BMJ Open Non-operative treatment for lumbar spinal stenosis with neurogenic claudication: an updated systematic review

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To cite: Ammendolia C. Hofkirchner C. Plener J. et al. Non-operative treatment for lumbar spinal stenosis with neurogenic claudication: an undated systematic review. BMJ Open 2022;12:e057724. doi:10.1136/ bmjopen-2021-057724

Prepublication history and additional supplemental material for this paper are available online. To view these files, please visit the journal online (http://dx.doi.org/10.1136/ bmjopen-2021-057724).

Received 27 September 2021 Accepted 15 December 2021

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ABSTRACT

Objectives Neurogenic claudication due to lumbar spinal stenosis (LSS) is a growing health problem in older adults. We updated our previous Cochrane review (2013) to determine the effectiveness of non-operative treatment of LSS with neurogenic claudication.

Design A systematic review.

Data sources CENTRAL, MEDLINE, EMBASE, CINAHL and Index to Chiropractic Literature databases were searched and updated up to 22 July 2020.

Eligibility criteria We only included randomised controlled trials published in English where at least one arm provided data on non-operative treatment and included participants diagnosed with neurogenic claudication with imaging confirmed LSS.

Data extraction and synthesis Two independent reviewers extracted data and assessed risk of bias using the Cochrane Risk of Bias Tool 1. Grading of Recommendations Assessment, Development and Evaluation was used for evidence synthesis.

Results Of 15 200 citations screened, 156 were assessed and 23 new trials were identified. There is moderatequality evidence from three trials that: Manual therapy and exercise provides superior and clinically important shortterm improvement in symptoms and function compared with medical care or community-based group exercise; manual therapy, education and exercise delivered using a cognitive-behavioural approach demonstrates superior and clinically important improvements in walking distance in the immediate to long term compared with self-directed home exercises and glucocorticoid plus lidocaine injection is more effective than lidocaine alone in improving statistical, but not clinically important improvements in pain and function in the short term. The remaining 20 new trials demonstrated lowquality or very low-quality evidence for all comparisons and outcomes, like the findings of our original review. Conclusions There is moderate-quality evidence that a multimodal approach which includes manual therapy and exercise, with or without education, is an effective treatment and that epidural steroids are not effective for the management of LSS with neurogenic claudication. All other non-operative interventions provided insufficient quality evidence to make conclusions on their effectiveness.

Strengths and limitations of this study

- This systematic review included a wide range of non-operative interventions commonly used in clinical practice.
- This review used consistent inclusion and exclusion criteria for neurogenic claudication, which included the corroboration of a diagnosis of lumbar spinal stenosis with imaging.
- This review used rigorous methods recommended by the Cochrane Back and Neck Pain Review Group including the use of Grading of Recommendations, Assessment, Development and Evaluation to synthesise and summarise the quality of the evidence.
- Only English studies were included in this review.
- Most studies had small samples sizes with heterogeneity in interventions tested, limiting ability to pool

PROSPERO registration number CRD42020191860.

INTRODUCTION

Lumbar spinal stenosis (LSS) causing neurogenic claudication is a highly prevalent and rapidly growing public health problem among older adults. It is characterised by bilateral or unilateral buttock pain and/or lower extremity discomfort, pain, weakness or heaviness precipitated by walking and prolonged standing and relieved by stooping forward and sitting.^{2 3} The underlying aetiology is usually age-related osteoarthritic changes to lumbar intervertebral discs, facets joints and ligaments leading to narrowing of the central and/or lateral spinal canals and compression and/or ischaemia of the spinal nerves.²⁴

Limited walking ability is the dominant impairment in neurogenic claudication and the most common reason for seeking care.⁵ Limited walking ability due to LSS is associated



with a significant decline in functional status, quality of life and independence in this population. ²⁵

Although LSS is the most common reason for spine surgery in older adults, most people with neurogenic claudication receive non-operative care.⁶ A course of non-operative care is also recommended prior to receiving surgical intervention.⁷ However, what constitutes effective non-operative care remains unknown. In 2013, we published a Cochrane review evaluating nonoperative treatment for LSS causing neurogenic claudication. 89 This review identified 21 randomised controlled trials (RCTs) assessing a variety of non-operative treatments. However, the quality of the evidence was deemed low or very low and therefore no conclusions could be made on the effectiveness of non-operative treatment for neurogenic claudication. The purpose of this study is to update this systematic review and the evidence for nonoperative treatments for neurogenic claudication. Our specific research question was: What non-operative interventions are effective in improving outcomes in patients with neurogenic claudication due to LSS?

METHODS

This systematic review was conducted and reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines. ¹⁰ We used methods recommended by the Cochrane Back Review Group. ¹¹

Patient and public involvement statement

Patients or the public were not involved in the conduct of this systematic review.

Population, interventions, comparison and outcomes criteria

The population of interest was individuals with imaging confirmed LSS (central or foraminal, with or without spondylolisthesis) and neurogenic claudication. Neurogenic claudication is a clinical diagnosis and was defined as buttock or leg pain and/or aching, numbness, tingling, weakness or fatigue with or without back pain, precipitated by standing or walking. There were no age restrictions. The interventions of interest included all non-operative treatments and the comparison was any treatment including surgery. Outcomes included at least one of the following measures: walking ability, pain intensity, physical function, quality of life or global improvement.

Search and study selection

We replicated and updated our original electronic data-base search (from 1966 to January 2011) up to 22 July 2020. The search was performed by an experienced librarian in CENTRAL (Cochrane Library 2011 issue1), MEDLINE, EMBASE, CINAHL and Index to Chiropractic Literature databases. The terms 'spinal stenosis', 'lumbar spinal stenosis', 'neurogenic claudication', 'lumbar radicular pain', 'cauda equina' and 'spondylosis' were combined with a highly sensitive search strategy to identify RCTs.

Reference lists of selected studies and previous reviews were also searched to identify additional articles. Online supplemental file 1 provides details on the full search strategies used for all databases.

Studies were included if they were RCTs published in peer-reviewed English journals, at least one arm of the trial provided data on effectiveness of a non-operative treatment and at least 80% of subjects had neurogenic claudication with imaging confirmed LSS. Studies evaluating subjects with radiculopathy caused by disc herniations without neurogenic claudication were excluded.

Studies with mixed populations were only included if separate data for subjects with neurogenic claudication due to LSS were provided.

Two pairs of reviewers independently screened all titles and abstracts identified by the search strategy. Full text of articles deemed to be potentially relevant were independently assessed by two reviewers who made the final decision for inclusion. A third reviewer was consulted if consensus was not reached.

Risk of bias assessment and data analysis

Two reviewers independently assessed methodological risk of bias and performed data extraction. Safety data (intervention side effects and/or complications) when available were also collected. The Cochrane Risk of Bias Tool 1 was used that included the 12-item criteria recommended by the Cochrane Back Review Group. Discrepancies in risk of bias scoring and data extraction were resolved with discussion and if necessary, with a third reviewer until consensus was reached. Reviewers who were authors of any of the included studies were recused from performing risk of bias assessment, data extraction, data analysis or synthesis of their own studies.

Low risk of bias was defined as fulfilling 6 or more of the 12 criteria including clearly described and appropriate randomisation (item A), and allocation concealment (item B), and with no severe flaws. A severe flaw was defined a priori as a serious methodological deficiency not captured by the 12-item criteria that significantly increases the risk of bias such as very high dropout or cross-over rates and sample sizes less than 30 subjects per treatment arm.

For each comparison, outcomes were analysed according to these follow-up time periods: immediate (up to 1 week following the intervention); short term (between 1 week and 3 months); intermediate (between 3 months and 1 year) and long term (1 year or longer). Outcome data were pooled, and meta-analyses were performed when trials were judged to be sufficiently homogeneous, both clinically and statistically.

Rehabilitation therapy was defined as treatment that used any combination of education, exercise instruction, manual therapy, heat and cold applications, electrotherapy, other physical therapy modalities, orthosis and other assistive devices. Multimodal treatment included various combinations of rehabilitation therapy



treatments, oral and other mediations and spinal injections, but not surgery.

Data synthesis

The quality of the evidence for each outcome and for each comparison was evaluated using Grading of Recommendations, Assessment, Development and Evaluation (GRADE). Development and Evaluation (GRADE). Overall quality of the evidence was based on performance against five domains: (1) risk of bias; (2) consistency of findings; (3) directness of comparisons; (4) precision of estimates and (5) other considerations such as selective reporting.

The quality of the evidence starts at high when there are consistent findings among at least 75% of RCTs with low risk of bias and consistent, direct and precise data and with no known or suspected publication bias. It downgrades a level for each domain not met. Treatment effects between comparators (more effective, less effective or no difference) were based on statistically significant and clinically important differences in outcomes.

High-quality evidence

All five domains are met; further research is very unlikely to change the confidence in the estimate of effect.

Moderate-quality evidence

One of the domains is not met; further research is likely to have an important impact on the confidence in the estimate of effect and may change the estimate.

Low-quality evidence

Two domains are not met; further research is very likely to have an important impact in the confidence of the estimate of effect and is likely to change the estimate.

Very low-quality evidence

Three or more domains are not met; there is great uncertainty about the estimate of effect.

Evidence provided by a single small trial was considered inconsistent and imprecise and thus provide 'low' or 'very low' quality evidence, depending on whether it was assessed as having a low or high risk of bias, respectively, and there were no other limitations. Studies with both low risk of bias and inappropriate or unclear randomisation and/or treatment allocation techniques were downgraded by two levels for the 'risk of bias' domain.

The results below are reported based on statistically significant differences between comparators for each outcome using data reported by authors. Differences considered clinically important will be specified when the quality of the evidence is moderate or higher. The minimal clinical important differences (MCIDs) used are listed in the online supplemental table 2. Adverse events for the new studies are detailed when reported by the authors.

RESULTS

Selection and description of included trials

We screened 15200 titles and abstracts and assessed 156 full-text articles. This resulted in 44 RCTs meeting the inclusion criteria, including 23 new trials. Figure 1 summarises original and updated screening results. Online supplemental table 1 describes the characteristics of all included trials. In total, 3792 participants (1765) males, 1836 females and 191 participants of undisclosed gender¹⁴ 15 were randomised to one of the 60 comparison groups. In total, 17 studies evaluated rehabilitation therapy or multimodal care, 14 16-31 11 assessed epidural injections, ³²⁻⁴² 7 evaluated oral medications, ^{15 43-48} 6 assessed calcitonin, ^{49–54} 2 evaluated acupuncture ^{55 56} and 1 assessed spinal manipulation.⁵⁷ Thirty-eight trials were conducted at tertiary care or university affiliated centres and six at medical/rehabilitation clinics. 18 24 35-38 The mean age of participants was 63.3 years. The duration of symptoms varied considerably among the studies with a mean ranging from 12 weeks to 15 years. Follow-up periods also varied significantly ranging from immediately following the intervention to 10 years post intervention.

Risk of bias of included studies

The median and mean number of criteria met was 7 of 12 (range 2–11), see table 1.

Although 31 studies met six or more criteria, only 9 were considered to have low risk of bias. ^{19 20 24 27 28 31 37 42 43 56} Among the remaining 22 studies that met six or more criteria, 13 failed to explicitly describe and/or use appropriate randomisation procedures, allocation concealment or both ^{16–18 30 32–34 39 41 48 52 54 57}; 3 had severe flaws due to high cross-over rates, ^{21 22 25} which made the intention-to-treat analyses uninterpretable and 6 had other serious flaws including premature stopping of the trial, ⁴⁷ large number of participants lost to follow-up ⁴⁰ and small sample size (less than 30 participants per arm). ^{26 29 46 55}

Evidence of effect of interventions

Overall, 53 of the 60 comparisons were examined in a single trial, most with small sample sizes. It was only possible to combine data from two trials (assessing surgery vs multimodal treatment) for one outcome in a meta-analysis. ^{19 22} The five other studies (all assessing calcitonin) ^{49-52 54} were combined qualitatively. The results of these pooled analyses were published in our previous reviews. ^{8 9} Heterogeneity in source population, intervention and outcome instruments precluded pooling of data from other trials. Online supplemental table 2, summary of GRADE assessment and outcomes, summarises the quality of the evidence for outcomes for each comparison.

Calcitonin

There were no new studies assessing calcitonin. The conclusion from our previous review was that there is very low-quality evidence from six trials $(N=231)^{49-54}$

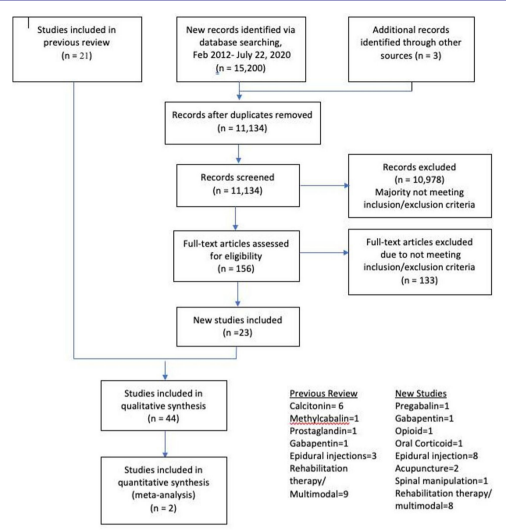


Figure 1 Study flow diagram.

that calcitonin is no better than placebo or paracetamol regardless of mode of administration or outcome assessed.

Oral medication

We identified four new studies assessing five oral medications. There is low-quality evidence based on one small cross-over trial (N=29)⁴⁶ that pregabalin does not improve pain, distance walked, function or global health status immediately following the intervention compared with placebo. Adverse events were reported in 64% of the pregabalin group, the most common being dizziness, compared with 35% in the placebo group.

A small trial evaluating gabapentin plus conservative care $(N=45)^{48}$ provides very low-quality evidence demonstrating no significant improvement in back/leg pain, disability scores or global health in the short term compared with conservative care plus botulinum toxin injection. Five patients (20.8%) reported mild-tomoderate pain at injection sites for a few days after botulinum toxin injections.

There is very low-quality evidence from one small trial $(N=24)^{47}$ that oxymorphone hydrochloride or

propoxyphene and acetaminophen are no better than placebo in the immediate term for all outcomes assessed.

A single small trial provided very low-quality evidence (N=61)¹⁵ that oral corticoids do not improve outcomes in the short term compared with placebo.

The original review identified three studies assessing oral medications and concluded that there is low-quality evidence that prostaglandins improve walking distance and leg pain in the short term compared with etodolac (a non-steroidal anti-inflammatory drug);⁴³ that there is very low-quality evidence that gabapentin improves walking distance and pain compared with placebo in the intermediate and long term⁴⁵ and that methylcobalamin (vitamin B 12) plus conservative treatment improves walking distance in the intermediate and long term compared with conservative treatment alone.⁴⁴

Rehabilitation therapy and multimodal treatment

We identified eight new studies evaluating 13 rehabilitation therapy and/or multimodal treatment approaches, with one study being compared with surgery.

Table 1 Risk of bias a	ssessme	nt for studie	s on non-c	Risk of bias assessment for studies on non-operative treatment for lumbar spinal stenosis with neurogenic claudication	ment for lum	bar spinal s	stenosis with	neurogenia	claudicati	nc			
Author	٨	В	c	D	E F		G H	_	_	J.	Х	٦	Total
Calcitonin													
Eskola <i>et al⁵¹</i>	خ		+	+	·		+		5	5	5	+	5
Porter and Hibbert ⁴⁹	خ		1	خ خ	+		· ·		1	خ	+	+	4
Porter and Miller ⁵⁰	خ		+	ذ ذ	-		+		خ	خ	خ	+	4
Podichetty et al ⁵²	5	٠	+	+	1		+		+	5	خ	+	9
Tafazal et al ⁵⁴	خ	. خ	+	+	+		+	,	ı	خ	خ	+	7
Sahin et al ⁵³	خ	٠	1	+	l		+	,	+	5	خ	+	4
Oral medications													
Prostaglandin													
Matsudaria ⁴³	+	+	,	+	+		÷		+	5	خ	+	7*
Methylcabalin													
Waikakul and Waikakul ⁴⁴	ı	۔ خ	1	+	+		÷	_	+	خ	خ	+	2
Gabapentin													
Yaksi <i>et al</i> ⁴⁵	خ	۔ خ	ı	1	٠.		+		خ	خ	خ	+	3
Pregabalin													
Markman e <i>t al</i> ⁴⁶	+	+	+	+	+		+		خ	+	ı	+	10‡
Gabapentin													
Park et al ⁴⁸	+		+	+	+		+		<i>-</i>	خ	ı	+	\$\$
Oxymorphone hydrochloride													
Markman e <i>t al⁴⁷</i>	+	+	+	+	+		+		خ	+	+	+	948
Oral corticoid													
Rodrigues and Natour ¹⁵	+	+	ځ	5	+		÷		خ	خ	خ	+	5
Rehabilitation therapy or multimodal care	imodal care												
Goren <i>et al²⁴</i>	+	+	ı	+	+		+		+	5	5	+	7*
$Koc et al^{23}$	خ	٠ -	1	+	+		+		+	5	5	+	5
Pua et al ²⁰	+	+	ı	+	+		+	_	+	5	ı	+	7*
Whitman et a/ ¹⁸	+		1	+	+		+	1	+	5	ċ	+	7
Minetama et al ³⁰	+		ı	+	+		+	_	5	+	+	+	\$\$
Schneider et al ³¹	+	+	1	+	1		+		+	5	+	+	*∞
Ammendolia et al ²⁷	+	+	ı	+	+		+	_	+	+	+	+	10*
Oğuz et al ¹⁴	خ	٠ -	1	c- -	ذ خ		+		خ	5	5	+	2
Homayouni <i>et al²⁶</i>	+	+	ı	+	+		+		I	+	5	+	7†
Marchand et al ²⁹	+	+	1	+	ر. د		+	1	5	1	+	+	7†
Kim et al ²⁸	+	+	+	+	+		+	_	5	+	+	+	11*
Spinal manipulation													
Passmore <i>et al⁵⁷</i>	ı	+	ı	+	+		+		+	+	+	+	18
													Continued



Table 1 Continued													
Author	A	В	ပ	۵	ш	ш	5	I	_	7	¥	_	Total
Acupuncture													
Kim et al ⁵⁵	+	+	ı	ı	I	ı	+	+	ı	+	+	+	7†
Qin et al ⁵⁶	+	+	+	1	+	+	+	+	+	·	+	+	10*
Epidural injections													
Cuckler et a/ ³²	خ	~	+	+	+	+	+	+	+	۔ ۔	+	+	6
Fukusaki <i>et al³⁴</i>	خ	<i>-</i>	5	5	+	+	+	+	+	خ	+	+	7
Zahaar ³³	<i>-</i>	<i>-</i>	+	خ.	+	+	+	+	+	-	خ	1	9
Brown ³⁵	+	ı	+	ı	ć	+	+	ı		٠	ı	+	5
Friedly <i>et al</i> , ³⁷ Friedly <i>et al</i> ⁶² and Makris ⁶⁴	+	+	+	+	+	+	+	+	٠.	+	+	+	*
Song et al ³⁶	5	۲.	5	5	۲	+	+	I	٠.	+	+	+	5
Milburn et a/38	<i>-</i>	~	+	1	+	1	+	1			ı	+	4
Hammerich et a/ ⁴⁰	+	+	ı	ı	+	ı	+	ن	٠.	ı	+	+	19
Sencan et a/41	+	<i>د</i> -	+	1	+	+	5	+	+	+	ځ	+	#8
Wei <i>et al</i> ⁴²	+	+	+	ı	I	+	I	+	٠.	+	+	+	*∞
Percutaneous epidural adhesiolysis	siolysis												
Karm et al ³⁹	+	5	+	ı	+	I	+	+		·	ı	+	‡9
Surgery versus physical therapy	apy												
Zucherman <i>et al,</i> ¹⁷ Zucherman <i>et al</i> ⁶⁰ and Hsu <i>et al</i> ⁵⁹	c-	+	1	ı	+	+		+	+		+	+	₩9<
Weinstein et $al,^{21}$ Weinstein et al^{98} , Abdu et al^{61}	+	+	1	1	+	+	+	+	ć			+	++ *** 9<
Amundsen et al ¹⁶	+	<i>د</i>	ı	ı	I	+	+	+	ı	٠	ı	5	4
Malmivaara et a/ ¹⁹	+	+	1	1	+	+	+	+	+	5	5	+	*∞
Weinstein $et al^{22}$, Weinstein $et al^{99}$, Lurie $et al^{58}$	+	+	I	I	+	I	+	+	<i>c</i> -	٠.	ı	+	119
Delitto et al ²⁵	+	+	1	1	+	5	+	1	+	1	+	+	7††

intervention? F. Was the drop-out rate described and acceptable? G. Were all randomised participants analysed in the group to which they were allocated? H. Are reports of the study free of suggestion of selective outcome responsible? I. Were cointerventions avoided or similar? K. Was the compliance acceptable in all groups? L. Was the timing of the outcome assessment similar in all groups? +: Yes, A: Was the method of randomisation adequate? B: Was the treatment allocation concealed? C: Was the patient blinded to the intervention? D: Was the care provider blinded to the intervention? E: Was the outcome assessor blinded to the -: No, ?: Unclear. BMJ Open: first published as 10.1136/bmjopen-2021-057724 on 19 January 2022. Downloaded from http://bmjopen.bmj.com/ on April 23, 2024 by guest. Protected by copyright.

Low risk of bias if six or more items met, including valid randomisation and treatment allocation techniques and no severe flaws.

^{1&}lt;30 participants per treatment arm.

FTreatment allocation unclear. §Premature end of study.

^{¶2-}year follow-up drop-out rate 30%, 1 year<20%; intention to treat inconsistent at 2-year follow-up. *Drop-out rate<20% at 1 year, >20% at 4 years.

^{††}Severe flaw due to high cross-over rates.



There is moderate-quality evidence from one trial (N=259)³¹ that manual therapy and exercise provides superior and clinically important short-term improvement in symptoms and function compared with medical care or community-based group exercise and that community-based group exercise improves physical activity in the short term compared with medical care. There were no reported serious adverse events in any group. There was a significantly greater rate of transient joint soreness associated with the manual therapy and exercise group (49%) compared with the community-based group exercise (31%) and medical care (6%) groups.

Another trial provides moderate-quality evidence (N=104)²⁷ that comprehensive care (manual therapy, education and exercise delivered using a cognitive-behavioural approach) demonstrates superior and clinically important improvements in walking distance in the immediate, short, intermediate and long term and compared with self-directed home exercise. This study also provides low-quality evidence that comprehensive care improves overall pain and function in the long term compared with self-directed home exercises. At 12 months, none of the 43 participants in the comprehensive group and 2 of the 46 participants in the self-directed group experienced adverse events. These adverse events were mostly attributed to a temporary increase in low back and/or leg pain.

There is low-quality evidence from one trial (N=34)²⁸ that a form of manual therapy (Mokuri Chuna), acupuncture and physician care, with or without a herbal remedy (Gang-Chuk Tang), improves low back pain in the intermediate term compared with oral aceclofenac, epidural steroids and physical therapy (heat and TENS).

A single study assessing supervised physical therapy (manual therapy, exercise and body weight-supported treadmill) (N=86)³⁰ provides low-quality evidence for improved symptoms, function and walking distance in the short term compared with home exercises.

There is very low-quality evidence from one study (N=120)¹⁴ that heat, transcutaneous electrical nerve stimulation (TENS) and home exercise instruction are no better than isokinetic exercise in the immediate, short and intermediate term for all outcomes and less effective than unloaded exercises in the immediate and short term. Unloaded exercise was also found to be superior to isokinetic exercise in the immediate and short term.

One small single study (N=47)²⁶ provides very low-quality evidence that aquatic exercise is more effective than physical therapy (exercise, ultrasound, heat and TENS) in improving pain and walking distance in the immediate term.

Another small single trial (N=40)²⁹ provides very low-quality evidence that a presurgical exercise programme improves postsurgical outcomes in the immediate, but not in the short or intermediate terms.

There is low-quality evidence from one study (N=169)²⁵ that a structured physical therapy programme (education

and exercises) provides similar outcomes to decompression surgery in the long term (2-year follow-up). Overall, 9 out of 82 participants receiving physical therapy reported adverse events consisting of worsening of symptoms, whereas 33 out 87 participants reported surgery-related complications, mainly attributable to reoperation, delay in wound healing and surgical site infection.

Our original review identified nine rehabilitation therapy/multimodal trials of which five were compared with surgical interventions. A meta-analysis was conducted for two of the surgical trials. Two of the original surgical trials have since published 8-year follow-up results (see below). All studies provide either low-quality or very low-quality evidence.

A meta-analysis⁸ that includes two trials²² ¹⁹ shows that laminectomy improves outcomes only at the 2-year follow-up compared with conservative care. One of these studies shows no difference in outcomes after an 8-year follow-up.⁵⁸

An interspinous surgical implant ^{17 59 60} was found to be superior to multimodal treatment (epidural injections, pain medication, education, exercise, back brace, heat/ice and massage). Another trial ¹⁶ provided inconclusive evidence when comparing laminectomy with or without fusion to lumbar orthosis and education.

Among patients with degenerative spondylolisthesis, one study²¹ shows no difference in outcomes with laminectomy when compared with conservative care, including after an 8-year follow-up.⁶¹

One study showed that exercise plus ultrasound is no better than exercise plus sham ultrasound but better than no treatment, and exercise plus sham ultrasound is better than no treatment. Other studies demonstrated that inpatient physical therapy (ultrasound, heat and TENS) is more effective than home exercise plus oral diclofenac, unweighted treadmill walking plus exercise is no better than cycling plus exercise and unweighted treadmill are more effective than flexion exercises, walking and sham ultrasound. 18

Epidural injections

We identified six new studies evaluating epidural injections. There is moderate-quality evidence from one study (N=400)³⁷ 62 that glucocorticoid plus lidocaine injection is better than lidocaine alone in improving pain and function at 3 weeks (short term) but not at 6 weeks (short term), 12 weeks (intermediate term) or 12 months (long term). The improved outcomes at 3 weeks were statistically significant but not considered to be of clinical importance. A follow-up subgroup analysis using patient-prioritised Roland-Morris Disability Questionnaire items did not change the results. A total 21.5% of patients in the glucocorticoid-lidocaine group and 15.5% in the lidocaine alone group reported one or more adverse events (p=0.08). Adverse events included headaches, fever, infection, dizziness, cardiovascular/lung problems, leg swelling and dural puncture.

A small study (N=29)³⁶ provided very low-quality evidence that an injection of lidocaine is no better than a saline injection for all outcomes in the short term.

There is very low-quality evidence from one study $(N=57)^{38}$ that steroid injections at the level of maximal stenosis improve pain and function in the immediate and short term compared with steroid injections at two levels cephalad to the maximum level of stenosis.

A small trial (N=54)⁴⁰ provided very low-quality evidence that steroid injections are no better than steroid injections combined with physical therapy (manual therapy and exercise) in improving pain or function in the short term but are more effective in improving pain in the intermediate and long term.

There is very low-quality evidence from one study $(N=67)^{41}$ that interlaminar steroid injection improves pain and walking distance in the intermediate but not in the short term compared with transforaminal steroid injection.

A three-arm trial (N=30)⁴² provided low-quality evidence that tumour necrosis factor (TNF) alpha inhibitor (etanercept) injections improved pain and function in the immediate, short and intermediate term compared with steroid or lidocaine injections and that steroid injections were no better than lidocaine for all outcomes and follow-up periods.

There is very low-quality evidence from one small trial (N=38)³⁵ that minimally invasive lumbar decompression surgery is no better than epidural steroid injections for all outcomes in the short term.

One small trial (N=44)³⁹ provided very low-quality evidence that an epidural inflatable balloon catheter (ZiNeu) improves pain and function in the intermediate term but not in the short term compared with a balloonless catheter (Racz). Minor and transient adverse events were reported equally in both groups (no data provided), mostly pain and paraesthesia at the injection site.

Our original review identified four trials evaluating seven epidural injection approaches, all with very low-quality evidence for all outcomes. Two trials demonstrated that translaminar³² or caudal³³ steroid injections were no better than placebo. Two other trials showed that translaminar epidural steroid plus a block was better than placebo or an epidural block alone,³⁴ that translaminar epidural block was better than placebo³⁴ and that interlaminar epidural steroid plus a block was better than home exercise plus diclofenac or inpatient physical therapy (ultrasound, heat and TENS).²³

Acupuncture

We identified two new studies assessing acupuncture. There is low-quality evidence from one trial $(N=80)^{56}$ that acupuncture improves back and leg pain, symptoms and function in the immediate, short and intermediate term compared with sham acupuncture. Overall, 3 out of 40 participants in the acupuncture group reported short-term pain at the insertion site (one also had a haematoma) and 5 out of the 40 participants in the sham group

reported non-serious back pain or fatigue. There is very low-quality evidence from a small trial (N=50)⁵⁵ that acupuncture plus usual care is no better than usual care alone in the short term for all outcomes.

Spinal manipulation

We identified one study assessing spinal manipulation. There is very low-quality evidence from a very small trial (N=14)⁵⁷ that spinal manipulation alone is no better than a wait list control in the immediate term for all outcomes.

DISCUSSION

We updated our systematic review on non-operative treatments for LSS causing neurogenic claudication and identified 23 new trials that were added to the previous 21 studies. The highest number of studies, 17/44, evaluated rehabilitation therapy/multimodal treatment, 11 assessed epidural interventions, 7 assessed oral medications, 6 assessed calcitonin, 2 evaluated acupuncture and 1 assessed spinal manipulation. Of the 60 comparisons that were evaluated, 5 comparisons from three trials^{27 31 37} provided moderate-quality evidence. The remaining comparisons provide either low-quality or very low-quality evidence. In our original review, all comparisons for all the interventions assessed were of low-quality or very low-quality evidence. This lack of moderate-quality or high-quality evidence limited our ability to make conclusions on the effectiveness of most non-operative treatments.

There is now moderate evidence that a multimodal structured 6-week programme consisting of manual therapy and exercise with or without education is an effective treatment approach^{27 31} for neurogenic claudication and that epidural steroid injections do not provide clinically important improvements in short-term or long-term outcomes compared with epidural lidocaine injections. However, given that these respective findings came from single studies, this evidence lacks consistency and therefore there is a possibility that replicating these trials in the future might result in substantially different conclusions. However, a recent clinical practice guideline for the management of LSS leading to neurogenic claudication concurred with our findings and recommended, based on moderate-quality evidence, multimodal care consisting of education with home exercises and manual therapy. 65 These guidelines also recommended against the use of epidural steroid injections, based on high-quality evidence. A recent systematic review and meta-analysis of RCTs evaluating conservative non-pharmacological therapies for degenerative LSS also concluded, based on low-tomoderate evidence, that manual therapy and supervised exercises significantly improve outcomes compared with self-directed or group exercises. ⁶⁶ A recent clinical update published in the British Medical Journal recommended supervised exercise and manual therapy as a first-line treatment for LSS and recommended against the use of epidural steroid injections.⁶⁷ More dated systematic

reviews did not recommend a combination of education, exercise and manual therapy as an effective treatment for LSS. ^{7 68 69} However, these reviews did not include the more recent higher quality trials^{27 31} evaluating this multimodal approach.

A multimodal approach to the treatment of LSS would appear to be a rational approach given the complexity of neurogenic claudication with underlying physical, functional and psychosocial factors impacting recovery.⁷⁰ There is also a plausible rationale for the lack of effectiveness of epidural steroid injections for neurogenic claudication since the dominant underlying pathophysiological mechanism appears to be neuroischaemia rather than neuroinflammation.4

Although we cannot make firm conclusions about the effectiveness of non-operative treatments for neurogenic claudication, this review is important because it provides important information regarding the state of current evidence regarding non-operative treatments. This can be used to inform clinical practice guidelines and aid clinicians and patients in making clinical decisions regarding treatment options. This is particularly important with respect to interventions that have higher risks and costs such as epidural injections and surgery. About 25% of all epidural injections are performed for LSS⁷¹ 72 yet the evidence from our current review and those of others^{73–75} do not support their use. The number and associated costs of surgical procedures for degenerative LSS are growing, especially decompression surgery with complex fusion. 76 77 LSS continues to be the most common reason for spine surgery in older adults.^{6 76} High-quality evidence for the effectiveness of surgery is also lacking based on our current review and the findings of other systematic reviews.⁷⁸ Clinical trials evaluating surgery for LSS are difficult to conduct due to challenges in recruitment and blinding (patient and practitioner) and high costs.⁸⁰ One ongoing clinical trial is comparing decompression surgery with sham surgery which should help to evaluate the potential role of the placebo effect of surgery for LSS.81

Oral medication is often the first-line treatment in primary care management of LSS.5 Pregabalin and gabapentin are commonly prescribed medications for LSS despite the growing evidence that these medications are not effective for back-related leg symptoms and may cause more harm than good.82-84

New to this updated review are clinical trials on acupuncture and spinal manipulation; however, the quality of the evidence was insufficient to make conclusions on their effectiveness. A systematic review and meta-analysis of RCTs and controlled clinical trials published in Chinese found no conclusive evidence for the effectiveness and safety of acupuncture for LSS.85 Passive unimodal treatments such as acupuncture and spinal manipulation are unlikely to provide long-term benefit but more likely to provide benefit when combined with a comprehensive approach to managing LSS,²⁷ not unlike recommendations for managing chronic low back pain.⁸⁶

This review is also important because it provides a comprehensive assessment and identification of significant knowledge gaps in this area to guide future research. This includes the need for higher quality studies that assess commonly used non-operative treatments particularly in primary care settings that are adequately powered and have low risk of bias and long-term follow-up. Future RCTs should follow the Consolidated Standards of Reporting Trials guideline⁸⁷ when planning trials and reporting study findings in an attempt to improve transparency and reduce bias.

The strengths of this review include the evaluation of a wide range of non-operative interventions and the use of consistent inclusion and exclusion criteria for neurogenic claudication, which included the corroboration of a diagnosis of LSS with imaging. The use of these criteria to define the study population increases the likelihood that participants in the included studies had the diagnosis of neurogenic claudication due to narrowing of the central canal or lateral foraminae. 88–90 Other strengths of this review include the use of rigorous methods recommended by the Cochrane Collaboration, the WHO and the Cochrane Back and Neck Pain Review Group. 13 This included the use of the GRADE method to synthesise and summarise the quality of the evidence.

Limitations of this review include the potential for language bias because only English articles were accepted. We also included studies with small samples sizes which are more prone to high risk of bias. 91 Over half of the included studies had less than 30 subjects per arm at baseline and none of these studies could be pooled because of high heterogeneity across studies. However, the exclusion of studies with small samples sizes in this review would not have changed our conclusions. The definition of a severe flaw and the cut-off point of 6 or more to differentiate trials of low from high risk of bias were arbitrary, therefore alternative definitions and cut-off points or the use of other risk of bias tools could have impacted the findings and conclusions of this review. The validity of MCIDs used in this review is unknown. Although most were derived from studies with neurogenic claudication, 63 92 93 others were based on an arbitrary improvement of at least 30%. 94 There are no agreed upon MCIDs in LSS and therefore different MCIDs thresholds could have potentially altered our conclusions. The location and severity of the stenosis on imaging was not deemed important in this review. Imaging findings often do not correlate with patient symptoms or severity and therefore imaging by itself is a not reliable diagnostic tool in this population.^{67 95 96} Neurogenic claudication is the clinical entity of interest in this review and, although usually caused by LSS, the diagnosis is made clinically without imaging.⁹⁷ Neurogenic claudication symptoms, by definitions, improve with flexion, due to the increased volume around the involved nerve roots irrespective of where the stenosis is located (eg, centrally or at the lateral recess). However, it is uncertain whether the effectiveness of some interventions, such as epidural steroid injections, is dependent on



location of the spinal stenosis. This is a different research question requiring future research.

CONCLUSIONS

There is moderate-quality evidence that a multimodal approach that includes manual therapy and exercise, with or without education, is a safe and effective treatment and that epidural steroids are not effective for the management of LSS causing neurogenic claudication. All other studies evaluating non-operative interventions provided insufficient quality evidence, limiting the ability to make conclusions about their effectiveness. With the growing prevalence and significant personal, social and economic burden of LSS, more high-quality evidence for non-operative interventions is urgently needed to guide clinical practice.

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Acknowledgements The authors would like to thank librarian Maggie Tiong from the Institute for Work & Health for her assistance in developing the search strategy and retrieving relevant articles and Carly Coelho for assistance in screening titles and abstracts

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Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests CA received research funding from the Canadian Chiropractic Research Foundation and the Arthritis Society. JY has received funding support from the Danish Foundation for Chiropractic Research and Postgraduate Education, the Ontario Chiropractic Association, the Canadian Memorial Chiropractic College, the National Chiropractic Mutual Insurance Company Foundation and the University of Southern Denmark. CC holds a Research Chair in Knowledge Translation in the Faculty of Health Sciences, Ontario Tech University, funded by the Canadian Chiropractic Research Foundation.

Patient consent for publication Not applicable.

Ethics approval This study does not involve human participants.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

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