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

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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 15200 citations screened, 156 were assessed and 23 new trials were identified. There is moderate.

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 data.

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.¹ 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

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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 database 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.¹¹ 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).^{12 13} 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 gender¹⁴¹⁵ 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-up⁴⁰ 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

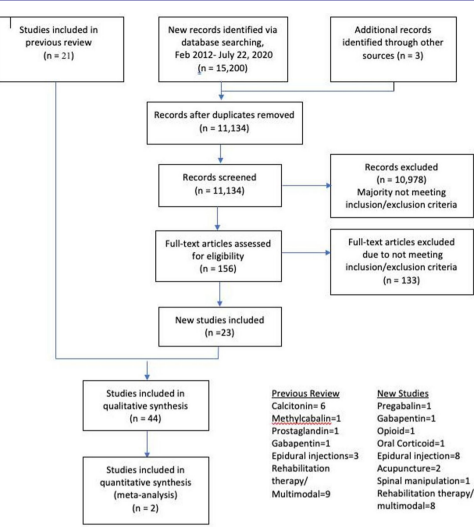


Figure 1 Study flow diagram.

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 $\left(N{=}24\right)^{47}$ that oxymorphone hydrochloride or

proposyphene 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.

Calcitonin												
Eskola <i>et al</i> ⁵¹ ?	6	+	+	+	2	+	1	2	2	2	+	5
Porter and Hibbert ⁴⁹ ?	ć	I	\$	ć	+	+	\$	I	ن د	+	+	4
Porter and Miller ⁵⁰ ?	ć	+	2	\$	I	+	+	ć	2	2	+	4
Podichetty <i>et al</i> ⁵² ?	د	+	+	+	1	+	1	+	۲	د	+	9
Tafazal e <i>t al</i> ⁵⁴ ?	ć	+	+	+	+	+	+	I	2	ć	+	7
Sahin <i>et al</i> ⁵³ ?	ć	I	I	+	1	ć	+	+	ć	ć	+	4
Oral medications												
Prostaglandin												
Matsudaria ⁴³ +	+	I	ļ	+	+	+	2	+	2	2	+	7*
Methylcabalin												
Waikakul and Waikakul ⁴⁴ -	ć	I	I	+	+	+	2	+	2	2	+	5
Gabapentin												
Yaksi <i>et al</i> ⁴5 ?	ć	I	I	I	ć	+	+	ć	2	2	+	e
Pregabalin												
Markman et al ⁴⁶ +	+	+	+	+	+	+	+	\$	+	I	+	10‡
Gabapentin												
Park <i>et al</i> ⁴⁸ +	ć	+	+	+	+	+	+	2	2	I	+	8‡
Oxymorphone hydrochloride												
Markman <i>et al</i> ⁴⁷ +	+	+	+	+	I	2	+	2	+	+	+	9†§
Oral corticoid												
Rodrigues and Natour ¹⁵ +	+	2	5	2	+	+	2	2	?	2	+	5
Rehabilitation therapy or multimodal care	l care											
Goren <i>et al</i> ²⁴ +	+	I	I	+	+	I	+	+	ć	ć	+	7*
Koc et al ²³ ?	ذ	I	I	+	+	+	I	+	\$	۲	+	5
Pua et al ²⁰ +	+	I	ļ	+	I	+	+	+	2	I	+	7*
Whitman <i>et al</i> ¹⁸ +	2	I	I	+	+	+	+	+	2	5	+	7
Minetama <i>et al</i> ³⁰ +	ż	I	I	+	+	+	+	2	+	+	+	8‡
Schneider <i>et al</i> ³¹ +	+	I	I	+	I	+	+	+	2	+	+	*0
Ammendolia <i>et al²⁷</i> +	+	I	I	+	+	+	+	+	+	+	+	10*
Oğuz et al ¹⁴ ?	2	I	I	د.	ړ	+	I	ć	2	2	+	2
Homayouni <i>et al</i> ²⁶ +	+	I	I	+	+	+	I	I	+	2	+	7†
Marchand <i>et al</i> ²⁹ +	+	I	I	+	ć	+	+	ذ	1	+	+	7†
Kim <i>et al</i> ²⁸ +	+	+	+	+	+	+	+	ړ	+	+	+	11*
Spinal manipulation												
Passmore <i>et al⁵⁷</i> –	+	I	I	+	+	+	I	+	+	+	+	8†
											0	Continued

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Table 1 Author

Risk of bias assessment for studies on non-operative treatment for lumbar spinal stenosis with neurogenic claudication

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Table 1 Continued													
Author	A	в	v	٥	ш	ш	IJ	т	_	Г	¥	_	Total
Acupuncture													
Kim et al ⁵⁵	+	+	I	I	I	I	+	+	I	+	+	+	7†
Qin et al ⁵⁶	+	+	+	I	+	+	+	+	+	I	+	+	10*
Epidural injections													
Cuckler <i>et al</i> ³²	ذ	ć	+	+	+	+	+	+	+	\$	+	+	6
Fukusaki <i>et al</i> ³⁴	ć	\$	6	2	+	+	+	+	+	2	+	+	7
Zahaar ³³	ć	ċ	+	\$	+	+	+	+	+	1	ć	1	6
Brown ³⁵	+	I	+	I	ć	+	+	I	ć	2	I	+	5
Friedly <i>et al.</i> ³⁷ Friedly <i>et al</i> ⁶² . and Makris ⁶⁴	+	+	+	+	+	+	+	+	~	+	+	+	11 *
Song <i>et al</i> ³⁶	2	2	2	\$	2	+	+	I	2	+	+	+	5
Milburn <i>et al</i> ³⁸	ć	ć	+	1	+	1	+	I	ć	I	1	+	4
Hammerich <i>et al</i> ⁴⁰	+	+	I	I	+	I	+	2	2	I	+	+	6†
Sencan et al ⁴¹	+	\$	+	I	+	+	2	+	+	+	2	+	8‡
Wei et al ⁴²	+	+	+	I	1	+	I	+	\$	+	+	+	*00
Percutaneous epidural adhesiolysis	iolysis												
Karm et al ³⁹	+	2	+	I	+	I	+	+	2	I	I	+	1 9
Surgery versus physical therapy	ру												
Zucherman <i>et al,¹⁷</i> Zucherman <i>et al⁶⁰</i> and Hsu <i>et al</i> ⁶⁹	2	+	I	I	+	+		+	+	6	+	+	>6¶
Weinstein <i>et al</i> , ²¹ Weinstein <i>et al</i> ⁶⁸ , Abdu <i>et al</i> ⁶¹	+	+	I	1	+	+	+	+	~	~	1	+	>6 *** 1†
Amundsen <i>et al</i> ¹⁶	+	2	I	I	I	+	+	+	I	2	I	2	4
Malmivaara et a/ ¹⁹	+	+	I	I	+	+	+	+	+	2	\$	+	*0
Weinstein <i>et</i> $a^{/22}$, Weinstein <i>et</i> $a^{/99}$, Lurie <i>et</i> $a^{/58}$	+	+	I	I	+	I	+	+	2	د.	I	+	6††
Delitto et al ²⁵	+	+	I	1	+	ć	+	1	+	I	+	+	7††
A: Was the method of randomisation adequate? B: Was the treatment allocation concealed? C: Was the patient blinded to the intervention? F: Was the dop-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 reporting? I: Were the groups similar at baseline regarding the most important prognostic indicators? J: Were cointerventions avoided or similar? K: Was the compliance acceptable in all groups? A: Yes, -0.0, 2: Unclear. Jou 7: 2: Unclear. Jou 7: 2: Unclear. Jou 6: 2: Constitution of the outcome assessment similar in all groups? A: Yes, -1.0 wisk of the study free of suggestion of selective outcome reporting? I: Were the state including valid randomisation and treatment allocation techniques and no severe flaws. Jou 7: Unclear. Jou 7: 2: Unclear. Jou 7: 2: Order of the outcome assessment similar in all groups? A: Yes, Towisk of the subord of the outcome assessment similar in all groups? A: Yes, Towisk of the study free of suggestion of selective outcome assessment similar in all groups? A: Yes, Towisk of the study free of suggestion and treatment allocation techniques and no severe flaws. Jour side of study. To participants per treatment am. To study the outcome assessment similar in all groups? A: Yes, Towisk of the study free of suggestion of selective outcome assessment similar in all groups? A: Yes, Towisk of the study free of suggestion of selective outcome assessment similar in all groups? A: Yes, Towisk of the study free of suggestion of selective outcome assessment similar in all groups? A: Yes, Towisk of the study free of suggestion of selective outcome assessment similar in all groups? A: Yes, Towisk of the study free of study. To study free of study. To study free of study. To study free of study. To study free of of study. To study free of study. To st	tion adequate trate describs ng the most ir ms met, inclui mm. 0%, 1 year<2 00% at 4 year voer rates.	 B: Was the d and accept nportant prog ding valid rant j%; intention s. 	treatment allocat table? G: Were al inostic indicators domisation and ti to treat inconsist	on concealed? C: I randomised partic ? J: Were cointerve reatment allocation tent at 2-year follow	7 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 participants analysed in the group to which they were allocated? H: Are reports of the study free of suggestion of selective outcome reporting? I: W terventions avoided or similar? K: Was the compliance acceptable in all groups? L: Was the timing of the outcome assessment similar in all groups ation techniques and no severe flaws.	inded to the inter the group to whi similar? K: Was t to severe flaws.	vention? D: Was tich they were allc he compliance a	the care provider retered? H: Are ret care provider retered? H: Are ret care coeptable in all gr	blinded to the ir oorts of the stud oups ? L: Was th oups?	tervention? E: War free of suggestio e timing of the out	s the outcome as: n of selective outc come assessmen come assessmen	sessor blinded tc come reporting? t similar in all grc	the I: Were the ups? +: Yes,

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 cognitivebehavioural 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 lowquality 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 lowquality 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 lowquality evidence.

A meta-analysis⁸ ⁹ 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 trial¹⁶ 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.²⁴ Other studies demonstrated that inpatient physical therapy (ultrasound, heat and TENS) is more effective than home exercise plus oral diclofenac,²³ unweighted treadmill walking plus exercise is no better than cycling plus exercise²⁰ and manual therapy, 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.⁶³ A follow-up subgroup analysis⁶⁴ 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.

Å 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 lowquality 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 LSS^{71 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.⁷⁸⁷⁹ 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

BMJ Open: first published as 10.1136/bmjopen-2021-057724 on 19 January 2022. Downloaded from http://bmjopen.bmj.com/ on April 19, 2024 by guest. Protected by copyright. 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 trans-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

parency and reduce bias.

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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Nonoperative treatment for lumbar spinal stenosis - 22 July 2020 update

Database: Ovid MEDLINE: Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE® Daily and Ovid MEDLINE® <1946-Present> Search Strategy:

- 1 randomized controlled trial.pt. (509927)
- 2 controlled clinical trial.pt. (93770)
- 3 Pragmatic clinical trial.pt. (1444)
- 4 random*.ti,ab. (1145458)
- 5 placebo.ab,ti. (215288)
- 6 drug therapy.fs. (2221199)
- 7 trial.ab,ti. (599425)
- 8 groups.ab,ti. (2097678)
- 9 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 (5031369)
- 10 (animals not (humans and animals)).sh. (4686362)
- 11 9 not 10 (4375594)
- 12 exp Constriction, Pathologic/ (30449)
- 13 limit 12 to yr="1976 1982" (1906)
- 14 exp Lumbar Vertebrae/ (52505)
- 15 limit 14 to yr="1966 1982" (4472)
- 16 exp Spinal Canal/ (7519)
- 17 limit 16 to yr="1966 1982" (1172)
- 18 exp Spinal Diseases/ (123399)
- 19 limit 18 to yr="1966 1982" (18365)
- 20 exp Spinal Stenosis/ (6116)
- 21 spinal stenosis.ti,ab. (5088)
- 22 (lumbar adj5 stenosis).ti,ab. (4268)
- 23 (spin* adj5 stenosis).ti,ab. (6620)
- 24 neurogenic claudication.ti,ab. (633)
- 25 exp Spinal Osteophytosis/ (4018)
- 26 exp Spondylosis/ (7484)
- 27 (lumb* adj5 spondyl*).ti,ab. (2886)
- 28 exp Cauda Equina/ (3250)
- 29 lumbar radicular pain.ti,ab. (218)
- 30 13 or 15 or 17 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 29 (44520)
- 31 9 and 30 (6508)
- 32 limit 31 to ed=20190920-20200731 (275)
- 33 limit 31 to yr=2019-2020 (545)
- 34 32 or 33 (583)

Database: Embase Classic+Embase <1947 to 2020 July 21> Search Strategy:

- ------
- 1 Randomized Controlled Trial/ (613507)
- 2 exp Controlled clinical trial/ (800817)
- 3 Controlled Study/ (7533843)
- 4 Double Blind Procedure/ (176652)

- 5 Single Blind Procedure/ (39549)
- 6 crossover procedure/ (64054)
- 7 placebo/ (362923)
- 8 Randomization/ (87513)
- 9 random*.ti,ab. (1563918)
- 10 placebo?.ti,ab. (314621)
- 11 allocat*.ti,ab. (155448)
- 12 assign*.ti,ab. (400691)
- 13 blind*.ti,ab. (436413)
- 14 (cross-over or crossover).ti,ab. (107060)
- 15 (compare or compared or comparing or comparison or comparative).ti,ab. (6802913)
- 16 (controlled adj7 (study or design or trial)).ti,ab. (355549)
- 17 ((singl* or doubl* or trebl* or tripl*) adj7 (blind* or mask*)).ti,ab. (250201)
- 18 trial.ti,ab. (878032)
- 19 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18 (12682849)
- 20 exp animals/ or exp invertebrate/ or animal experiment/ or animal model/ or animal tissue/ or animal cell/ or nonhuman/ (29761121)
- 21 human/ or normal human/ or human cell/ (22533987)
- 22 20 and 21 (22470134)
- 23 20 not 22 (7290987)
- 24 19 not 23 (9386132)
- 25 exp vertebral canal stenosis/ (12543)
- 26 (spin* adj5 stenosis).ti,ab. (9011)
- 27 (lumbar adj5 stenosis).ti,ab. (5728)
- 28 (neurogenic adj2 claudication).ti,ab. (1047)
- 29 (Spin* adj2 Osteophytosis).ti,ab. (26)
- 30 exp cauda equina/ (4498)
- 31 lumbar radicular pain.ti,ab. (316)
- 32 (lumb* adj5 spondyl*).ti,ab. (4037)
- 33 exp spondylosis/ (9560)
- 34 spondylolisthesis/ (9419)
- 35 25 or 26 or 27 or 28 or 29 or 30 or 31 or 32 or 33 or 34 (36443)
- 36 24 and 35 (11296)
- 37 limit 36 to yr=2019-2020 (1405)
- 38 limit 36 to dd=20190920-20200731 (282)
- 39 37 or 38 (1426)

CENTRAL via CRS Web

- 1 MESH DESCRIPTOR Spinal Stenosis EXPLODE ALL AND CENTRAL: TARGET 423
- 2 (spin* NEAR5 stenosis) AND CENTRAL:TARGET 1189
- 3 lumb* NEAR5 stenosis AND CENTRAL:TARGET 871
- 4 neurogenic claudication AND CENTRAL:TARGET 168
- 5 MESH DESCRIPTOR Spinal Osteophytosis EXPLODE ALL AND CENTRAL:TARGET 86
- 6 MESH DESCRIPTOR Spondylosis EXPLODE ALL AND CENTRAL:TARGET 374
- 7 lumb* NEAR5 spondyl* AND CENTRAL:TARGET 400
- 8 MESH DESCRIPTOR Cauda Equina EXPLODE ALL AND CENTRAL:TARGET 15
- 9 lumbar radicular pain AND CENTRAL:TARGET 93

10 #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 AND CENTRAL:TARGET 1932

- 11 2019:YR AND CENTRAL:TARGET 105034
- 12 2020:YR AND CENTRAL:TARGET 30634
- 13 #11 OR #12 135668
- 14 #13 AND #10 209

CINAHL

S43 S41 OR S42 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 242 S40 AND EM 20190919-20200731 S42 Search modes - Boolean/Phrase Interface - EBSCOhost **Research Databases** Search Screen - Advanced Search Database - CINAHL Plus with Full Text 192 Limiters - Published Date: 20190901-20200731 S41 S40 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 161 S28 AND S39 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases \$40 Search Screen - Advanced Search Database - CINAHL Plus with Full Text 3,036 S29 OR S30 OR S31 OR S32 OR S33 OR S34 OR S35 OR S36 OR S37 OR S38 539 Search modes -Boolean/PhraseInterface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 6,262 Search modes - Boolean/Phrase Interface - EBSCOhost Research S38 lumb* W5 spondvl* Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 796 Search modes - Boolean/Phrase Interface - EBSCOhost Research S37 MH "Spondylolysis" Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 486 S36 MH "Spondylolisthesis" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 1,438 "lumbar radicular pain" Search modes - Boolean/Phrase Interface - EBSCOhost Research \$35 Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 125 S34 MH "Cauda Equina" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 368 S33 Search modes - Boolean/Phrase Interface - EBSCOhost Research MH "Spinal Osteophytosis" Databases Search Screen - Advanced Search

Database - CINAHL Plus with Full Text 310 S32 "neurogenic claudication" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 243 S31 lumb* W5 stenosis Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 1,768 S30 spin* W5 stenosis Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 3,656 S29 MH "Spinal Stenosis" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 2.741 S28 S26 NOT S27 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 2,433,818 S27 MH "Animals" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 87,894 S26 S7 OR S12 OR S19 OR S25 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 2,461,016 S25 S20 OR S21 OR S22 OR S23 OR S24 Search modes - Boolean/Phrase Interface - EBSCOhost **Research Databases** Search Screen - Advanced Search Database - CINAHL Plus with Full Text 1,686,740 S24 volunteer* Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 52,797 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases S23 prospectiv* Search Screen - Advanced Search Database - CINAHL Plus with Full Text 525,699 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases \$22 control* Search Screen - Advanced Search Database - CINAHL Plus with Full Text 1,275,002 followup stud* Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases S21 Search Screen - Advanced Search Database - CINAHL Plus with Full Text 203 S20 follow-up stud* Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 12,011 S13 OR S14 OR S15 OR S16 OR S17 OR S18 Search modes - Boolean/Phrase Interface -\$19 **EBSCOhost Research Databases**

Search Screen - Advanced Search

1,539,358 Database - CINAHL Plus with Full Text MH "Prospective Studies+" Search modes - Boolean/Phrase Interface - EBSCOhost Research S18 Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 444,171 MH "Evaluation Research+" Search modes - Boolean/Phrase Interface - EBSCOhost Research S17 Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 248,871 S16 MH "Comparative Studies" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 331,705 S15 latin square Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 248 S14 MH "Study Design+" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 1,351,924 S13 MH "Random Sample" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 34,389 Search modes - Boolean/Phrase Interface - EBSCOhost Research S12 S8 OR S9 OR S10 OR S11 Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 431,064 S11 random* Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 414,911 \$10 placebo* Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 66,332 S9 MH "Placebos" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 12,827 S8 MH "Placebo Effect" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 2,282 S7 S1 OR S2 OR S3 OR S4 OR S5 OR S6 Search modes - Boolean/Phrase Interface - EBSCOhost **Research Databases** Search Screen - Advanced Search Database - CINAHL Plus with Full Text 404,557 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases 56 triple-blind Search Screen - Advanced Search

Database - CINAHL Plus with Full Text 379 Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases S5 single blind Search Screen - Advanced Search Database - CINAHL Plus with Full Text 15,679 S4 double blind Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 58,644 S3 clinical W3 trial Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 250,481 "randomi?ed controlled trial*" Search modes - Boolean/Phrase Interface - EBSCOhost Research S2 Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 169,418 S1 MH "Clinical Trials+" Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Plus with Full Text 303,246

PEDro

Yield: 12 Abstract and title: stenosis AND Body part: lumbar spine, sacroiliac joint or pelvis AND Method: clinical trial Year: Since 2012

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S2	, Peer Review only, Publication Type:Controlled Clinica	l Trial	9	2020-0	7-22
10:21:48					
S3	, Peer Review only, Publication Type:Randomized Cont	rolled Tr	ial	288	2020-
07-22 10:22:09					
S4	All Fields:random* OR All Fields:placebo* OR All Fields:	sham, P	eer Revie	ew only	1029
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S5	All Fields:\"clinical trial\" OR All Fields:\"controlled tria	I∖", Peer	Review	only	481
2020-0	7-22 10:28:38				
S6	All Fields:versus OR All Fields:vs., Peer Review only	196	2020-0	7-22 10	28:57
S7	All Fields:double-blind OR All Fields:\"double-blind\", P	eer Revi	ew only	107	2020-
07-22 10:29:22					
S8	All Fields:single-blind OR All Fields:\"single-blind\", Pee	er Review	/ only	456	2020-
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Type:Controlled Clinical Trial OR, Peer Review only, Publication Type:Controlled Clinical Trial OR, Peer Review only, Publication Type:Randomized Controlled Trial OR All Fields:random* OR All Fields:placebo* OR All Fields:sham, Peer Review only OR All Fields:\"controlled trial\", Peer Review only OR All Fields:versus OR Al

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S10 Subject:\"spinal stenosis\" OR All Fields:\"spinal stenosis\", Peer Review only 83 2020-07-22 10:30:56

S11 Subject:\"Spinal Osteophytosis\" OR Subject:\"Spondylosis\" OR

Subject:\"Spondylolisthesis\", Peer Review only 83 2020-07-22 10:31:27

S12 Subject:\"Cauda equina\" OR All Fields:\"lumbar radicular pain\", Peer Review only

14 2020-07-22 10:32:34

S13 Subject:\"spinal stenosis\" OR All Fields:\"spinal stenosis\", Peer Review only OR Subject:\"Spinal Osteophytosis\" OR Subject:\"Spondylosis\" OR Subject:\"Spondylolisthesis\", Peer Review only OR Subject:\"Cauda equina\" OR All Fields:\"lumbar radicular pain\", Peer Review only

172 2020-07-22 10:32:47

\$14, Peer Review only, Publication Type:Clinical Trial OR , Peer Review only, PublicationType:Controlled Clinical Trial OR , Peer Review only, Publication Type:Randomized Controlled Trial OR AllFields:random* OR All Fields:placebo* OR All Fields:sham, Peer Review only OR All Fields:\"clinical trial\"OR All Fields:\"controlled trial\", Peer Review only OR All Fields:versus OR All Fields:vs., Peer Review onlyOR All Fields:double-blind OR All Fields:\"double-blind\", Peer Review only OR All Fields:vs., Peer Review onlyOR All Fields:\"single-blind OR All Fields:\"double-blind\", Peer Review only OR All Fields:\"spinal stenosis\" OR All Fields:\"spinalstenosis\", Peer Review only OR Subject:\"Spinal Osteophytosis\" OR Subject:\"Spondylolisthesis\", Peer Review only OR Subject:\"Cauda equina\" OR All Fields:\"lumbarradicular pain\", Peer Review only262020-07-22 10:32:57

S15 , Year: from 2019 to 2020, Peer Review only 325 2020-07-22 10:33:21

S16 , Peer Review only, Publication Type:Clinical Trial OR , Peer Review only, Publication Type:Controlled Clinical Trial OR , Peer Review only, Publication Type:Randomized Controlled Trial OR All Fields:random* OR All Fields:placebo* OR All Fields:sham, Peer Review only OR All Fields:\"clinical trial\" OR All Fields:\"controlled trial\", Peer Review only OR All Fields:versus OR All Fields:vs., Peer Review only OR All Fields:double-blind OR All Fields:\"double-blind\", Peer Review only OR All Fields:\"single-blind OR All Fields:\"double-blind\", Peer Review only OR All Fields:\"single-blind OR All Fields:\"spinal stenosis\" OR All Fields:\"spinal stenosis\" OR All Fields:\"spinal stenosis\" OR Subject:\"Spondylolisthesis\", Peer Review only OR Subject:\"Cauda equina\" OR All Fields:\"lumbar radicular pain\", Peer Review only AND, Year: from 2019 to 2020, Peer Review only

Supplemental Table 2. Non operative interventions for neurogenic claudication due to lumbar spinal stenosis: A summary of GRADE assessment and outcomes (60 comparisons)

						Walking	ability/pain/functio	n/quality of life mea	isures	GRADE
Studies	Risk of Bias	Consistency	Directness	Precision	Selective Reporting	Immediate up to 1w	Short-term >1w - 3m	Intermediate 3m – 1yr	Long term <u>>1</u> yr	
						Calcitonin				
				C	alcitonin ir	njection vs. placeb	o injection			
Eskola	High	No	Yes	No	Yes	v 1	= TWT	= TWT	= TWT	+000
1992		No	Yes	No			= VAS	= VAS	= VAS	+000
Porter 1983	High	No	Yes	No	Yes		? Distance walked	? Distance walked		+000
Porter	High	No	Yes	No	Yes		= Distance walked			+000
1988	Ũ	No	Yes	No			= VAS			+000
				Cal	lcitonin na	sal spray vs. place	bo injection			
Podichetty	High	No	Yes	No	Yes		= Distance walked			+000
2004		No	Yes	No			= Time walked			+000
		No	Yes	No			= SF-36			+000
		No	Yes	No			= VAS			+000
Tafazal	High	No	Yes	No	No		= Shuttle walk			+000
2007		No	Yes	No			= VAS leg			+000
		No	Yes	No			= VAS back			+000
		No	Yes	No			= ODI			+000
		No	Yes	No			= Global			+000
	-					cal therapy vs. par	racetamol plus phy	sical therapy		
Sahin	High	No	Yes	No	No		= Distance walked			+000
2009		No	Yes	No			= VAS			+000
		No	Yes	No			= RMDI			+000
						Oral Medication				
				0	ral prostag	glandin vs. Etodlad	c (NSAID)			
Matsudaira	Low	No	Yes	No	Yes		> Distance walked #			++00
2009		No	Yes	No			? SF-36			+000
		No	Yes	No			=LBP			++00
		No	Yes	No			> Leg pain			++00
		No	Yes	No			> Global #			++00
			Methyloo	cobalami	n (vit B12)	plus conservative	e care vs. conserva	tive care		
Waikakul 2000	High	No	Yes	No	No			> Distance walked #	> Distance walked #	+000

	Gaba	pentin pl	us physical	therapy,	corset &	NSAIDS vs. placel	bo plus physical th	nerapy, corset & N	SAIDS	
Yaksi 2007	High	No No No	Yes Yes Yes	No No No	No		= VAS	> Distance walked > VAS	> Distance walked # > VAS #	+000 +000
					Prega	balin vs. active pla	icebo			
Markman	High	No	Yes	No	No	= NPS rest/final				+000
2015		No	Yes	No		= Distance walked				+000
		No	Yes	No		= Recovery time				+000
		No	Yes	No		= Global				+000
		No	Yes	No		< RMDQ				+000
	<u> </u>		Gat	papentin	plus cons	ervative vs. conserv	vative plus botulir	lum		
Park	High	No	Yes	No	No		= NPS (Back/leg)			0000
2017	_	No	Yes	No			= ODI			0000
		No	Yes	No			= Global			0000
				0	xvmorph	one hydrochloride y	vs. placebo			
Markman	High	No	Yes	No	No	= NPS rest/final				0000
2015 - 2	8	No	Yes	No		= Distance walked				0000
		No	Yes	No		= Recovery Time				0000
		No	Yes	No		= ZCQ (s)				0000
		No	Yes	No		= ZCQ (f)				0000
		No	Yes	No		= Global				0000
						ne/acetaminophen	vs. placebo			
Markham	High	No	Yes	No	No	= NPS rest/final				0000
2015 - 2		No	Yes	No		= Distance walked				0000
		No	Yes	No		= Recovery Time				0000
		No	Yes	No		= ZCQ (s)				0000
		No	Yes	No		< ZCQ (f) #				0000
		No	Yes	No		= Global				0000
			Oxy	morphor	e hydrocl	nloride vs. propoxy	phene/acetaminor	ohen		
Markham	High	No	Yes	No	No	= NPS rest/final				0000
2015 - 2	-	No	Yes	No		= Distance walked				0000
		No	Yes	No		= Recovery Time				0000
		No	Yes	No		= ZCQ (s)				0000
		No	Yes	No		> ZCQ (f) #				0000
		No	Yes	No		= Global				0000
					Ora	l corticoid vs. place	ebo			

Rodrigues	High	No	Yes	No	No	= SF-36		0000
2014	mgn	No	Yes	No	110	= RMDO		0000
2011		No	Yes	No		$= 6 \min \text{ walk}$		0000
		No	Yes	No		<vas #<="" td=""><td></td><td>0000</td></vas>		0000
				Reha	bilitation	Therapy and Multimodal Car	e	
			E			und vs. exercise plus sham ultra		
Goren	low	No	Yes	No	No	= TWT		++00
2010		No	Yes	No		= VAS back		++00
		No	Yes	No		= VAS leg		++00
		No	Yes	No		= ODI		++00
				E	xercise plu	s ultrasound vs. no treatment		
Goren	Low	No	Yes	No	No	= TWT		++00
2010		No	Yes	No		= VAS back		++00
		No	Yes	No		> VAS leg #		++00
		No	Yes	No		> ODI		++00
						ham ultrasound vs. no treatment	- /	
Goren	Low	No	Yes	No	No	= TWT		++00
2010		No	Yes	No		= VAS back		++00
		No	Yes	No		> VAS leg #		++00
		No	Yes	No		> ODI #		++00
						s. home exercise program plus o		
Koc	High	No	Yes	No	Yes	= TWT	= TWT	+000
2009		No	Yes	No		= VAS	= VAS	+000
		No	Yes	No		= RMDI	= RMDI	+000
		No	Yes	No	1 .11 1	= NHP	= HNP	+000
	-	1				king plus exercise vs. cycling pl		
Pua	Low	No	Yes	No	No	= Distance wall	ked	++00
2007		No	Yes	No		= ODI		++00
		No	Yes	No		= RMDI		++00
		No No	Yes Yes	No No		= VAS = Global		++00 ++00
	M						11. 1 1	
	-	r				treadmill vs. flexion exercise, w	aiking and sham ultr	
Whitman	High	No	Yes	No	No	= TWT		+000
2006		No	Yes	No		> Global #		+000
		No	Yes	No		= ODI		+000
		No	Yes	No		= NPRS		+000
				Supe	ervised phy	vsical therapy vs home exercises		
				~	in the phy	in the py is meane thereises		

Minetama	High	No	Yes	No	No		> ZCQ (F) #			+000
2019	riigii	No	Yes	No	INO		>ZCQ (I') #			+000 $+000$
2019		No	Yes	No			> Distance walked #			+000
		No	Yes	No			> NPS (leg)			+000
		No	Yes	No			> SF-36 PF			+000
		No	Yes	No			> SF-36 BP			+000 +000
		No	Yes	No						+000 +000
				No			= Daily Steps			+000 +000
		No	Yes		marcal the area	py & exercise vs r	nadical care			+000
Schneider	L	N	V	Yes	No	py & exercise vs i	> ZCQ #	= ZCQ		+++0
	Low	No	Yes		No		= SPWT			+++0 +++0
2019		No	Yes	Yes				= SPWT		
		No	Yes	Yes	1.1 (· ·	= PA	= PA		+++0
			T		19	k exercise vs. com		1	1	
Schneider	Low	No	Yes	Yes	No		> ZCQ #	= ZCQ		+++0
2019		No	Yes	Yes			= SPWT	= SPWT		+++0
		No	Yes	Yes			= PA	= PA		+++0
					Communit	ty exercise vs. mee	lical care			
Schneider	Low	No	Yes	Yes	No		= ZCQ	= ZCQ		+++0
2019		No	Yes	Yes			= SPWT	= SPWT		+++0
		No	Yes	Yes			> PA	= PA		+++0
			Cor	nprehens	ive therap	y and exercise vs.	self-directed exerc	ise		
Ammendolia	Low	No	Yes	Yes	No	> SPWT #	> SPWT #	> SPWT #	> SPWT #	+++0
2018		No	Yes	Yes		> 30% SPWT	> 30% SPWT	> 30% SPWT	>30% SPWT	+++0
		No	Yes	Yes		> 50% SPWT	= 50% SPWT	= 50% SPWT	> 50% SPWT	+++0
		No	Yes	Yes		> ZCQ (s)	= ZCQ (s)	= ZCQ (s)	> ZCQ (f) #	++00
		No	Yes	Yes		= ZCQ (f)	= ZCQ (f)	= ZCQ (f)	> ZCQ (s) +	++00
		No	Yes	Yes		= ODI	= ODI	> ODI (walk)	ZCQ (f)	++00
		No	Yes	Yes		> NPS (back) #	= NPS (back)	= NPS (back)	= ODI	++00
		No		Yes		= NPS (leg)	= NPS (leg)	= NPS (leg)	= NPS (back)	++00
						= SF-36 BP	= SF-36 BP	= SF-36 BP	> SF-36 BP #	++00
						= SF-36 PF	> SF-36 PF #	= SF-36 PF	>SF-36 PF #	++00
				S	tandard ex	ercise vs. isokineti	c exercises			
Oğuz	High	No	Yes	No	Yes	= VAS	= VAS	= VAS		0000
2013	8	No	Yes	No		= ODI	= ODI	= ODI		0000
-		No	Yes	No		= TWT	=TWT	=TWT		0000
				S	Standard ex	kercise vs. unloade	d exercise		•	
Oğuz	High	No	Yes	No	Yes	<vas< td=""><td>< VAS</td><td>= VAS</td><td></td><td>0000</td></vas<>	< VAS	= VAS		0000
2013	8	No	Yes	No		< ODI	= ODI	= ODI		0000
		No	Yes	No		= TWT	= TWT	= TWT		0000
		110	105	INU		1 // 1	1 1 1	1 1 1 1		0000

				Iso	okinetic ex	ercises vs. unloaded	d exercises			
Oğuz	High	No	Yes	No	Yes	< VAS	= VAS	= VAS		0000
2013		No	Yes	No		< ODI	< ODI	= ODI		0000
		No	Yes	No		< TWT #	< TWT	= TWT		0000
	-	•		<u> </u>		nerapy exercise vs.				
Homayouni	High	No	Yes	No	Yes	> VAS #	= VAS			0000
2015		No	Yes	No		> Distance walked	= Distance walked			0000
			Pre-surgica	al exerci	se progran	vs. routine preope	rative hospital ma	nagement		
Marchand	High	No	Yes	No	Yes	> NPS (leg) #	= NPS (leg)	= NPS (leg)		0000
2019		No	Yes	No		> Duration walked #	= Duration walked	= Duration walked		0000
Gang-C	huk Tang	(herbal co	oncoction),	daily M	okuri Chu	na therapy, daily ac	upuncture, physic	ian consultation v	s. oral aceclofe	nac,
				ep	idural ster	oid injection, physic				
Kim	Low	No	Yes	No	Yes		= VAS (leg)	= VAS (leg)		+000
2019		No	Yes	No			= VAS (back)	> VAS (back) #		+000
		No	Yes	No			> OCS	= OCS		+000
		No	Yes	No			> Distance walked	> Distance walked		+000
Mo	okhuri Chu	una, acupt	incture, and	d physici	an consult	ation vs. oral acecle	ofenac, epidural s	teroid injection, pl	nysical therapy	
Kim	Low	No	Yes	No	Yes	>VAS (low back)#	= VAS (leg)	>VAS (leg) #		+000
2019 No Yes No $= VAS (back) + 000$										
		No	Yes	No			= OCS	= OCS		+000
		No	Yes	No			= Distance walked	= Distance walked		+000
					Sp	inal Manipulation	l			
				Ι	umbar spi	nal manipulation vs	s. waiting			
Passmore	High	No	Yes	No	No	= NPS (Back)				0000
2017	Ũ	No	Yes	No		= NPS (Leg)				0000
						Acupuncture				
				Ac	cupuncture	with usual care vs.	usual care			

Kim	High				No		6 weeks:			
2016	mgn	No	Yes	No	110		= ODI			0000
2010		No	Yes	No			= SF-36 BP			0000
		No	Yes	No			= SF-36 PF			0000
		No	Yes	No			= LBP			0000
		No	Yes	No			= Leg pain			0000
		No	Yes	No			= Distance walked			0000
							3 months:			
		No	Yes	No			= ODI			0000
		No	Yes	No			= SF-36 BP			0000
		No	Yes	No			= SF-36 PF			0000
		No	Yes	No			= LBP			0000
		No	Yes	No			= Leg pain			0000
		No	Yes	No			= Distance walked			0000
					Acupunc	ture vs. sham acup	uncture			
Qin	Low	No	Yes	No	No	> RMDQ	> RMDQ	> RMDQ		++00
2020		No	Yes	No		> NRS (back) #	> NRS (back) #	> NRS (back)		++00
		No	Yes	No		> NRS (leg) #	> NRS (leg) #	> NRS (leg) #		++00
		No	Yes	No		> SSS-S #	> SSS-S #	> SSS-S #		++00
		No	Yes	No		> SSS-F #	> SSS-F #	> SSS-F #		++00
		No	Yes	No		= SPWT	= SPWT	= SPWT		++00
					E	pidural Injection				
			Tra	nslamina	r epidural	steroid injections v	vs. placebo injectio	ons		
Cuckler	High	No	Yes	No	No	= Global			=global	+000
1985										
			Translan	ninar epio	lural stero	ids plus epidural bl	lock vs. placebo ir	jections		
Fukusaki	High	No	Yes	No	No	> Distance walked #	= Distance walked			+000
1988	_									
		T	ranslamina	ar epidura	al steroids	plus epidural block	x vs. epidural bloc	k injections		
Fukusaki	High	No	Yes	No	No	= Distance walked	= Distance walked			+000
1988										
				Т	ranslamin	ar epidural block v	s. placebo			
Fukusaki	High	No	Yes	No	No	> Distance walked #	= Distance walked			+000
1988										
		Intralamin	ar epidura	l steroid	plus epidu	ral block vs. home	exercise program	plus oral diclofer	nac	
Koc	High	No	Yes	No	Yes		= TWT	= TWT		+000
			3.7	3.1	3.7		> TIAC II	TILO		1000
2009		No	Yes	No	Yes		> VAS #	= VAS		+000

		No	Yes	No	Yes		> NHP	= HNP		+000
		Iı	ntralamina	epidura	l steroid pl	lus epidural block v	s. in-patient physi	cal therapy		
Koc	High	No	Yes	No	Yes		= TWT	= TWT		+000
2009	e	No	Yes	No	Yes		= VAS	= VAS		+000
		No	Yes	No	Yes		= RMDI	= RMDI		+000
		No	Yes	No	Yes		= NHP	= HNP		+000
				Cau	dal epidura	al steroids vs. place	bo injections			
Zahaar 1991	High	No	Yes	No	No	= Global			= Global	+000
			Ν	/ild lum	oar decom	pression vs. epidura	al steroid injection	l		
Brown	High	No	Yes	No	No		= VAS			0000
2012		No	Yes	No			= ODI			0000
		No	Yes	No			= ZCQ			0000
1		No	Yes	No			12 weeks:			
							= VAS			0000
		No	Yes	No			= ODI			0000
		No	Yes	No			= ZCQ			0000
				I	idocaine	vs. glucocorticoid–	lidocaine			
Eni - 11 2014	T			1	No	vs. glucocorticolu-	3 weeks:	12 weeks:	12 months:	
Friedly 2014,	Low	N	V	Yes	INO					
2017		No No	Yes	Y es Yes			< RMDQ	= RMDQ	= RMDQ	+++0
		INO	Yes	Y es			< NPS (leg) 6 weeks:	= NPS (leg) 6 months:	= NPS (leg)	+++0
		No	Yes	Yes			= RMDO	= RMDQ		+++0
		No	Yes	Yes						$^{+++0}_{+++0}$
		No	Yes	r es			= NPS (leg)	= NPS (leg)		+++0
							Makris 2016			
							3 weeks:			
Makris 2016	Low	No	Yes	No	Yes		< RMDQ using SIP			0000
101atrib 2010	2011	110	105	110	105		Weights			0000
		No	Yes	No	Yes		< RMDQ Patient-			0000
		110	105	110	105		Prioritized			0000
							(LESSER)			
							6 weeks:			
		No	Yes	No	Yes		< RMDQ using SIP			
		110	105	1,0	105		Weights			0000
		No	Yes	No	Yes		= RMDQ Patient-			0000
		110	105	110	105		Prioritized			0000
							(LESSER)			0000
			I				(LESSER)			

Song 2016	High	No			No					
	U	No		1	INO		1 month:			
			Yes	No			= VAS			0000
		No	Yes	No			= FRI			0000
							3 months:			
		No	Yes	No			= VAS			0000
		No	Yes	No			= FRI			0000
	Fluoros	scopically	guided lur	nbar ILE	SIS at the	level of maximal s	tenosis vs. two int	ervertebral levels	cephalad	
Milburn	High	1 2			No	1 week:	4 weeks:			
2014	0	No	Yes	No		> NPS (walking) #	> NPS (walking) #			0000
		No	Yes	No		> RMDQ #	> RMDQ			
		No	Yes	No			12 weeks:			0000
							= NPS (walking)			
		No	Yes	No			> RMDQ			0000
		No	Yes	No			, i i i i i i i i i i i i i i i i i i i			0000
			•	Epidura	al steroid inj	ection (ESI) Vs. ESI	& physiotherapy	•		•
Hammerich	High	No	Yes	No	No		= ODI	= ODI	= ODI	0000
2019	-	No	Yes	No			= NPS	> NPS #	> NPS #	0000
		No	Yes	No			> SF-36 ER #	= SF-36 ER	= SF-36 ER	0000
		No	Yes	No			> SF-36 EWB	= SF-36 EWB	= SF-36 EWB	0000
		No	Yes	No			> SF-36 GH	= SF-36 GH	= SF-36 GH	0000
			Ι	nterlami	har vs. tran	sforaminal epidura	al steroid injection			
Sencan 2020	High				Yes	= NPS	3 weeks:	3 months:		
	8	No	Yes	No			= NPS	> NPS		0000
		No	Yes	No			= ODI	= ODI		0000
		No	Yes	No			> BDS	> BDS		0000
		No	Yes	No			= Distance walked	> Distance walked #		0000
		N.	V	No						0000
		No No	Yes	No No						0000
			Yes							
		No	Yes	No						0000
		No	Yes	No						0000
				TNF alp	ha inhibit	or (Etanercept) vs.	steroid injection			
Wei 2020	Low	No	Yes	No		> VAS #	1, 3 months:	6 months:		++00
		No	Yes	No			> VAS #	> VAS #		++00
		No	Yes	No			> ODI #	> ODI #		++00
				TNF	alpha inh	ibitor (Etanercept)	vs. lidocaine			

BMJ	Open
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Wei 2020	Low	No	Yes	No		> VAS #	1, 3 months:	6 months:		++00	
		No	Yes	No			>VAS #	> VAS #		++00	
		No	Yes	No			> ODI #	> ODI #		++00	
	Steroid vs. lidocaine injection										
Wei 2020	Low	No	Yes	No		= VAS	1, 3 months:	6 months:		++00	
		No	Yes	No			= VAS	= VAS		++00	
		No	Yes	No			= ODI	= ODI		++00	
			•]	Percutane	ous Epidural Adh	esiolysis		•		
	Balloon-less catheter (Racz) vs. inflatable balloon catheter (ZiNeu)										
Karm 2018	High				No		1 month:	6 months:			
	Ū.	No	Yes	No			= NPS (back)	< NPS (back) #		0000	
		No	Yes	No			= NPS (leg)	< NPS (leg) #		0000	
		No	Yes	No			= ODI	< ODI		0000	
							3 months:				
		No	Yes	No			= NPS (back)			0000	
		No	Yes	No			= NPS (leg)			0000	
		No	Yes	No			= ODI			0000	
					Surger	y vs. Physical The	rapy				
				Intersp	inous spac	er (X_Stop) vs. nor	n operative care				
Zucherman	High	No	Yes	No	No		> ZCQ(S)#	> ZCQ(S)#	> ZCQ(S)#	+000	
2004, 2005,	_	No	Yes	No			> ZCQ(F)#	> ZCQ(F)#	> ZCQ(F)#	+000	
Hsu 2006							> SF-36 PF	> SF-36 PF	> SF-36 PF#	+000	
							> SF-36 BP	> SF-36 BP	> SF-36 BP#	+000	
							> SF-36 GH	> SF-36 GH	> SF-36 GH	+000	
							> SF-36 ER	> SF-36 ER	> SF-36 ER#	+000	
		La	minectomy	/ +/- fusi	on vs. non	operative care for	degenerative spon	dylolisthesis			

***	TT: 1	N		N	N					
Weinstein	High	No	Yes	No	No		= SF-36 BP, PF	= SF-36 BP, PF	2 years:	
2007, 2009		No	Yes	No			= ODI	= ODI	= SF-36 BP, PF	+000
Abdu 2018		No	Yes	No			= LBPBS	= LBPBS	= ODI	+000
		No	Yes	No			= LPBI	= LPBI	= LBPBS	+000
		No	Yes	No			= SBS	= SBS	= LPBI	+000
									= SBS	+000
		No	Yes	No					4 years:	
		No	Yes	No					= SF-36 BP, PF	+000
		No	Yes	No					= ODI	+000
		No	Yes	No					= LBPBS	+000
		No	Yes	No					= LPBI	+000
									= SBS	+000
		No	Yes	No					8 years:	
		No	Yes	No					= SF-36 BP, PF	+000
		No	Yes	No					= ODI	+000
		No	Yes	No					= LBPBS	+000
		No	Yes	No					= LPBI	+000
									= SBS	+000
				Lam	inectomy -	+/- fusion vs. non o	perative care			
Amundsen	High	No	Yes	No	No		?* Pain severity	?* Global	?* Pain severity	+000
2000	8	No	Yes	No			5		? Global	+000
Malmivaara	Low	No	Yes	No	No			= TWT	= TWT	++00
2007		No	Yes	No				= SW	= SW	++00
N=94		No	Yes	No				> VAS leg walk #	> VAS leg walk	++00
		No	Yes	No				> VAS LB walk #	#	++00
		No	Yes	No				> ODI	> VAS LB walk	++00
		110	100	110				021	#	
									> ODI	++00
Weinstein	High	No	Yes	No	No		= SF-36 BP	= SF-36 BP	2 years:	+000
2008, 2010,	mgn	No	Yes	No	110		= SF-36 PF	= SF-36 PF	> SF-36 BP **	+000
Lurie 2015		No	Yes	No			= LBPBS	= LBPBS	#	+000
Euric 2015		No	Yes	No			= LPBI	= LPBI	= SF-36 PF	+000
		No	Yes	No			= SBS	= SBS	= LBPBS	+000
		No	Yes	No			= ODI	= ODI	= LPBI	+000
		INU	105	INU			- ODI	- ODI	= SBS	+000
									= ODI	1000
									4 years:	+000
			1						4 years: =SF-36 BP **	+000 +000
			1							
			1						= SF-36 PF	+000
									= LBPBS	+000
L									= LPBI	+000

									= SBS 8 years: = SF-36 BP = SF-36 PF = ODI = Stenosis Index	+000 +000 +000 +000
	Laminectomy, facet resection, neuroforaminotomy vs. physical therapy									
Delitto	High				No				2 years:	
2015		No	Yes	No					= SF-36	+000
		No	Yes	No					= ODI	+000

> favours intervention (first comparison), < favours control (second comparison), = no difference between intervention and control groups, TWT= Treadmill Walking Test, VAS= Visual Analog Scale for Pain Intensity, RMDI= Roland-Morris Back Disability Index, NHP= Nottingham Health Profile, Global= Patient Perceived Improvement, SR= Selective Reporting, ODI= Oswestry Back Disability Index, ?= insufficient data, LBP= Low back Pain Severity Scale, Leg pain= Leg Pain Severity Scale, ? SF-36=No data on overall score, improvement in some subscales, NPRS= Numeric Pain Rating Scale, SF-36 BP= SF-36 Bodily Pain Subscale, SF-36- PF= SF-36 Physical Function Subscale, SF-36 ER= SF-36 emotional role subscale, SF-36 EWB= SF-36 emotional well-being subscale, SF-36 GH= SF-36 General health subscale, LBPBS= Low Back Pain Bothersome Scale, LPBI= Leg Pain Bothersome Index, SBS= Stenosis Bothersome Scale, SW= Subjective Walking, VAS leg= Visual Analog Scale for Leg Pain, VAS LB= Visual Analog Scale for Low Back Pain, VAS leg walking= Visual Analog Scale for Leg pain while walking, SIP= sickness index profile, BDS= Beck Depression Score, LESSER= Lumbar Epidural Steroid Injection for Spinal Stenosis Extended Research, PA= Physical Activity, FRI= Functional Rating Index, TWT= Total Walking Time, SSS= Spinal Stenosis Questionnaire, ?*= no between group statistical comparisons, **= SF-36 BP significantly better at 2 years but not 4 years.</p>

GRADE evidence; +00= Very low GRADE evidence, ++00= Low GRADE, +++0= Moderate GRADE evidence, ++++= High GRADE evidence # between group difference meeting the MCID. The MCID used were: ≥ 1.25 points for back pain and ≥ 1.5 points for leg pain on 0 to 100-point Visual Analogue Scale (VAS) and 0 to 10-point Numerical Rating Scale (NRS) for back pain (58), ≥ 5 points on 0- to 24-point Roland-Morris Disability Questionnaire (RMDQ) (59), ≥ 8 points for conservative treatment and ≥ 12 points for surgery on 0- to 100-points for Oswestry Disability Index (ODI) (60), ≥ 0.1 points for the functional component and 0.36 points for symptom component of the Zurich Claudication Questionnaire (ZCQ) (58), ≥ 0.38 points for combined symptoms and functional scores of the ZCQ (92), $\geq 30\%$ between-group difference for walking distance, global improvement and SF36 subscales (61).

Supplemental Table 1. Characteristics of included studies

Study	Participants and	Int	erventions		tcomes/Follow-	Results
	Settings			up		(Group 1 is reference group)
					tonin	
Eskola 1992	39 subjects with an average of 6 years of pain, average age of 56.6 years of age, 20 males and 19 females. Setting: Orthopaedic hospital in Finland.	1) 2)	100IU Calcitonin injection every other day for 4 weeks (n=20) Placebo treatment (Miacalcic Sandoz 100IU) every other day for 4 weeks (n=19)	5) Fo 4,	VAS Treadmill test Coping with ADLs Digitest Ergojump Blood tests blow-up: 1, 3, 6 and 12 onths	Between group WMD and 95% CI Pain (VAS) (mm): -0.050 (-0.053 to -0.047) Walking distance (meters): -18.5 (-240.37 to 203.37) Adverse events: The calcitonin injection group reported minor nausea and rash in 89% of the subjects.
Podichetty 2004	55 subjects with an average age of 68.5 years and an average of 36.2 weeks of the condition in the intervention group and 29.8 weeks in the placebo group, 33 males and 22 females. Setting: Spinal center in the United States	1) 2)	400 IU intranasal calcitonin daily for 6 weeks followed by open label 6- week extension (n=36) Placebo nasal spray daily for 6 weeks, followed by open label 6- week extension, during which all patients received 400IU calcitonin (n=19)	1) 2) 3) 4) 5) 6) 7) Fol we	VAS Walking capacity ODI Stenosis specific questionnaire Satisfaction with pain levels, functional status, and treatment received SF-36 Symptom diary	Between group MD, 95% CI, p values 12 weeks: Pain VAS (mm): 0.5 (-0.85 to 1.93): p=0.44, Walking time (seconds): 42.2 (-86.9 to 170.4): p=0.51 Walking distance (feet): 163.3 (-311.16 to637.84); p=0. 0.49 SF-36 MCS: -4.22 (-10.41 to1.97) ; p=0.18 SF-36 PCS: 0.43 (-3.73 to 4.59); p= 0.84
Donton	11 aubie eta uriti	1)	100 III salman salaitanin initi			Insufficient data marridad ta calculata maan difference in
Porter	41 subjects with	1)	100 IU salmon calcitonin injection	1)	Walking chart	Insufficient data provided to calculate mean difference in

1983	10 in a double blind RCT crossover, 37 males and 4 females with mean age of 55.4 years. Setting: Infirmary in	four times per week, sometimes with Maxalon for nausea (n=5) 2) Matching placebo (n=5) Only responders randomized	and ability to walk more than 1 mile 2) ODI Follow-up: 10 weeks	 walking distance or ODI among the 10 patients enrolled in RCT. Adverse events: The calcitonin injection group reported minor nausea and rash in 40% of the subjects.
Porter 1988	England 42 subjects, 35 male, 7 female, average age of 53.6 years in 20 subjects and 56.7 years in 22 subjects, median duration of back pain reported was 11 years for 19 subjects, and 14 years for 22 subjects. Median duration of claudication was 1.25 years for 20 subjects and 4.5 years for 22 subjects. Setting: Infirmary in England	 100 IU of salmon calcitonin injected subcutaneously 4 times per week for 8 weeks (n=20) 1 ml of saline injected 4 times per week for 8 weeks (n=22) 	 VAS Claudication threshold 3 level mobility assessment Analgesic requirements 3 level sleep disturbance Treatment Treatment success defined as 100% improvement in walking distance and able to walk 800 m. Follow-up: 4 and 8 weeks	Difference in median score from baseline between groups Pain score (VAS) (mm): 4 weeks: -9 8 weeks: -5.5 Walking distance until symptoms onset (meters): 4 weeks: -14 8 weeks: 42 Walking distance until pain prevents walking (meters): 4 weeks: -41 8 weeks: -99 No significant between group differences. No p values or 95% CI provided.
Sahin 2009	45 subjects 31 males and 14 females, average	 200 IU intranasal calcitonin daily for 8 weeks (n=23) 	 VAS Walking capacity 	Percent change between groups: 8 weeks: VAS at rest: 4.7%, p>0.05

and Rehabilitation Medicine Department in Turkey			
Tafazal40 subjects, 301)2007males, 10females, averageof 67 years in the2)	Placebo nasal spray NaCl for 4 weeks (n=20) 200 IU nasal salmon calcitonin for 4 weeks (n=20)	 VAS Shuttle walking test 4-point subjective outcome of overall assessment (excellent, good, fair, poor) ODI Modified Somatic Perception Questionnaire Modified Zung Depression Score 	4 weeks: Between group MD 95% CI ODI: -0.7 (1.7 to -3.5) LBOS: -3.0 (-0.6 to -4.7) VAS leg (mm): -10 (-4.0 to -13) VAS back (mm): -6.0 (-6 to -12) Shuttle walk distance (m): -13 (-7 to -35) 16 weeks: between group MD, p values ODI: 0.1, p=0.44; LBOS: 0.7, p=0.93; VAS leg (mm): -4, p=0.66; VAS back (mm): 16, p=0.03; Shuttle walking distance (m): -11, p=0.39

			Ora	l Me	dication	
Matsudaira 2009	79 subjects, 24 males and 24 females, with an average age of 69.6 years in the Limaprost group and 72.2 in the Etodolac group. Setting: Orthopaedic surgery in a medical faculty	1) 2)	Oral prostaglandin E1 derivative (15 g Limaprost) 3 times daily for 8 weeks (n=39) 400 mg of etodolac (NSAID) twice daily for 8 weeks (n=40)	1) 2) 3) 4) 5) 6) 7)	SF-36 Verbal pain rating scales Walking distance LBP severity Leg pain severity Leg numbness severity Treatment satisfaction	SF-36 subscales MD, p values 8 weeks: physical function: 9.4, p=0.01, role physical: 13.7, p=0.03, bodily pain: 15.5, p<0.01: General health: 6.6, p=0.08; vitality: 11.3, p=0.02; social functioning: 8.0, p=0.17; role emotional: 10.2, p=0.07; mental health: 12.2, p<0.01. Secondary outcomes not provided in a way that MD can be extracted: 8 weeks: low back pain: p=0.77; leg pain p=0.08; Leg numbness: p<0.01; walking distance p<0.01; patient subjective improvement p<0.01; patient satisfaction p<0.01 all in favor of limaprost
	in Japan				llow-up: 8 eks	Adverse events: 5% of subjects in both groups reported gastrointestinal upset.
Waikakul 2000	152 subjects, 68 males and 84 females with an average age of 66.8 years. 44 of the subjects had symptoms for less than one month, 98 had symptoms for more than one month. Setting: Hospital in Thailand	1) 1) 2)	Conservative treatment consisting of education, activity modification, exercise and physical therapy. NSAIDs, muscle relaxants, and analgesics as necessary. Vitamin B1, B6, and B12 3 times per day (n=82) Conservative treatment plus Methlcobalin ESAI, 1.5mg per day in 3 divided doses after meals for 6 months (n=70)	1) 2) 3) Fo	Presence of pain on spinal motion Claudication distance Medication intake (NSAIDs, muscle relaxants, and steroids) llow-up: every mth for two	gastometry of the systemWalking distancePercent able to walk > 1000 meters6 mo: 71.3% vs. 88.6%, $p < 0.05$ 12 mo: 81.3% vs. 97.1%, $p < 0.05$ 18mo: 83.8% vs. 97.1% $p < 0.05$ Adverse events: There were no reported adverse effects in subjects in methylocabalin group
Yaksi 2007	55 subjects, 22 males, 33 females, average age of 50.8 years. Setting: Hospital	1)	900 mg of gabapentin per day increased weekly by 300 mg to a maximum of 2400 mg (n=28) Placebo (n=27)		VAS – low back and leg pain during movement Walking distance	Between group difference, p values Pain (VAS) (mm) no raw data 3 rd mo 3.4 vs. 1.9, p =0.039 4 th mo 4.1 vs.2.0, p =0.006 Walking Ability, no raw data

	department of physical medicine and rehabilitation in Turkey	Both groups received physical therapy exercises, a lumbosacral corset with steel bracing and NSAID treatments	 3) Presence or absence of motor and/or sensory deficits Follow-up: 15 days, 1, 2, 3, 4 months 	Grp 1: longer walking distance at end of 2^{nd} mo (p < 0.05), 3^{rd} mo (p <0.05) and 4^{th} mo (p <0.005) Adverse events: some subjects randomized to the gabapentin group (no data specified) experienced mild to moderate drowsiness and/or dizziness.
Markman 2015	29 participants, 20 males, 9 females, Eligible subjects were older than 50 years (mean 70.1 years) with at least one level of radiographically confirmed lumbar spinal stenosis and symptoms of neurogenic claudication for at least 3 months. Setting: Hospital in Rochester, New York	 Pregabalin group (n=14) Active placebo (Diphenhydramine) (n=15) Cross over study after 7 day wash out period. Pregabalin was started at 75 mg PO twice daily or diphenhydramine, 6.25 mg) and increased on day 4 to 150 mg PO twice daily (12.5 mg diphenhydramine) for 7 days. Pregabalin was decreased to 75 mg PO twice daily (6.25 mg diphenhydramine) on day 11 for 3 days of tapering. 	 NRS - time to first moderate pain symptom during a 15- minute treadmill test (Tfirst) (NRS - greater than 4) Follow-up: day 10 of intervention period 	Between group MD, 95% CI, p values Treadmill testing pain at rest (NRS) 0.29 (0.41 to 0.98): p=0.40 Treadmill testing final pain (NRS) 0.25 (-0.44 to 0.94): p=0.46 Treadmill testing distance walked (m) -24.06 (-75.63 to 27.52): p=0.35 Treadmill testing recovery time (min) -0.79 (-1.86 to 0.28): p=0.14 Treadmill testing patient global assessment of pain -0.08 (-0.45 to 0.29): p=0.67 Treadmill testing RMDQ 1.50 (0.38 to 2.62): p=0.01 Adverse events: Complications were reported in 64% of subjects in group 1, the most common being dizziness, compared to 35% in group 2.
Park 2017	45 subjects, 21 in GPN Group (17 female, 4 males, mean age 66.1± 10.5), and 24 in BTX group (15 female and 9 males, mean age	 Conservative treatments plus gabapentin (group GPN): Gabapentin 300 to 1200mg/d - titrated to patient characteristics, comorbidities, and reported side effects (n=21) Conservative treatments plus BTX 	 NRS - back/leg pain intensity Cramp frequency (no./wk) Cramp severity (0-4 	No statistically significant difference between groups and lack of reporting of quantitative data Adverse events: Five patients (20.8%) in group 2 reported mild to moderate pain at injection sites for a few days.

	66.2±8.2) Setting: Outpatient department for interventional pain management in Korea	 injection (group BTX): The BTX (botulinum toxin type A [Nabota]) dose was 100U in 5mL of 0.9% saline injected into the gastrocnemius medialis and lateralis. (n=24) Conservative treatments: education, exercise, analgesic medication, injection therapy including epidural injections, and physical therapy 	 criteria) 6) Insomnia severity – (ISI 0-28) 7) ODI 8) Patient global impression of change Follow-up: 2 weeks, 1 and 3 months. 	
Markman 2015 - 2	24 participants, 12 males and 12 females, (mean age 72 years) LSS by imaging with symptoms of neurogenic claudication Setting: Translational Pain Research Center at a University in Rochester, New York	 Oxymorphone hydrochloride (Opana IR, 5 mg) (n=8) Propoxyphene/acetaminophen (Darvocet, 100 mg/650 mg) (n=8) Placebo: 3 separate visits (random order with at least 3 day washout periods) (n=8) 	 NRS (at rest) NRS (final pain rating) AUC 4) 4) Distance walked (m) Recovery time (min) ZCQ Patient global assessment of pain RMDQ ODI Follow-up: Study was prematurely terminated 	Between group MD, 95% CI, p values Treadmill testing pain at rest (NRS) Grp 1 vs Grp 3: -0.24 (-0.72 to 0.65): $p-0.89$ Grp 2 vs Grp 3: -0.27 (-0.95 to 0.41): $p=0.32$ Grp 1 vs Grp 2: 0.23 (-0.45 to 0.92): $p=0.40$ Treadmill testing final pain (NRS) Grp 1 vs Grp 3: 0.2 (-0.74 to 1.14): $p=0.60$ Grp 2 vs Grp 3: 0.53 (-0.40 to 1.46): $p=0.16$ Grp 1 vs Grp 2: -0.33 (-1.26 to 0.61): $p=0.39$ Treadmill testing distance walked (m) Grp 1 vs Grp 3: -12.41 (-63.01 to 38.20): $p=0.54$ Grp 1 vs Grp 3: -23.41 (-73.60 to 26.79): $p=0.25$ Grp 1 vs Grp 3: -0.03 (-0.19 to 0.13): $p=0.61$ Grp 2 vs Grp 3: 0.01 (-0.15 to 0.17): $p=0.85$ Grp 1 vs Grp 3: 0.04 (-0.20 to 0.11): $p=0.47$ Grp 2 vs Grp 3: 0.04 (-0.16 to 0.09): $p=0.47$ Grp 2 vs Grp 3: 0.11 (-0.01 to 0.23): $p=0.03$ Grp 1 vs Grp 3: -0.03 (-0.52 to 0.47): $p=0.90$ Grp 1 vs Grp 3: -0.03 (-0.52 to 0.47): $p=0.90$ Grp 1 vs Grp 3: -0.03 (-0.52 to 0.47): $p=0.90$ Grp 1 vs Grp 3: -0.03 (-0.52 to 0.47): $p=0.90$ Grp 1 vs Grp 3: -0.03 (-0.52 to 0.47): $p=0.90$ Grp 1 vs Grp 3: -0.03 (-0.52 to 0.47): $p=0.52$

						Grp 1 vs Grp 2: -0.15 (-0.64 to 0.34): p=0.44
						The study was prematurely terminated because of the removal of propoxyphene/acetaminophen from the US market.
Rodrigues 2014	61 patients with lumbar canal stenosis (50–75 years; canal area < 100 mm ² at L3/L4, L4/L5, and/or L5/S1on MRI; and claudication within 100 m). 31 in the corticoid group (mean age 58.23 (6.38), and 30 in the placebo group (mean age 58.33 (6.19)) Setting: Hospital in São Paulo, Brazil	1)	mg/kg of oral corticoids daily, with a dose reduction of one-third per week for 3 weeks (n=31) Control group was administered placebo for the same period (n=30)	and	llow-up: 3, 6 1 12 weeks	Between group comparison VAS (6 weeks) Corticoid vs Placebo: 1.53 p=0.02 (in favour of placebo)
	· · · · · · · · · · · · · · · · · · ·		Rehabilitation The			
Goren 2010	45 subjects, 13 males, 32 females, average ages in groups of 57.4, 49.13, and 53.06. 7 subjects with pain duration of 3-6 months, 7 with pain duration of	2)	Stretching and strengthening exercises for lumbar, abdominal, leg muscles as well as low intensity cycling exercises were given as therapeutic exercises. Ultrasound was applied with 1mHz, 1.5W/cm2 intensity, in continuous mode on the back muscle for 10 minutes (n=17) Same as group 1 with Ultrasound on	2) 3) 4)	VAS (out of 10) Treadmill test at 3 km/h for maximum of 15 minutes or 750m. ODI Analgesic consumption	 Pain (VAS) (mm) within group MD 3 weeks: Grp 1: -2.2 for back pain; -1.47 for leg pain Grp 2: -1.94 for back pain; -2.47 for leg pain Grp 3: 0.40 for back pain; 0.54 for leg pain Between groups differences Leg pain: Grp 1> Grp 3 (p<0.01), Grp 2> Grp 3 (p<0.01) Walking Ability (within group MD)
	6-12 months, and	2)	off- mode (n=17)		Physiatrist	3 weeks: Grp 1: 94.30 seconds

	31 with pain		assessment	Grp 2: 114.94 seconds
	duration of	3) No exercise-no treatment (n=16)		Grp 3: -66.10 seconds
	greater than 12		Follow-up: End of	No significant change between groups
	months.		3-week treatment	
			period only	Disability (ODI) (within group MD)
	Setting:		period only	3 weeks:
	Rehabilitation			Grp 1: -3.94
	center in Turkey			Grp 2: -7.8
				Grp 3: -3.6
				GIP 5. 5.0
				ODI between groups differences
				Grp 1> Grp 3 (p<0.05), Grp 2> Grp 3 (p<0.05)
Koc	29 subjects, 21	1) Conservative inpatient physical	1) VAS	No raw data provided.
2009	male, 8 female,	therapy program 5 days a week for 2	2) Treadmill	No significant between group differences for all outcomes and
	average ages of	weeks. PT included applications of	walk test	follow-ups except:
	62.6, 61.1, and	ultrasound 1.5 W/cm2 for 10min, hot	3) Nottingham	
	53.1 years in the	pack for 20min, and TENS for 20min	Health Profile	Pain (VAS)
	three groups,	to the lumbar region $(n=13)$	4) RMDI	2 weeks: Grp 2 less pain than Grp 3 $p=0.008$
	average pain		5) Functional	
	duration of 5.7	2) Lumbar epidural steroid injections,	testing	Disability (RMDI)
	years, 5.0 years,	10 ml of solution containing 60mg of	including	2 weeks: Grp 2 less disability than Grp 3 $p=0.007$
	and 5.7 years in	triamcinolon acetonide (1.5 mL), 15	finger to floor	
	the three groups.	mg of 0.5% bupivacain hydrochloride	distance, sit-	
	and an or groups.	(3 mL), and 5.5 mL of physiologic	to-stand, and a	Quality of Life (Nottingham Health Profile) (no data
	Setting: Medical	saline (0.9%NaCl) was injected in	weight	provided)
	school	3.5minutes. (n=10)	carrying test	Grp 2 had significantly higher improvement than Grp 3 at 2
	department of		earlying test	weeks in mobility subgroup scores.
	physical	3) Control group (n=10)	Follow-up: 2	weeks in meening subgroup secres.
	medicine and		weeks, 1, 3 and 6	Adverse events: 1 subject reported angina pectoralis and 1
	rehabilitation in	All patients included were trained to	months	reported gastric complaints (group not specified).
	Turkey	pursue a home-based therapeutic exercise	monus	reperieu guerne comprante (group net specificu).
	1 01110 9	program performed twice daily for a		
		period of 6 months, and oral diclofenac		
		sodium 75mg was administered to all		
		patients twice daily for 2 weeks		
Pua	68 subjects, 35	1) Unweighted treadmill training:	1) VAS for pain	Pain (VAS) (mm) MD and 95% CI
2007	males, 33	Weeks 1 and 2, participants walked	over past	6 weeks: 2 (-5 to 10)
2007	males, 55	Teers I and 2, participants walked	over past	0 weeks. 2 (5 to 10)

	females, average age of 58 years, 12 week median pain duration Setting: Hospital in Singapore	2)	with a relatively pain-free gait which translated to 30–40% of body weight. In weeks 3 to 6, participants were encouraged to walk at a moderate intensity. The duration of each treadmill session was limited by participant tolerance or to a maximum of 30 minutes. 2x per week for 6 weeks = 12 sessions (n=33) Cycling on upright bicycle: During weeks 1 and 2, participants cycled at their comfortable pace at 50 to 60 rpm. Participants were instructed to assume a flexed posture. In weeks 3 to 6, participants were encouraged to exercise at a moderate intensity and the duration of each cycling session was limited by participant tolerance or to a maximum of 30 minutes. 2x per week for 6 weeks for 12 sessions (n=35)		perceived benefit on a 6- point scale ODI RMDI	Disability (ODI), OR, 95% CI 6 weeks: OR 1.10 (0.41 to 2.98) Patient perceived benefit, OR, 95% CI 6 weeks: OR 0.50 (0.17 to 1.48) Walking ability (≥800 m), OR, 95% CI 6 weeks: OR 1.14 (0.44 to 2.94) Adverse events: 1 subject in treadmill group reported increase in pain.
Whitman 2006	58 subjects, 31 males, 27 female, 29 (group 1) with an average age of 70 years, 29 (group 2) with an average age of 68.9, median low back pain duration of 108 months in Group 1's 29 subjects and 60 months in Group 2's 29	1)	Flexion Exercise and Walking Group: 45-60 minutes twice per week for 6 weeks. Lumbar flexion exercises along with self-pace treadmill walking program, and sub- therapeutic ultrasound. The duration of each treadmill session was based on that patient's tolerance on that specific day and could extend up to 45 minutes. (n=29) Manual Therapy, Exercise and Walking Group: 45-60 minutes twice per week for 6 weeks - Manual	1) 2) 3) 4) 5) 6)	Medication consumption	Patient Global Assessment (somewhat better or greater) 6 weeks: 41% vs. 79% p<0.01

	subjects, lower extremity median pain duration of 48 months in Group 1's 29 subjects and 24 months in Group 2's 29 subjects. Setting: University in the United States	physical therapy (thrust and non thrust) to the thoracic and lumbar spine, pelvis, and lower extremities and specific exercises at discretion based on the underlying impairments. Patients received specific exercises to address impairments in mobility, strength, and/or coordination. Exercises were performed in the clinic and as part of a home exercise program. Patients also underwent a bodyweight supported treadmill ambulation program using a cable and trunk harness system to unload a specific amount of weight from the patient while the patient walks as comfortably as possible on a treadmill (n=29).	Stenosis Scale 7) Additional use of health care resources Follow-up: 6 weeks, 1 year, long term mail survey (averaging 29 months)	Long term: 1.8 (0.6 to 3.0) vs. 2.0 (0.7 to 3.4) Between group MD not statistically significant at any follow-up period Walking Ability (improvement in meters) within group MD, 95% CI 6 weeks: 176.5 (-9.5 to 362.4) vs. 339.7 (218.4 to 461) 1 year: 130.4 (-55.3 to 316.2) vs. 209.8 (67.5 to 352.1) Between group improvement not statistically significant at any follow-up Disability (ODI) within group MD 6 weeks: 6.55 (1.87 to 11.23) vs. 10.48 (6.5 to 14.4) 1 year: 5.03 (1.71 to 8.35) vs. 7.14 (1.5 to 12.8) Between group differences not statistically significant at any follow-up
Minetama 2019	86 patients, 39 men and 47 women, average age 72.7 years 43 patients (20 men and 23 women, average age 72.3 years to the PT group 43 patients (19 men and 24 women, average age 73.2 years) to the HE group. Duration symptoms 20 months	 Physical therapy + home exercise program (n=43) Home exercise (HE) program alone (n=43) Supervised physical therapy twice a week for 6 weeks, including manual therapy, individually tailored stretching and strengthening exercises, cycling, and body weight-supported treadmill walking. The manual therapy included manipulation, stretching, and massaging of the thoracic and lumbar spine, pelvis, and lower extremities. The individually tailored muscle exercises included those for the trunk (eg, abdominal planks, side bridge, and/or back extension) and lower 	 ZCQ Satisfaction SPWT (m) NRS JOABPEQ- acquired points SF-36 HADS PCS PASS-20 TSK-11 Daily steps Follow-up: 6 weeks 	Between group MD, 95% CI ZCQ - Symptom severity -0.4 (-0.6 to -0.2): statistically significant ZCQ - Physical function -0.4 (-0.6 to -0.2): statistically significant SPWT (m) 455.9 (308.5 to 603.2): statistically significant NRS - Leg pain -1.4 (-2.5 to -0.3): statistically significant SF-36 - Physical functioning 9.2 (2.1 to 16.3): statistically significant SF-36 - Bodily pain 10.4 (3.3 to 17.5): statistically significant Daily steps 723.4 (199.1 to 1,283.5): statistically significant

	Setting: Spine care center at a university hospital in Japan	extremities (eg, unloading hip and/or knee exercise with ankle weight and/or standing squats). The typical dosage for strengthening exercises was a total of 2 to 3 sets with 10 repetitions, each of 6- second contraction. The typical duration of stretching was three repetitions of 30 seconds.		
		All patients in both groups were asked to take a daily walk that did not exacerbate their lower extremity symptoms using a pedometer and walking diary and to perform a HE program consisting of lumbar flexion exercises including three 30-second bouts of both single and double knee-to-chest exercises, ten 6- second bouts of trunk raises and bridging in the supine position, and a 4-point kneeling exercise at least twice daily.		
Schneider 2019	259 subjects, 122 males and 137 women with an average age of 72.4, 68 patients had symptoms for less than 6 months, 191 had symptoms for greater than 6 months Setting: Outpatient research clinic in	 Medical care (MC) (n=88) Group exercise (GE) (n=84) Manual therapy + exercise (MTE) (n=87) Medical Care: 3 visits to a physical medicine physician over 6 weeks. Primarily prescription of oral medications in any combination of nonnarcotic analgesics, anticonvulsants, antidepressants. Optional referral for epidural steroid injections if inadequate pain relief by oral 	 SSS SPWT Physical Activity Follow-up: 2 and 6 months 	Between group MD, 95% CI SSS (2 months) GE vs MC: 0.4 (-1.3 to 2.1) MTE vs MC: -2.0 (-3.6 to -0.4) MTE vs GE: -2.4 (-4.1 to -0.8) SPWT (2 months) GE vs MC: 79.9 (-74.5 to 234.5) MTE vs MC: 122.9 (-25.7 to 271.6) MTE vs GE: 43.0 (-111.8 to 197.9) Physical activity (2 months) GE vs MC: 28.7 (2.7 to 54.7) MTE vs MC: 20.4 (-4.5 to 45.3) MTE vs GE: -8.3 (-34.5 to 17.6) SSS (6 months) GE vs MC: -0.5 (-2.3 to 1.3)
	Pittsburgh	medication, severe neurogenic claudication, and/or patient preference.		MTE vs MC: -1.1 (-2.8 to 0.6) MTE vs GE: -0.6 (-2.4 to 1.2)

		Physician rendered general guide and on gentle stretching and advice to stay active. Group Exercise: Supervised exercise classes at 2 local senior community centers. 2x 45-min classes/week, 6 weeks. Taught by senior fitness instructors. Participants self-select level of exercise based on fitness level (easy to medium)			SPWT (6 months) GE vs MC: 86.5 (-75.7 to 248.8) MTE vs MC: 73.8 (-84.1 to 231.7) MTE vs GE: -12.7 (-175.6 to 150.1) Physical activity (6 months) GE vs MC: 21.3 (-6.9 to 49.4) MTE vs MC: -2.9 (-30.1 to 24.3) MTE vs GE: -24.2 (-52.5 to 4.0) Adverse events: There were no reported serious adverse events in any group. There was a significantly greater rate of transient
		Manual Therapy + Exercise: 2x 45minute sessions per week, 6 weeks by either 2 chiropractors or 2 physiotherapists. Sessions included 3 interventions: 1. Warm-up procedure on stationary bicycle 2. Manual therapy procedures (lumbar distraction, hip, lumbar/sacroiliac joint and neural mobilizations 3. Individualized instruction in spinal stabilization exercises and home stretching Practitioner determined what muscles required stretch/strengthening and appropriate exercises added to program.			joint soreness associated with group 3 (49%) compared with group 2 (31%) and group 1 (6%).
Ammendolia 2018	104 patients, 45 males and 59 females, 48 in comprehensive group and 51 in self-directed group, with an average age of 69.4	 Comprehensive (n=48) Self-directed (n=51) Comprehensive: Chiropractor providing 2x/week of 15-20-minute treatment sessions over a 6-week period followed by a single (booster) session, 4 weeks later. 	1) 2) 3)	SPWT Distance Clinical Significance - 30% improvement in SPWT no. (%) Clinical	Between group MD, 95% CI, p values SPWT 8 wks: 345.4 (150.0 to 540.7): p=0.00 3 mo: 304.1 (77.9 to 530.3): p=0.01 6 mo: 421.0 (181.4 to 660.6): p=0.00 12 mo: 473.2 (203.9 to 742.4): p=0.00 30% improvement in SPWT 8 wks: 24 (6-40): p=0.01 3 mo: 21 (4-38): p=0.02

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	(comprehensive)	Education: Self-management strategies	Significance -	6 mo: 19 (2-35): p=0.02
	and 71.7 (self-	via cognitive behavioral approach.	50%	12 mo: 22 (4-39): p=0.02
	directed)	Body repositioning (pelvic tilt) when	improvement	50% improvement in SPWT
	neurogenic	standing and walking.	in SPWT no.	8 wks: 26 (8-42): p=0.01
	claudication >3	Exercises:	(%)	3 mo: 19 (-1.0 to 36): p=0.06
	months, imaging-	Standardized set of exercises	4) ZCQ-S	6 mo: 17 (-2 to 35): p=0.09
	confirmed canal	demonstrated gradually over 6 weeks and	5) ZCQ-F	12 mo: 24 (5-40): p=0.01
	narrowing, walk	was a part of structured home exercise	6) ZCQ-S +	ZCQS
	>20m and not	program. Cycling, muscle stretching,	ZCQ-F	8 wks: -0.19 (-0.37 to -0.02): p=0.03
	surgical	strengthening, conditioning for back and	7) ODI	3 mo: -0.15 (-0.37 to 0.08): p=0.19
	candidates in	lower extremity fitness and to facilitate	8) ODI walk	6 mo: -0.02 (-0.22 to 0.19): p=0.87
	next 12 months	lumbar flexion	9) NRS Back	12 mo: -0.22 (-0.47 to 0.02): p=0.07
		Manual therapy: Spinal manipulation;	10) NRS Leg	ZCQF
	Setting:	joint, soft tissue and neural mobilization;		8 wks: -0.02 (-0.22 to 0.17): p=0.81
	Academic	lumbar flexion-distraction; and manual	Follow-up: 8	3 mo: -0.18 (-0.39 to 0.03): p=0.09
	hospital	muscle stretching applied each visit.	weeks, 3, 6, and 12	6 mo: -0.11 (-0.33 to 0.11): p=0.34
	outpatient clinic	Participants received an instructional	months	12 mo: -0.27 (-0.49 to 0.04): p=0.02
	in Toronto	video and workbook and pedometer.		ZCQS+ZCQF
				8 wks: -0.24 (-0.56 to 0.07): p=0.13
		Self-directed: Instructional Video,		3 mo: -0.36 (-0.75 to 0.03): p=0.07
		workbook, pedometer and a single 15-to		6 mo: -0.23 (-0.58 to 0.12): p=0.20
		30-minute training session with an		12 mo: -0.48 (-0.90 to -0.06): p=0.03
		experienced independent licensed		ODI
		chiropractor, independent of the		8 wks: -0.02 (-0.07 to 0.02): p=0.30
		comprehensive program,		3 mo: -0.04 (-0.09 to 0.01): p=0.13
		Training session: Describe 6-week		6 mo: -0.02 (-0.07 to 0.02): p=0.34
		program, review workbook, explain		12 mo: -0.03 (-0.08 to 0.02): p=0.30
		pedometer use and recording of weekly		ODI Walk
		walking steps.		8 wks: -0.2 (-0.6 to 0.1): p=0.14
		Video and workbook: Educational		3 mo: -0.4 (-0.9 to 0.03): p=0.07
		information and the same exercise		6 mo: -0.9 (-1.3 to -0.4): p<0.001
		instruction and self-management		12 mo: -0.2 (-0.7 to 0.2): p=0.32
		strategies received by the comprehensive		NRS Back
		group		8 wks: -1.4 (-2.2 to -0.5): p=0.002
				3 mo:-0.6 (-1.4 to 0.3): p=0.23
				6 mo: -0.7 (-1.7 to 0.3): p=0.16
				12 mo: -0.4 (-1.3 to 0.4): p=0.32

				3 mo: 9.2 (1.1 to 17.3): p=0.027 6 mo: 5.8 (-2.1 to 13.6): p=0.15 12 mo: 8.2 (0.2 to 16.2): p=0.045 Adverse events: At 12 months, 0 participants out of 43 in group 1 and 2 out of 46 participants in group 2 experienced adverse events that were mostly attributed to a temporary increase in low back and/or leg pain.
in grou an ave 57.1 yo in grou an ave 55.8 yo and gro an ave 57.4 yo LSS sy	bup 1 with erage age of years old, 30 bup 2 with erage age of years old roup 3 with erage age of years old, symptoms, wing by	Standard exercise group (n=30) sokinetic exercise program (n=30) Jnloading exercise group (n=60) roups physician-guided (5x/week for eks) then at-home (3x/week) dard Exercise: 15 sessions of S, hot packs with home exercise tection. netic exercise: 20 minutes/day, 5 pns/week for a total of 15 sessions	 VAS ODI Beck Depression Inventory Follow-up: 4, 12 and 24 weeks	Between group MD, p value VAS After treatment: Grp 1 vs Grp 2:0.37, p>0.05 Grp 1 vs Grp 3: 1.36, p<0.05 Grp 2 vs Grp 3: 0.99, p<0.05 4^{th} week: Grp 1 vs Grp 2: 1.43, p>0.05 Grp 1 vs Grp 3: 1.17, p<0.05 Grp 2 vs Grp 3: -0.26, p>0.05 12 th week: Grp 1 vs Grp 2: 0.93, p>0.05 Grp 1 vs Grp 3: 0.71, p>0.05 Grp 2 vs Grp 3: -0.22, p>0.05 24 th week:

department of	rates of 60°/sec, 120°/sec, 180°/sec with	Grp 1 vs Grp 3: 0.46, p>0.05
physical	70° of body movement (50° flexion to	Grp 2 vs Grp 3: -0.62, p>0.05
medicine and	20° extension)	ODI
rehabilitation in	Each session had 3 sets, each set had 5	After treatment:
Turkey	repetitions at described velocity, with 20s	Grp 1 vs Grp 2: -0.8, p>0.05
Turkey	rest between each set.	Grp 1 vs Grp 2: -0.0, p> 0.05 Grp 1 vs Grp 3: 1.8, p< 0.05
	Test between each set.	Grp 2 vs Grp 3: 2.6, p<0.05
	Unloaded exercise: 5 sessions of	4 th week:
	unloading exercise per week, for a total	Grp 1 vs Grp 2: 1.5, p>0.05
		Grp 1 vs Grp 2: 1.5, p>0.05 Grp 1 vs Grp 3: 2.6, p>0.05
	of 15 sessions with a physician. Walking	
	with unloading exercise devise: session $1.5 = 450$ / he downight accession (15 =	Grp 2 vs Grp 3: 1.1, p<0.05 12 th week:
	1-5 = 45% body weight, session $6-15 = 20%$	
	30% body weight. Treadmill walking at	Grp 1 vs Grp 2: 1, p>0.05
	1.2 km/hr for 20 minutes, or until pain	Grp 1 vs Grp 3: 1.3, p>0.05
	due to neurogenic claudication was felt.	Grp 2 vs Grp 3: 0.3, p>0.05
	Subjects advised to follow exercise	24^{th} week:
	program s at home at least 3x/week after	Grp 1 vs Grp 2: 0.4, p>0.05
	discharge.	Grp 1 vs Grp 3: 0.5, p>0.05
		Grp 2 vs Grp 3: 0.1, p>0.05
		Total Gait Duration
		After treatment:
		Grp 1 vs Grp 2: 64.6, p>0.05
		Grp 1 vs Grp 3: -50.5, p>0.05
		Grp 2 vs Grp 3: -115.1, P<0.05
		4 th week:
		Grp 1 vs Grp 2: 45.9, p>0.05
		Grp 1 vs Grp 3: -18.4, p>0.05
		Grp 2 vs Grp 3: -64.3, p<0.05
		12 th week:
		Grp 1 vs Grp 2: 52.23 p>0.05
		Grp 1 vs Grp 3: -0.67 p>0.05
		Grp 2 vs Grp 3: -52.9 p>0.05
		24 th week:
		Grp 1 vs Grp 2: 35.2, p>0.05
		Grp 1 vs Grp 3: 1.9, p>0.05
		Grp 2 vs Grp 3: -33.3, p>0.05

Homayouni	47 subjects, 23	1)	Treatment in therapeutic pools with	1) VAS	All between group comparisons
2015	male, 24 female,		water temperature of 29-30 degrees	2) Walking	Walking ability
	24 in group one,		Celsius. Every aquatic session	ability	Grp 1 > Grp 2: p=0.02
	mean age 55.56,		started with warm up and ended with		VAS
	12 male, 12		cool down, with duration of 10-15	Follow-up:	Grp 1 > Grp 2 p=0.001
	female, 23 in		min for each of them. Participants	Immediately after	
	group two, mean		should have attended aquatic	therapy, 3 months	
	age 55.68, 11		physical therapy sessions every other		
	male, 12 female		day for a total duration of 24		
			sessions. Each session included		
	Setting:		ambulation, side walking, chain		
	University-based		walking, forward walking with		
	pain clinics in		kickboard, stretching of each muscle		
	Iran		group including adductors,		
			abductors, flexors and extensors of		
			the hip, knee flexors and ankle		
			plantar flexors and dorsiflexors.		
			Other interventions were mini-squat,		
			pelvic curl, pelvic tilt, and knee to		
			chest, double knee lift, and deep-		
			water exercise. (n=25)		
		2)	Passive modalities by physical		
			therapists including continuous mode		
			ultrasound (US) 1.5W/ cm2 for 10		
			min and hot pack and trans-electrical		
			nerve stimulation (TENS) for 20 min		
			to the lumbar region. Also, the		
			therapists instructed the patients in		
			this group to perform trunk muscle		
			endurance, William's and stretching		
		1	exercises. The patients were treated		
			using these passive modalities and		
			were given exercises under		
			supervision of physiotherapists for		
			10 sessions. They were instructed to		
			perform the learned exercises 30 min		

			a day at home in the following weeks until the end of the eighth week. (n=25)		
Marchand 2019	40 participants, 17 females and 23 males, 20 in the intervention group with an average age of 66.7 years old and 20 in the control group with an average age of 71.5 years old, with history and diagnostic imaging of LSS Setting: Regional hospital in Quebec	1) 2)	Exercise 3x week / 6 weeks prior to surgery (n=20) Regular hospital preoperative management with back posture education (n=20)	NRS (Pain Intensity) ROM (Active) Muscle strength (N-m) Walking capacity (seconds)	Between group MDNRS (leg)Preoperative: -2.1, $p<0.05$ Postoperative: 1.1, $p>0.05$ 3 months: 1.1, $p>0.05$ 6 months: 0.3, $p>0.05$ ROM (active)Preoperative: 5, $p<0.05$ Postoperative: -6, $p>0.05$ Muscle StrengthPreoperative: 5.1, $p>0.05$ Walking DurationPreoperative: -14.5, $p>0.05$
Kim 2019	34 subjects, mean age 64 (5.3), women 24 (66.7) Setting: Hospital in Seoul, South Korea	1)	MT1 group: 110 g of Gang-Chuk Tang was administered 3 times a day (Gang-Chuk Tang is an herbal concoction consisting of Eucommiae Cortex, Achyranthis Radix, Rhizoma Cibotii, Sorbus commixta, G. thunbergii, Saposhnikovia Radix, and Acanthopanacis Cortex in equal portions) Daily Mokhuri Chuna therapy (relaxation and mobilization of lumbar joint and back muscle) Daily acupuncture treatment on LI4, ST36, LV3, BL22, BL23, BL24, BL25, and Ashi points. Consultation on precautions related to daily	VAS for leg pain VAS for low back pain Oxford Claudication Scoring Walking distance	All between group comparisons VAS leg pain (post treatment) MT2 (28.82±27.46) vs CMT (51.82±25.34) groups: P=0.04 VAS leg pain (6 months) MT1 (48.91±23.08) vs CMT (72.27±16.72) groups: P=0.01 MT2 (42.36±21.29) vs CMT groups: P=0.003 VAS low back pain (6 months): MT2 (30.00±13.48) vs CMT (60.82±18.62) groups: P=0.001 Oxford Claudication Scoring (3 months) MT1 (18.75±6.52) vs CMT (25.82±6.24) groups: p=0.02 Walking distance (3 months) MT1 vs CMT: p=0.03 Walking distance (6 months) MT1 vs CMT: p=0.01

		2)	activity and stepwise walking training for the entire 4 weeks of therapy. (n=12) MT2 group: Mokhuri Chuna, acupuncture, and physician consultation were offered in the same manner and dosage as the MT1 group with the exception that all herbal medications were withheld. (n=11) CMT group: Oral analgesic therapy (aceclofenac 100 mg twice daily and eperisione hydrochloride 50 mg three times daily for 28 days) and three interlaminar epidural steroid injections (5 mg of dexamethasone per injection) at the level of the affected spinal region over a 4-week period were administered. Physiotherapy including heating pad, and transcutaneous electrical nerve simulator, and deep tissue heating therapy five times per week for 4 weeks. (n=11)		The primary outcome of this pilot study was safety as measured by the type and incidence of adverse events (AEs).
				Manipulation	
Passmore 2017	14 patients with degenerative LSS (n=14); Swiss Spinal Stenosis score of M=63.2, standard deviation [SD] = 15.9) (mean age 59.0 (10.6)), 7 in the SM group (4	1)	Spinal manipulation group: received bilateral high-velocity; low- amplitude spinal manipulation directed toward the lumbar region (by a licensed chiropractor with more than 10 years of clinical experience) (n=7) Non Intervention Group: Waited 5 minutes if they were assigned to the	 Movement time NPS (Back) NPS (leg) ROM Follow-up: Immediately after intervention	There was no significant difference between groups for all outcomes. 1. Grp 1 vs. Grp 2, p=0.739 2. Grp 1 vs. Grp 2, p> 0.05 3. Grp 1 vs. Grp 2, p> 0.05 4. Grp 1 vs. Grp 2, p> 0.05

female, 3 male) (mean age 59.1 (9.3)), 7 in the NI group (3 female, 4 male) (mean age 58.9 (12.6)) Setting: rehabilitation hospital in Winnipeg, Manitoba	no intervention group (n=7)		
l		cupuncture	1
Kim 201650 participants mean age of 62.0±9.8 years, acupuncture (n=26), age 65.0±8.7, male / female 12/14, control (n=24), age 58.9±10.2, male / female 10/14. Mean duration of symptoms 33mSetting: Hospital in Yangsan, South Korea	 Acupuncture: 269 acupuncture sessions were administered during the study. 81% (n=21) of patients received at least 10 acupuncture sessions. Electrical acupuncture was applied at least once and bilaterally at back shu points (BL23, BL24, BL25 or BL26) or Jiaji points at L2– L5 spinal levels. Other frequently used points were BL57, BL60, GB39, GB34 and tender points located in the lower extremities (n=26) Control: In total, 255 physical therapy sessions were provided to patients in the control group at their request. 92% (n=22) of patients received at least 10 physical therapy sessions (median 11, range 1–13). (n=24) 	 ODI SF-36 bodily pain SF-36 physical function LBP bothersomene ss LBP intensity Leg pain bothersomene ss Leg pain intensity Self-reported pain-free walking distance (m) Follow-up: 6 weeks, 3 months 	Between group MD, 95% CI ODI 6 wk: -2.2 (-7.0 to 2.6) 3 mo: -2.5 (-8.9 to 3.8) SF-36 BP 6 wk: -8.6 (-18.6 to 1.3) 3 mo: 3.2 (-8.3 to 14.7) SF-36 PF 6 wk: 0.1 (-7.6 to 7.9) 3 mo: 1.3 (-8.3 to 10.9) LBP bothersomeness 6 wk: -0.6 (-11.4 to 10.1) 3 mo: -7.4 (-19.6 to 4.8) LBP intensity 6 wk: -5.1 (-15.5 to 5.3) 3 mo: -13.5 (-26.2 to -0.7) Leg pain bothersomeness 6 wk: -7.4 (-18.4 to 3.7) 3 mo: -9.2 (-21.6 to 3.2) Leg pain intensity 6 wk: -11.5 (-0.9 to -22.0) 3 mo: -12.6 (-24.6 to -0.6)

						None statistically significant
Qin 2020	80 participants	1)	Acupuncture: Applied by	1)	RMDQ	RMDQ
	assigned with 70		acupuncturists with 5 years of	2)	NRS back	4 wk: -3.6 (-5.2 to -1.9): p<0.001
	completing the 8-		Chinese medical university program	3)	NRS Leg	8 wk: -2.6 (-3.7 to -1.4): p<0.001
	week treatment		and at least 2 year of clinical	4)	SSS	3 mo: -2.3 (-3.9 to -0.7): p=0.005
	course (38 in acu		experience. Sterile disposable steel		Symptoms	6 mo: -1.8 (-3.6 to -0.3): p=0.086
	group and 32 in		needles (Hwato Acupuncture,		subscale	NRS Back
	sham acu group).		Suzhou, China; 0.30 £ 40 mm/0.30 £	5)	SSS physical	4 wk: -1.7 (-2.4 to -0.9): p<0.001
	Mean age of		75 mm) were inserted through		function	8 wk: -2.3 (-3.0 to -1.5): p<0.001
	61.5±7.9 years		adhesive pads. Participants		subscale	3 mo: -1.7 (-2.6 to -0.8): p<0.001
	with 34 males		underwent 3 treatments weekly over	6)	SSS	6 mo: -1.2 (-2.1 to -0.3): p=0.007
	and 46 females.		8 weeks, and each session persisted		satisfaction	NRS Leg
	Duration of		for 30 minutes. To maintain "De qi,"		subscale	4 wk: -2.0 (-2.6 to -1.3): p<0.001
	symptoms <3mo		a sensation of numbness and	7)	Self-paced	8 wk: -2.9 (-2.6 to -1.3): p<0.001
	=14 (17.5%), 3-		soreness, acupuncture manipulation		walk test	3 mo: -2.4 (-3.3 to -1.4): p<0.001
	12 mo = 1(1.3%),		(twirling, lifting, and thrusting on			6 mo: -2.1 (-3.0 to -1.2): p<0.001
	1 to 5 $y = 24$		needles) was performed every 10		llow-up: 4	SSS Symptoms Subscale
	(30%), >5 y =41		minutes during the treatment.		eks, 8 weeks	4 wk: -0.6 (-0.8 to -0.4): p<0.001
	(51.3%)				d of treatment),	8 wk: -0.9 (-1.2 to -0.6): p<0.001
		2)	Sham acupuncture: Chosen		nonths, 6	3 mo: -0.9 (-1.2 to -0.6): p<0.001
	Setting:		acupoints, treatment duration, and	mo	nths	6 mo: -1.0 (-1.3 to 0.6): p<0.001
	2 Clinical Sites -		frequency of sessions were the same			SSS Physical Function Subscale
	Department of		as in the acupuncture group.			4 wk: -0.5 (-0.8 to -0.3): p<0.001
	Acupuncture and		Participants in the sham cohort were			8 wk: -0.8 (-1.1 to -0.5): p<0.001
	Neurology,		treated using a pragmatic placebo			3 mo: -0.7 (-1.0 to -0.4): p<0.001
	Guang'anmen		needle on the same acupoints, which			6 mo: -0.7 (-1.1 to -0.4): p<0.001
	Hospital		is similar to the Streitberger needle			Self-Paced Walk Test
	Department of		design (Supplementary Materials).			4 wk: p=0.648
	Acupuncture and		Acupuncturists pretended to			8 wk: p=0.29
	Neurology,		manipulate the needle every 10			3 mo: p=030
	Beijing Fengtai		minutes, but "De qi" was not sought.			6 mo: p=0.133
	Hospital of					
	Integrated					Adverse events: 3 participants in group 1 reported pain after
	Traditional and					needle insertion and 1 had a hematoma. 3 participants in group 2
	Western					reported back pain and 2 reported fatigue. All adverse events
	Medicine.					were reported as mild or moderate, and none required medical
						intervention.

		Epidu	ural injections	
Cuckler 1985	73 subjects in total, 37 with spinal stenosis, 36 with acute herniated nucleus pulposus, 37 males, 36 female, average age of 48.5 years in the experimental group and 49.5 years in the placebo group. Experimental group average 36.6 months in symptom duration, placebo group averaged 29.4 months. Setting: Orthopaedic surgery department in the	 Steroid group: 2ml of sterile water containing 80mg of methylprednisolone acetate combined with 5ml of 1% procaine was injected into the epidural space in the region between the 3rd and 4th lumbar vertebrae with the patient in the lateral decubitus position lying on the side of the painful limb (n=42), 20 with stenosis). Placebo group: 2ml of saline combined with 5ml of 1% procaine was injected into the epidural space in the region between the 3rd and 4th lumbar vertebrae with the patient in the lateral decubitus position lying on the side of the painful limb. (n=31, 17 with stenosis) All patients were advised to take mild analgesics (aspirin or acetaminophen) during the post-injection period. Second injection given if less than 50% improvement after 24 hours - considered 	 Subjective percentage of improvement with 75% required to be considered a treatment improvement, if less than 50% after 24 hours was considered a treatment failure Re-injection rates Surgery rates Follow-up: 24 hours, every 3 months up to 30 months, averaging 20.2 months in the steroid group and 21.5 months in the 	Patient Global Assessment (improved by at least 75%) 24 hours: 33% (steroid) vs. 21% (saline) p>0.05 Long term: 33% (saline) vs. 14% (saline) p>0.05
Fukusaki 1988	United States 53 subjects, 38 males and 15 female. Group 1 averaged 70 years of age and 79 days of symptoms on average, group 2 averaged 69 years of age and	 treatment failure Epidural injection with 8 ml of saline, repeated twice in the first week (n=16) Epidural injection with 8 ml of 1% mepivacaine, repeated twice in the first week. (n=18) Epidural injection with a mixture of 8 ml of 1% mepivacaine and 40 mg 	 control group. 1) Walking distance which was graded according to distance (excellent, good, or poor) Follow-up: 1 week, 1 month, 3 	Walking distancePercent excellent effect = mean of > 100m in walking distance1 week: 12.5 % (saline) vs. 55% (block) vs. 63.2% (block +steroid); block or block + steroid > saline, p< 0.05;

	an average of 82 days of symptoms, group 3 averaged 72 years of age and 94 days of symptoms on average Setting: Anaesthesia department in Japan	of methylpredniso in the first week. (months	all follow-up periods, p>0.05 Adverse events: no reported complications
Zahaar 1991	30 subjects, 37 male and 26 female. Steroid group averaged 46.5 years of age and 36.6 months of symptoms, control group averaged 49 years of age and 29.4 months of symptoms Setting : Medical facility in Egypt	 (n=18) 2) Control: 2x2ml of injected into epidu completed with sto 30ml. (n=12) 	etate suspension, 4% Volume erile saline to 30ml carbocaine, 4% ral space. Volume erile saline to	 Subjective percentage of improvement where 75% or more was deemed successful and surgery after injection was considered a failure. Follow-up: 24 hours, then every three months up to 36 mo averaging 20.2 mo in the steroid group and 21.5 mo control group. 	Patient Global Assessment (improved by at least 75%) 24 hours: 55% (steroid injection) vs. 50% (control) p> 0.05 Up to 36 mo: 38% (steroid injection) group vs. 33.3% (control) p>0.05 Failures (%) (required surgery) Up to 36 mo: 61% (steroid injection) vs. 66.6% (control) p>0.05
Friedly 2014, 2017 Makris 2016	400 patients, 221 females and 179 males, 200 in the lidocaine group	 Lidocaine + gluco of 0.25-1% lidocai 3 mL triamcinolor betamethasone (6- 	ne followed by 1- ne (60-120mg),	1) RMDQ 2) NRS (Leg Pain)	Between group MD, 95% CI, p values RMDQ 3 weeks: -1.8 (-2.8 to -0.9): p<0.001 6 weeks: -1.0 (-2.1 to 0.1): p=0.07

	with an average age of 68.1 years old and 200 gluocorticoid- lidocaine group with an average age of 68 years old, LSS by CT or MRI. 26% patients symptoms greater than 5 years. Setting: 16 medical centers across the United States	 dexamethasone (8-10mg) or methylprednisone (60-120mg)) (n=200) 2) Lidocaine group (0.25-1% lidocaine alone) (n=200) Physician option for intralaminar and/or transformaminal techniques 	 Follow-up: 3, 6, and 12 weeks, 6 and 12 months Makris 2016 subgroup 1) RMDQ using SIP Weights 2) RMDQ patient-prioritized (LESSER) Follow-up: 3 and 6 weeks 	12 wk: 0.1 (-1.0 to 1.3): p=0.84 6 mo -0.00 (-1.1 to 1.1): p=0.99 12 mo: -0.4 (-1.6 to 0.9): p=0.55 NRS (Leg pain) 3 weeks: -0.6 (-1.2 to -0.1): p=0.02 6 weeks: -0. (=0.8 to 0.4): p=0.48 12 wk: 0.1 (-0.5 to 0.7): p=0.70 6 mo: -0.2 (-0.8 to 0.4): p=0.47 12 mo: 0.1 (-0.5 to 0.7): P=0.75 Subgroup Analysis RMDQ using SIP weight 3 wks: -1.9 (-2.9 to -0.7): p<0.001 6 wks: -1.1 (-2.2 to -0.1): p=0.04 RMDQ patient prioritized (LESSER) 3 wks: -1.8 (-2.8 to -0.8): p<0.001 6 wks: -1.0 (-2.0 to 0.1): p=0.08 Adverse events: A total 21.5% of patients in group 1 and 15.5% in group 2 reported one or more adverse events (p=0.08) that included headaches, fever, infection, dizziness, cardiovascular/lung problems, leg swelling and dural puncture.
Song 2016	29 subjects, 14 males and 15 women with an average age of 58.3 and 61.7 between groups, history of intermittent claudication and lower limb radicular pain or paresthesia	 Lidocaine spinal injection, 40 mg triamcinolone mixed with 10 mL 0.5% lidocaine was used under the guide of fluoroscopy (n=15) Saline spinal injection using same volume (n=14) 	 VAS FRI Follow-up: 1 and 3 months 	No significant difference between groups. VAS 1-month p= 0.696, 3 months p= 0.891 FRI 1-month p=0.983, 3 months p=0.743

	Setting: Rehabilitation clinic in Korea			
Milburn 2014	57 patients met inclusion criteria, agreed to participate, and were enrolled. 20 patients were male; 37 were female. Mean patient age was 65.3 years (range, 32-88 years). Average duration of symptomatology (pain and/or disability) was 42 months. The mean degree of canal narrowing at the most stenotic level was 6.1 mm (range, 2.5-9.1 mm). The most common maximally stenotic intervertebral level was L4-L5	 Fluoroscopically guided lumbar ILESI performed either at: 1) The level of maximal stenosis (n=30) 2) Two intervertebral levels cephalad, corresponding to a less stenotic level (n=27) Injection was performed with a 20-gauge Tuohy needle using a loss of resistance technique. The injectate consisted of 2 mL of 40 mg/mL methylprednisolone (Pfizer), 2 mL of bupivacaine 0.25% (Hospira), and 2 mL of normal saline for a total injectate volume of 6 mL. 	 NRS - Pain with Ambulation RMDQ Follow-up: 1, 4 and 12 weeks 	All between group comparisons NRS (pain with ambulation) 1 wk: Grp 1 lower pain compared to Grp 2, p=0.045 4 wk: Grp 1 lower pain compared to Grp 2, p=0.049 12 wk: Grp 1 lower compared to Grp 2, p=0.001 4 wk: Grp 1 lower compared to Grp 2, p=0.009 12 wk: Grp 1 lower compared to Grp 2, p=0.003

	(n ¹ /42) followed by L3-L4 (n ¹ /411) and L5-S1 (n ¹ /44). Setting: Clinic in New Orleans, Louisiana			
Brown 2012	38 patients, 21 males and 17 females, 21 in mild group with an average age of 74.2 years and 17 in ESI group with an average age of 78.7 years, symptomatic LSS patients with painful lower limb neurogenic claudication, able to walk at least 10 feet unaided, (ODI) score > 20 Setting: Pain management clinic in Florida	 Epidural steroid (80 mg triamcinolone acetate) (n=17) Mild lumbar decompression (n=21) 	 VAS ODI ZCQ Patient Satisfaction (0-10) Follow-up: 6 and 12 weeks	VAS 6 and 12 weeks P=0.54 ODI p=0.86 ZCQ p>0.05 Patient satisfaction p>0.05
Hammerich 2019	54 patients total, age 67.2 ± 9.7 , 27 male, 27 female, 31 in ESI group, 23 in ESI plus PT. Mean duration of	 ESI (n=31) ESI + PT (n=23) ESI: 1.5 mL of steroid at each site injected with maximal involvement using transforaminal approach. 	 ODI NRS current SF-36 emotional role SF-36 emotional role well-being 	Between group MD, 95% CI, p values ODI 10 wks: -1.08 (-8.10 to 5.94) p=0.80 6 mo: -4.70 (-11.72 to 2.32) p=0.27 12 mo: -2.72 (-9.74 to 4.30) p=0.52 NRS 10 wks: -1.68 (-3.08 to -0.29) p=0.07

	symptoms 14 m Setting: Clinics in Colorado, Texas, South Carolina and New Hampshire	PT: 8-10 sessions PT manual therapy and exercise. Walking program and/or stationary bike, stretching and strengthening exercises.	 5) SF-36 general health perception Follow-up: 10 weeks, 6 and 12 months 	6 mo: -1.99 (-3.38 to -0.60) p=0.04 12 mo:-2.44 (-3.80 to -1.08) p=0.00 SF-36 Emotional role 10 wks: -28.53 (-49.05 to -8.01) p=0.03 6 mo: -11.25 (-31.77 to 9.27) p=0.39 12 mo: -10.67 (-31.19 to 9.85) p-0.41 SF-36 Emotional well-being 10 wks: -11.26 (-19.52 to -2.99) p=0.02 6 mo: 2.69 (-5.57 to 10.95) p=0.59 12 mo: -5.76 (-14.02 to 2.50) p=0.24 SF-36 General Health Perception 10 wks: -8.99 (-17.20 to -0.78) p=0.05 6 mo: -5.56 (-13.77 to 2.65) p=0.23 12 mo: -5.10 (-13.31 to 3.11) p=0.27
Sencan 2020	67 patients. The median age 62.5 years with 18 males and 49 females. Median duration of symptoms was 29 and 24 months in the ILESI and bilateral TFESI groups, respectively Setting: University department Pain Medicine, Istanbul Turkey	 Interlaminar: ILESI, fluoroscopy guided with 1 to 2 mL contrast dye with mixture of 80 mg methylprednisolone acetate, 2 mL saline solution, and 2 mL (0.5%) bupivacaine solution Transforaminal: TFESI, fluoroscopy guided with 1 to 2 mL contrast dye with mixture of 80 mg methylprednisolone acetate, 2 mL saline solution, and 2 mL (0.5%) bupivacaine solution 	 NPS ODI Beck depression scale Walk distance Follow-up: after treatment, 3 weeks and 3 months 	Between Group Median Differences (data not provided), p values NPS after treatment: p=0.14 3 wks: p=0.28 3 mo: p=0.047 ODI 3 wks: p=0.93 3 mo: p=0.65 Beck Depression Scale 3 wks: p=0.048 3 mo: p=0.03 Walking Distance 3 wks: p=0.23 3 mo: p= 0.048
Wei 2020	90 patients. Mean age about 65 years, 45 females, 45	 Epidural injection with 2.0mL of lidocaine and 10 mg of TNF-a inhibitor (etanercept) on the affected spinal nerves. 	 VAS (leg) ODI Follow-up: after 	Between Group Mean Differences (data not provided), p values Grp 1 vs Grp 2 VAS

	males, mean duration of symptoms about 2.8 months Setting: University Hospital Jiangsu China	2) 3)	Epidural administration with 2mL of lidocaine mixed with 2mL of steroid (diprospan) Epidural injection 4.0mL of lidocaine only.		atment, 1,3, 6 nths	after treatment, 1, 3 and 6 mo, Grp 1 greater reduction, p<0.05 ODI 1, 3 and 6 mo, Grp 1 greater reduction, p<0.05 Grp 1 vs Grp 3 VAS after treatment, 1, 3 and 6 mo, Grp 1 greater reduction, p<0.05 ODI 1, 3 and 6 mo, Grp 1 greater reduction, p<0.05 Grp 2 vs Grp 3 VAS after treatment, 1, 3 and 6 mo, no significant difference, p>0.05 ODI 1, 3 and 6 mo, no significant difference, p>0.05 ODI 1, 3 and 6 mo, no significant difference, p>0.05 ODI
Karm 2018	44 patients total, 20 in the RACZ group (age 66.1 +-12.2, male 9 (45.0%), and 24 in the ZiNeu group (Age 65.5 +-6.4 18 females, 26 males. Setting: Single- center, academic, outpatient interventional pain management clinic in Korea	1) 2)	PEA Using a Balloon-less Catheter (Racz) (n = 20) Percutaneous Epidural Decompression and Adhesiolysis Using an Inflatable Balloon Catheter (ZiNeu) (n = 24)	2) 3) Fol	NRS (back pain) NRS (leg pain) ODI llow-up: 1, 3 1 6 months	1, 5 and 6 mo, no significant difference, p>0.05 Between group MD, 95% CI, p values NRS-11 (Back pain) 1 mo:-0.38 (-1.81 to 1.06): p=0.61 3 mo: -1.13 (-2.63 to 0.38): p=0.14 6 mo: -2.02 (-3.58 to 0.45): p=0.01 NRS-11 (Leg pain) 1 mo: 0.73 (-0.40 to 1.85): p=0.21 3 mo: -0.69 (-1.89 to 0.52): p=0.26 6 mo: -1.88 (-3.15 to 0.61): p=0.00 ODI (%) 1 mo: -6.13 (-13.88 to 1.61): p=0.12 3 mo: -6.63 (-14.75 to 1.48): p=0.11 6 mo: -13.74 (-22.18 to 5.30): p=0.00 Adverse events: Minor and transient adverse events were reported equally in both groups (no data provided), mostly pain and paresthesia at the injection site.
				Surg	gery	
Zucherman 2004, 2005, 2006	191 subjects, 57% male and 43% female in the X STOP group. 52% male	1) 2)	X STOP Interspinous Process Decompression System (n=100) Non-operative treatment: Subjects received an epidural steroid injection	1) 2) 3)	SF-36 ZCQ Worker's compensation claims	Patient global assessment (Good result)2 yrs: 73.1% (surgery) vs. 35.9% (control) (P< 0.001)

operative group. Average age of 70 years in the X STOP group and 69.1 years in the non-operative group. Average of 3.5 year symptom duration in the X STOP group and 4.7 years in the non-operative group. Setting: Spine center in the	well as NSAIDS, analgesic agents, and physical therapy. Physical therapy consisted of education on back care and modalities such as ice packs, heat packs, massage, stabilization exercises, and pool therapy. Braces such as abdominal binders and corsets were permitted, but body jackets and chair back braces were not. (n=91)	changes Follow-up: Surgery: 7 (2 yr) Control: 19 (2 yr)	 "Clinically relevant improvement (patients)": 2 yrs: 60.2% (surgery) vs. 18.5% (control) (P< 0.001) Symptoms Severity score†† Surgery better at 6 w, 6 mo, 1 and 2 yr (graphs) (P<0.001) 2 yrs: MPC 44.3% (surgery) vs0.4% (control) (P < 0.001) "Clinically relevant improvement (as measured by patients)": 2 yrs: 57% (surgery) vs. 14.8% (control) (P < 0.001) ZCQ (global success) 6 mo: 52% (surgery) vs. 9% (control) (P value not reported) 1 yr: 59% vs 12% (P value not reported) 2 yrs: 48.4% (surgery) vs. 4.9% (control) (P < 0.001) Quality of life (SF-36) At all post treatment time points (6 w, 6 mo, 1 yr, 2 yr), the mean domain scores documented in the X STOP group were significantly greater than those in the non operative group, with the exception of the mean General Health, Role Emotional, and
2007, 2009, image-confirmed Abdu 2018 degenerative spondylolisthesis:	 Assigned to surgery (standard laminectomy with or without fusion) (n=159) Assigned to non-surgical treatment: Usual non-operative care (n=145) 	 SF-36 bodily pain SF-36 bodily function low back pain bothersomeness scale Leg pain bothersomeness scale ODI Subjective self- 	Mental Component <i>Summary scores at 2 years</i> Adverse events: No complications were reported in group 2. In group 1, complications were reported in 11% of subjects including spinous process fracture, coronary ischemia, respiratory distress, hematoma, and 1 death (pulmonary edema) All between group comparisons using Intention-to-Treat analysis SF-36 Bodily Pain, DMC, 95% CI 2 yrs: 1.5 (-4.2 to 7.3) 4 yrs: -2 (-8.6 to 4.6) 8 yrs: p=0.85 SF-36 Bodily Function, DMC, 95% CI 2 yrs: 1.9 (-3.7 to 7.5) 4 yrs: -3.1 (-9.2 to 3.0) 8 yrs: p=0.31 Disability (ODI), DMC, 95% CI 2 yrs: 2.2 (-2.3 to 6.8)

	age of 64.7 years			r	eported	4 yrs: 4.1 (-0.8 to 9.1)
	in the surgical				mprovement,	8 yrs: p=0.039
	group and 68.2				atisfaction with	8 yis. p=0.039
	years in the non-				urrent	Other outcomes (patient's satisfaction; Stenosis Bothersomeness
	•			-		Index, Leg Pain Bothersomeness Scale; and Low Back Pain
	surgical group.				ymptoms and	
	Subjects had			-	are	Bothersomeness Scale) were not provided separately for the
	symptoms for at				Stenosis	randomized cohort.
	least 12 weeks				othersomeness	Adverse events: group 1 reported 14% intraoperative
				11	ndex	complication mostly and dural tears and 19% postsurgical
	Setting: multi-					complications including 1 death, 11% required additional
	centred				llow-up: 6	surgeries at 2 years,
	orthopaedic				eks, 3 and 6	
	departments in			mo	nths, 1, 2, 4 and	
	the United States			8 y	ears	
Amundsen	100 subjects, 54	1)	Surgery: Partial or total	1)	VAS	Patient global assessment (Good result)
2000	male, 46 female,		laminectomy, medial facetectomy,	2)	Verbal Rating	1 yr: RR 2.07 (0.98 to 4.38)
	median age of 59		discectomy, and/or removal of		Scale	4 yrs: RR 1.94 (1.14 to 3.31)
	(males were 1.5		osteophytes from the vertebral	3)	Subjective	10 yrs: RR 3.18 (0.97 to 10.41)
	years higher than		margins or facet joints. No fusions.		change	Pain (none or mild)
	females).		(n=13)		(better, worse,	1 yr: NR
	Median back pain		(),		or unchanged)	4 yrs: RR 3.33 (0.77 to 14.33)
	duration was 14	2)	Conservative therapy: Lumbar	4)	Work status	10 yrs: RR 1.59 (0.55 to 4.55)
	years, median	_,	orthosis use for 1 month worn during	5)	Subjective	
	duration of		the day for all activities plus	5)	rating from	Other outcomes (claudication or walking distance; level of daily
	sciatica was 2		instruction and back school." (n=18)		evaluating	activity; and neurologic deficits) were not reported separately
	years.		instruction and back school. (if 10)		physician and	for the randomized cohort.
	years.				study team	for the fandomized conort.
	Setting:				(Excellent,	
	0				Fair,	
	Neurology department in a				,	
					Unchanged,	
	hospital in				Worse)	
	Norway			17.1	n	
					llow-up:	
					nonths, 1, 4 and	
					years	
Malmivaara	94 subjects, 22%	1)	Segmental decompressive surgery	1)	11 point	All between group comparisons
2007	of surgical		with facetectomy (n=50)		numerical pain	Leg pain, MD, 95% CI

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subjects were male, 45% of non-operative subjects were male. Nonoperative group had average age of 62.9 years, surgical group had average age of 63.9 years. Surgical group averaged 14 years since onset of symptoms, nonsurgical group average 16 years since onset of symptoms. Minimum of 6 months of symptoms for study inclusion. Setting: Research Center in Finland	 2) Non-operative treatment: NSAIDS when indicated and seen one to three times by a physiotherapist, in addition to the standard visit at each follow-up. The physiotherapist gave all patients educational brochure. The patients were encouraged to use their back in a normal way. Pain-relieving body postures were taught as well as basic ergonomics related to lifting and carrying. Individually structured programs included trunk muscle endurance and stretching-type exercises. Additional individual physiotherapy consisting of passive treatment methods (such as ultrasound and transcutaneous nerve stimulation). (n=44) The patients in the surgical group also received the brochure and the instructions described above. 	 rating scale for back and leg pain 2) Walking ability (distance without a break) also via treadmill test 3) General health status on a 5 point scale (very good, quite good, average, quite poor or very poor. 4) ODI 5) Ability to complete certain activities of daily 6) living without difficulty, some difficulty, marked difficulties or not at all 7) Radiographic examination 	1 yr: 1.69 (0.41 to 2.96) 2 yr: 1.51(0.25 to 2.77) Back pain, MD, 95% CI 1 yr: 2.33 (1.12 to 3.55) 2 yrs: 2.13(0.98 to 3.28) Disability (ODI), MD, 95% CI 1 yr: 11.3 (4.3to 18.8) 2 yrs: 7.8 (0.8 to14.9) > 10 points reduction (ODI): RR, 95% CI 1 yr: 2.16 (1.31 to 3.57) 2 yrs: 1.36 (0.88 to 2.10) Walking disability (walking distance <1.250 m), RR, 95% CI 1 yr: 0.93 (0.61 to 2.03) 2 yrs: 1.08 (0.70 to 2.42) Walking disability (walking distance <400 m), RR, 95% CI 1 yr: 0.91 (0.51 to 4.24) 2 yrs: 1.18 (0.67 to 4.72)
		Follow-up: 6 months, 1 and 2 years	

Weinstein 2008, 2010,	289 in the RCT, 365 in the	1)	Assigned to surgery: Standard laminectomy with or without fusion	1)	SF-36 bodily pain	All between group comparisons using Intention-to-Treat Analysis
Lurie 2015	observational cohort. 62% male		(n=138)		SF-36 bodily function	SF-36 Bodily Pain, DMC, 95% CI 2 yrs: 7.8 (1.5to 14.1)
	in the surgical groups, 59% male in the non-	2)	Assigned to non-surgical treatment: Usual non-operative care - recommended to include at least	3)	Low back pain bothersomene ss scale	4 yrs: 0.3 (-6.4 to 7) 8 yrs: p=0.25 SF-36 Bodily Function, DMC, 95% CI
	surgical groups. Average age of 63.8 in the		active physical therapy, education or counseling with home exercise instruction, and the administration of	4)	Leg pain bothersomene ss scale	2 yrs: 0.1 (-6.4 to 6.5) 4 yrs: -3.2 (-9.9 to 3.6) 8 yrs: p=0.89
	surgical group, 66.1 in the non- surgical group.		NSAIDs, if tolerated (n=151)	5) 6)	ODI Subjective self-reported	Disability (ODI), DMC, 95% CI 2 yrs: -3.5 (-8.7 to 1.7) 4 yrs: 0.2 (-5.2 to 5.7)
	60% in the surgical group and 55% in the				improvement, satisfaction with current	8 yrs: p=0.87 Other outcomes (patient's satisfaction; Stenosis Bothersomeness
	non-surgical group had symptoms for			7)	symptoms and care, Stenosis	Index, Leg Pain Bothersomeness Scale; and Low Back Pain Bothersomeness Scale) were not provided separately for the randomized cohort.
	over 6 months.			')	bothersomene ss index	Adverse events: In group 1, 10% of patients required
	Setting: multi- centred- orthopaedic			Follow-up: 6 weeks, 3 and 6		transfusions intraoperatively and 5% postoperatively. The most common surgical complication was dural tear, in 9% of patients. At 2 years, reoperation had occurred in 8% of
	departments in the United States.			yea		subjects.
Delitto 2015	169 patients, 88 males and 81 females, 87	1)	Surgical decompressive laminectomies, partial facet resection, and neuroforaminotomies	fun	SF-36 physical action	2 years -SF-36 Physical Function, MD and 95% CI 0.9 (7.9 to 9.6)
	surgical group with an average age of 66.6 years	2)	(n=87) PT program: lumbar flexion	Fo	llow-up: 2 years	Adverse events: 9 out of 82 participants in group 2 reported adverse events consisting of worsening of symptoms whereas 33 out 87 participants in group 1 reported surgery related
	old and 82 PT group with an average age of 69.8 years old,	,	exercises, exercises and education (n=82)			complications, mainly attributable to reoperation, delay in wound healing and surgical site infection.
	LSS by computed					

tomography -		
criteria of Wiesel		
and colleagues		
(18) or magnetic		
resonance		
imaging - criteria		
of Boden and		
colleagues (2)		
Setting:		
Neurologic and		
orthopedic		
surgery		
departments and		
physical therapy		
clinics in western		
Pennsylvania		· · ·

ADLs = Activities of Daily Living, AUC = Area under the pain-intensity curve, BTX = Botox, CI = Confidence Interval, DMC = Difference in mean change from baseline, ESI = Epidural Steroid Injection, FRI = Functional Rate Index, GRP = Group, HADS =Hospital Anxiety and Depression Scle, IU = International Units, JOABPEQ = Japanese orthopaedic association back pain evaluation questionnaire, LBOS = Low Back Outcome Score, LBP = Low Back Pain, m = Meters, MCS = Mental Component Score, MD = Mean Difference, mm = Millimeters, Mo = Months, MPC = Mean Percent Change, NRS = Numerical Pain Rating Scale, NR = Not Reported, ODI = Oswestry Disability Index, OR = Odds Ratio, PASS-20 = Pain Anxiety Symptoms Scale, PCS = Physical Component Score, RCT = Randomized Controlled Trial, RMDI = Roland Morris Disability Index, ROM = Range of Motion, RR = Relative Risk, SBI = Stenosis Bothersomeness Index, SPWT = Self-Paced Walking Test, SSS = Spinal Stenosis Questionnaire, TSK-11 = Tampa Scale-11, VAS = Visual Analogue Scale, WMD = Weighted Mean Difference, ZCQ = Zurich Claudication Questionnaire