the study.
dMean age of the 45 included subjects who started the study.
eIt is stated in the main text that the outcomes were also analysed after 3 months but FU data were only shown after 30 days.
fMean age of the 86 included subjects who started the study.
Table 5 lists the side effects reported in all the RCTs included in this review. The most frequent side effects in the trials using oral therapy concerned the gastrointestinal system (diarrhoea, nausea, vomiting, flatulence and constipation). Injection-related side effects, but no gastrointestinal side effects, were reported in the trial using intravenous treatment (35). There was no statistically significant difference in the frequency of gastrointestinal side effects ($P$ value 0.07). The mean difference was 0.13 (95% CI: −0.51 to 0.78), in favour of the placebo group. One severe side effect was reported: a stroke 31 days following a placebo infusion (35).

Simulations

As NLC outcome measures varied considerably from one study to another, standard meta-analysis could not be used to combine the results of these small trials. Simulations were therefore used, the results of which are shown in Figure 2.

The difference between the median number of cramps per week in the magnesium group and the median in the placebo group was 0.345 (quantile 2.5%: −0.133, quantile 97.5%: 0.875). In other words, a median patient would get one cramp less every 3 weeks when magnesium is taken rather than a placebo. The three studies in a pregnant population had a median difference of 0.807 (quantile 2.5%: 0.015, quantile 97.5%: 2.146).
<table>
<thead>
<tr>
<th>Reference</th>
<th>Sequence generation</th>
<th>Allocation concealment</th>
<th>Selection and attrition bias</th>
<th>Blinding of participants and research staff</th>
<th>Blinding of outcome assessors</th>
<th>Sample size estimate</th>
<th>Description of study population, design and measurements</th>
<th>Intention to treat analysis and assessment of statistical significance</th>
<th>Role of funding body or industry</th>
<th>Reporting of side effects</th>
<th>Main study limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frusso (1999) (24)</td>
<td>Unclear risk</td>
<td>Unclear risk</td>
<td>High risk</td>
<td>Unclear risk</td>
<td>High risk</td>
<td>High risk</td>
<td>Unclear risk</td>
<td>Unclear risk</td>
<td>Yes</td>
<td>Small sample size, short-term outcomes, much information lacking, cross-over design</td>
<td></td>
</tr>
<tr>
<td>Roffe (2002) (34)</td>
<td>Unclear risk</td>
<td>Low risk</td>
<td>High risk</td>
<td>Low risk</td>
<td>Unclear risk</td>
<td>High risk</td>
<td>Unclear risk</td>
<td>High risk</td>
<td>Yes</td>
<td>Small sample size, short-term outcomes, loss to follow-up, cross-over design</td>
<td></td>
</tr>
<tr>
<td>Nygaard (2008) (9)</td>
<td>Unclear risk</td>
<td>Unclear risk</td>
<td>High risk</td>
<td>Unclear risk</td>
<td>High risk</td>
<td>High risk</td>
<td>Unclear risk</td>
<td>Unclear risk</td>
<td>Yes</td>
<td>Small sample size, short-term outcomes, loss to follow-up</td>
<td></td>
</tr>
</tbody>
</table>
Table 4. Results of RCTs of magnesium in nocturnal leg cramps (Mg = magnesium group, Plac = placebo group, nb = number, NK = information not provided in the paper, NoD = no difference between the groups according to the paper)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Nb of cramps or nb of days and/or nights with cramps, mean (SD or 95% CI) if not specified</th>
<th>Severity index of cramps, mean (SD) if not specified</th>
<th>Duration of cramps in minutes, mean (SD) if not specified</th>
<th>Sleep disturbances index recorded on numerical scale, mean (SD) if not specified</th>
<th>Nb (%) of subjects saying that the treatment was effective</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mg</td>
<td>Plac</td>
<td><em>P</em> value</td>
<td>Mg</td>
<td>Plac</td>
</tr>
<tr>
<td>Dahle (1995) (19)</td>
<td>Median nb of days and/or nights decreased from every other day to once a week</td>
<td>Median nb of days and/or nights decreased from every other day to twice a week</td>
<td>&lt;0.05</td>
<td>-40.1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-20.4&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Frusso (1999) (24)</td>
<td>11.8 (7.6)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>11.1 (7.3)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.59</td>
<td>2.0 (1.7)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>2.1 (2)&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Roffe (2002) (34)</td>
<td>e</td>
<td>e</td>
<td>0.07</td>
<td>NoD</td>
<td>NoD</td>
</tr>
<tr>
<td>Nygaard (2008) (9)</td>
<td>Nb of days and/or nights in 2 weeks: 9.5 (5.1)</td>
<td>Nb of days and/or nights in 2 weeks: 7.7 (4.7)</td>
<td>0.27</td>
<td>13.2 (6.5)&lt;sup&gt;f&lt;/sup&gt;</td>
<td>11.4 (8.5)&lt;sup&gt;f&lt;/sup&gt;</td>
</tr>
<tr>
<td>Garrison (2011) (35)</td>
<td>-2.4/week (4.4)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>-1.7/week (3.3)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0.51</td>
<td>NoD</td>
<td>NoD</td>
</tr>
</tbody>
</table>

<sup>a</sup> Median nb of days and/or nights decreased from every other day to once a week
<sup>b</sup> Median nb of days and/or nights decreased from every other day to twice a week
<sup>c</sup> Median nb of days and/or nights decreased from every other day to twice a week
<sup>d</sup> Median nb of days and/or nights decreased from every other day to once a week
<sup>e</sup> Median nb of days and/or nights decreased from every other day to twice a week
<sup>f</sup> Median nb of days and/or nights decreased from every other day to twice a week
<table>
<thead>
<tr>
<th>Reference</th>
<th>Nb of cramps or nb of days and/or nights with cramps, mean (SD or 95% CI) if not specified</th>
<th>Severity index of cramps, mean (SD) if not specified</th>
<th>Duration of cramps in minutes, mean (SD) if not specified</th>
<th>Sleep disturbances index recorded on numerical scale, mean (SD) if not specified</th>
<th>Nb (%) of subjects saying that the treatment was effective</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mg</td>
<td>Mg</td>
<td>Mg</td>
<td>Mg</td>
<td>Mg</td>
</tr>
<tr>
<td></td>
<td>Plac</td>
<td>Plac</td>
<td>Plac</td>
<td>Plac</td>
<td>Plac</td>
</tr>
<tr>
<td></td>
<td><em>P</em> value</td>
<td><em>P</em> value</td>
<td><em>P</em> value</td>
<td><em>P</em> value</td>
<td><em>P</em> value</td>
</tr>
<tr>
<td>Supakatisant (2012) (36)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
<td>% Of subjects with 50% reduction of severity index: 69.8 (56–83.5)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
</tr>
<tr>
<td></td>
<td>−1.0/week (1.1)</td>
<td>−0.5/week (0.5)</td>
<td>NK</td>
<td>NK</td>
<td>NK</td>
</tr>
<tr>
<td>Rosenbaum (unpublished) (37)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 86 (75.6–96.3)</td>
<td>% Of subjects with 50% reduction of severity index: 69.8 (56–83.5)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
<td>% Of subjects with 50% reduction of nb of cramps: 60.5 (45.8–75.1)</td>
</tr>
</tbody>
</table>

*Compared to baseline; severity index recorded on numerical scale from 0 to 100 (0 = ‘insignificant’, 100 = ‘extremely painful’); it is not stated whether these are means or medians.

*Time unit not stated (probably month).

*It is only stated that severity index was recorded on numerical scale without additional information.

*Sleep disturbances index recorded on numerical scale from 0 to 10 (0 = ‘no sleep disturbances’, 10 = ‘could not sleep because of the cramps’).

*In the group starting with Mg: median of 9 (95% CI: 5–13) cramps per week in Phase 1 (Mg) and 8 (95% CI: 4–14) cramps per week in Phase 2 (placebo); in the group starting with placebo: median of 9 (95% CI: 5–13) cramps per week in Phase 1 (placebo) and 5 (95% CI: 4–8) cramps per week in Phase 2 (Mg); mean nb of leg cramps per week for each of the 12 weeks of the trial shown only graphically (exact nb and CIs not given).

*Severity index recorded on numerical scale from 0 to 4 (0 = ‘no pain’, 4 = ‘extreme pain’), then these scores were added and the mean value for each group was calculated. 

*Difference of reduction in the nb of cramps between the two groups: 0.7 cramps/week (95% CI: −3.1 to 1.7, *P* = 0.51); difference of reduction in the nb of cramps between the two groups in %: 5.5% (95% CI: −35.0% to 23.9%, *P* = 0.71).

*Mean % change of nb of cramps: 79.0% (70.2–87.8) in Mg group and 32.4% (3.4–61.5) in placebo group (*P* value 0.003); % of subjects without cramps: 48.8% (33.8–63.7) in Mg group and 27.9% (14.4–41.3) in placebo group (*P* value 0.04).
1.207), whereas the four others dealing with much older populations showed a median difference of 0.362 (quantile 2.5%: −0.386, quantile 97.5%: 1.148).

Discussion

Magnesium is widely prescribed in Europe and across the world to treat NLC (1). Yet uncertainty remains as to whether this treatment is beneficial (1,3,30). The aim of this systematic review was to clarify whether magnesium therapy is more efficient than placebo in treating NLC. This critical appraisal of seven RCTs showed weak evidence in favour of the effectiveness of magnesium in pregnant women, and unclear results in general populations. Meta-analysis using simulations confirmed the absence of a statistically significant effect of magnesium on the median number of leg cramps per week in general populations. A weak, but statistically significant effect was evidenced in pregnant populations. Yet the effect seemed small and the clinical significance of these findings is uncertain. Oral magnesium was associated with gastrointestinal side effects and intravenous magnesium with burning at the intravenous site.

Young et al. (3,30) previously suggested that oral magnesium had different effects in pregnancy than in the general population. The treatments they investigated in their two systematic reviews included magnesium, calcium, multivitamins and mineral supplements, sodium chloride and vitamin E for pregnant women, and quinine, analgesics, antiepileptic drugs, magnesium, vitamin E, compression hosiery and stretching exercises in the general population. Magnesium showed a statistically significant effect on the proportion of pregnant women reporting leg cramps, but no effect on NLC in the general population. This conclusion; however, rested on low-quality evidence due to the very low number of RCTs included and serious methodological limitations.

Our systematic review confirms that magnesium substitution is potentially beneficial in pregnant women but not in general, older populations. Garrison et al. (35) argues that this may be explained by differences in oral bioavailability of magnesium, the rate of magnesium absorption falling with age and by different

Figure 2. Estimation of the median difference in the number of leg cramps per week between the placebo and the intervention group (with the quantiles 2.5% and 97.5%) obtained by simulation. The sizes of the squares that stand for the median effects are proportional to the number of patients in the study. Studies marked with * included only pregnant women.
Magnesium therapy in nocturnal leg cramps

underlying aetiologies of NLC. Indeed, as pregnancy is a physiological state of low serum magnesium (36), pregnancy-related NLC is more likely to be due to magnesium depletion. Magnesium therapy could therefore be more efficient in this population.

High risk of bias and small sample sizes in all the studies included in this review limit the strength of the evidence it provides. In particular, since insufficient information was available in Dahle’s and Nygaard’s papers regarding blinding (see Table 5), the results showing a potential effect of magnesium in pregnant women could be due to a placebo effect if blinding was inadequate. Follow-up periods were short, even though subjects treated with oral magnesium often require relatively long treatments, because absorption of dietary magnesium decreases when dosage increases (ranges from 80% absorption at a low dietary intake to 10% at a high intake) and the steady state within tissue pools of magnesium is known to occur slowly (18,39). Finally, two studies included in the review used a crossover design. Although this may have provided more precise comparisons, using fewer subjects to achieve the same precision (each individual being considered as his/her own control), these studies carried additional risks of bias related to possible carry-over effects (38).

The rate of magnesium absorption falls with age and varies according to the type of magnesium preparation involved (35,40). The studies included individuals of many different ages and exposed them to a variety of different magnesium formulations. This may also explain why, despite biological plausibility and a general tendency of most studies to show a larger reduction in cramps in the intervention compared to the placebo groups, pooled results could not provide evidence in favour of magnesium treatment. We can hypothesize that the lack of effectiveness partly results from the limited bioavailability of oral magnesium, mainly in elderly people. Another explanation for the negative results relates to selection bias with the inclusion of many subjects whose leg complaints may have been confused with disorders not known to be related to magnesium deficiency such as restless leg syndrome or vascular claudication, for example (1). None of the trials mentioned all these disorders in the exclusion criteria (and in one study, no exclusion criteria were stated).

As expected, oral magnesium was often associated with gastrointestinal side effects. However, the symptoms were relatively minor and the same reactions occurred in the placebo groups, though generally less frequently. In addition, no serious and/or life-threatening adverse reaction was reported in relation to

Table 5. Side effects reported in RCTs of magnesium in nocturnal leg cramps (nb = number, NK = information not provided in the paper)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Magnesium group Nb followed up</th>
<th>Placebo group Nb followed up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dahle (1995)(^{a}) (19)</td>
<td>Slight nausea (infrequent)</td>
<td>Slight nausea (infrequent), severe nausea (1)</td>
</tr>
<tr>
<td>Frusso (1999)(^{\circ}) (24)</td>
<td>Diarrhoea/nausea/vomiting (10.7%)</td>
<td>Diarrhoea/nausea/vomiting (10.1%)</td>
</tr>
<tr>
<td>Roffe (2002)(^{\circ}) (34)</td>
<td>Diarrhoea (14), constipation (6), nausea/indigestion/flatulence (2), skin peeling (1)</td>
<td>Diarrhoea (8), constipation (11), nausea/indigestion/flatulence (4), bruising (1), headache (1)</td>
</tr>
<tr>
<td>Nygaard (2008)(^{b}) (9)</td>
<td>Gastrointestinal side effects (nausea, flatulence, diarrhoea, intestinal air) (6)</td>
<td>Gastrointestinal side effects (nausea, flatulence, diarrhoea, intestinal air) (6)</td>
</tr>
<tr>
<td>Garrison (2011)(^{\circ}) (35)</td>
<td>Asymptomatic hypotension (3), facial flushing (9), light-headedness (2), burning on the intravenous site (12) (five received normal saline solution to improve tolerability)</td>
<td>Facial flushing (7), on Day 31 post-infusions: stroke (1)</td>
</tr>
<tr>
<td>Supakatisant (2012)(^{\circ}) (36)</td>
<td>Nausea (11), diarrhoea (6)</td>
<td>Nausea (6), diarrhoea (1)</td>
</tr>
<tr>
<td>Rosenbaum (unpublished)(^{\circ}) (37)</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

\(^{a}\)95% CI and/or \(P\) values not provided in the paper. 
\(^{\circ}\)Total nb of subjects followed up: 42. 
\(^{\circ}\)Diarrhoea (\(P\) value 0.1); constipation (\(P\) value 0.2); nausea/indigestion/flatulence (\(P\) value 0.6). 
\(^{b}\)Nausea (\(P\) value 0.3); diarrhoea (\(P\) value 0.1).
magnesium treatment. These data confirm that oral magnesium is safe.

**Strengths and limitations of this review**

To our knowledge, this is the first systematic review of RCTs in which a meta-analytical approach was used to examine the evidence in relation to magnesium therapy to treat NLC. Our review methods followed PRISMA guidelines. We searched trial registration databases in addition to publication databases, and interrogated experts in the field, thus limiting the risk of excluding unpublished trials from our review.

Potential limitations relate to missing information in the papers published before 2000 and in the unpublished trial, which we were unable to complete by contacting the authors. As the trials we retrieved were not sufficiently similar in terms of study population, design, type of treatment and outcome measures, we could not use traditional meta-analytical methods to pool the results. We used simulations to overcome the problem. As most outcome distributions were not normal, we had to consider the difference of medians between the placebo and the intervention population in these simulations, rather than the difference in means. This approach leads to higher uncertainty and potential sources of bias across studies. Also, our meta-analysis did not allow a weighting proportional to the precision of the estimates, but used a more basic weighting based on the number of patients.

**Implications for clinical practice and future research**

The evidence that can be extracted from the current review suffers from the relatively low methodological quality of the included trials. Yet it suggests magnesium treatment is efficient in treating NLC in pregnant women, though a difference of less than two cramps per week compared to placebo, as computed in our meta-analysis, is probably not clinically significant. Although magnesium remains the cornerstone of NLC treatment across Europe and much of the world, this review highlights the lack of evidence that this treatment is truly effective outside pregnancy. Thus, there is no support for prescribing this treatment in everyday practices.

Physiopathology, patient feedback and clinical experience all concur to suggest that more research, less affected by bias, is needed to guide clinical practice in the future. New well-designed parallel group (or cross-over group, but with appropriate washout periods) RCTs are urgently needed, since NLC are common and alter the quality of life of the patients suffering from this condition. They should include either pregnant women or non-pregnant adults, exclude not only other conditions mimicking cramps (in particular, restless leg syndrome, periodic limb movements, vascular claudication and peripheral neuropathy) but also secondary causes of NLC which are unlikely to respond to magnesium substitution (e.g. NLC associated with other electrolytic abnormalities, such as hypocalcaemia or hyponatraemia). They should have sufficient power to allow a small-to-moderate real effect of magnesium therapy to be identified and have long-term follow-ups to allow time for the treatment to be efficient (>3 months). Primary care physicians in particular, who see the majority of patients presenting with this condition, should be encouraged to include patients in well-designed trials that address the limitations of previous studies and provide reliable data in favour or against the routine use of magnesium as treatment for patients suffering from NLC. The main outcome that could be assessed in these future trials could be the decrease of the mean (or median) number of NLC per week in the two groups, compared to baseline. These studies could also address a trade-off between a potential reduction of cramps and increase in gastrointestinal side effects related to the magnesium therapy. It would probably be useful to compare two different doses of the drug with the placebo in order to document the smallest effective dose.

In conclusion, this systematic review shows unclear results concerning the efficacy of magnesium therapy in the treatment of NLC in general populations and weak evidence in favour of this treatment for pregnancy-related NLC. This uncertainty results from relatively low methodological qualities of the trials included in the review. Further research using better designed RCTs is necessary to confirm whether physicians may continue to use this treatment for their patients’ benefit. Before these trials are available, the routine use of this treatment is not warranted, though magnesium therapy may be considered in pregnancy-related NLC.

**Supplementary material**

Supplementary material is available at Family Practice online.

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Ethical approval: we did not seek ethical approval for this study as it mainly involved the assessment of manuscripts published in the public domain.

Conflict of interest: the authors have nothing to declare.

References